SIERRA Multimechanics Module: Aria

Patrick K. Notz, Samuel R. Subia, Matthew M. Hopkins, Harry K. Moffat,
David R. Noble, Tolulope O. Okusanya

Prepared by
Sandia National Laboratories
Albuquerque, New Mexico 87185 and Livermore, California 94550

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Abstract

Aria is a Galerkin finite element based program for solving coupled-physics problems described by systems of PDEs and is capable of solving nonlinear, implicit, transient and direct-to-steady state problems in two and three dimensions on parallel architectures. The suite of physics currently supported by Aria includes thermal energy transport, species transport, and electrostatics as well as generalized scalar, vector and tensor transport equations. Additionally, Aria includes support for manufacturing process flows via the incompressible Navier-Stokes equations specialized to a low Reynolds number ($Re < 1$) regime. Enhanced modeling support of manufacturing processing is made possible through use of either arbitrary Lagrangian-Eulerian (ALE) and level set based free and moving boundary tracking in conjunction with quasi-static nonlinear elastic solid mechanics for mesh control. Coupled physics problems are solved in several ways including fully-coupled Newton’s method with analytic or numerical sensitivities, fully-coupled Newton-Krylov methods and a loosely-coupled nonlinear iteration about subsets of the system that are solved using combinations of the aforementioned methods. Error estimation, uniform and dynamic $h$-adaptivity and dynamic load balancing are some of Aria’s more advanced capabilities. Aria is based upon the Sierra Framework.
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Chapter 1

Introduction

1.1 Overview

The SIERRA Multimechanics Module: Aria, henceforth referred to as Aria for brevity, is an application implementing the finite element method (FEM) for solving systems of partial differential equations (PDEs). Foremost, Aria’s development targets applications which involve incompressible flow (Navier-Stokes for $Re < 1$). However, the general design of Aria lends itself to the solution of systems of PDEs describing physical processes including energy transport, species transport with reactions, electrostatics and general transport of scalar, vector and tensor quantities in two and three dimensions both transient and direct to steady state. Moreover, different regions of the physical domain (i.e., the input mesh) may have either different materials and/or different collections of physics (viz., PDEs) defined on them. These systems of equations may be solved alone, in a segregated but coupled algorithm (“loosely coupled”) or as a single, fully-coupled system. Additionally, Aria can be loosely coupled to the quasi-statics solid mechanics code Adagio using the coupling application Arpeggio.

Aria is able to accommodate meshes that utilize linear and quadratic elements in two and three dimensions. In two dimensions, Aria supports quadrilateral (4 and 9 node) and triangular (3 and 6 node) elements. In three dimensions, Aria supports hexahedral (8 and 27 node) and tetrahedral (4 and 10 node) elements. Moreover, meshes may be comprised of combinations of these elements (i.e., both quadrilateral and triangular elements in two dimensions).

The physical coordinates and mesh displacements are always interpolated in accordance with the input mesh, but other solution degrees of freedom may be interpolated using a lower order basis function. For example, if the input mesh is composed of 9 node (quadratic) quadrilateral elements, then the physical coordinates and mesh displacements (if active) will be interpolated using quadratic basis functions, whereas other degrees of freedom, e.g., temperature or voltage, could use linear shape functions.

Additional information concerning the project may be found at the Aria’s home page, Aria Users Home-page [1], and at Aria’s Trac wiki web site, Aria Trac Wiki [2]. Both of those web sites currently require access to Sandia’s internal restricted network.

1.2 A Brief History of Aria

In many respects, the predecessor code to Aria is the highly successful Goma code. Although none of the Goma code is included in Aria, Goma has inspired many of the algorithm, design and implementation decisions in Aria. Several of the staff members who contributed to Goma have also contributed to Aria, either as developers or users and testers.

Like Goma, Aria’s primary nonlinear algorithm is a full Newton-Raphson method with analytical sensitivities. However, Aria takes aspects of the implementation further by adding versatility in how the sensitivities are constructed and providing options for both finite-difference, and automatic differentiation sensitivities.
Furthermore, Aria has a richer suite of nonlinear solver capabilities including matrix-free Newton-Krylov, methods and even loose coupling. Like its muse Goma, Aria includes a large suite of physics and capabilities which can be included in an analysis.

On December 5, 2000 Phil Sackinger initiated the Opera code which was to serve as a test bed for what would become Aria. The Opera code was a patterned after the Calore code which is a thermal analysis GFEM code, but differed in many respects. At that time, no applications had been written on top of the Sierra Framework that used multiple degrees of freedom (e.g., velocity AND pressure) or that used multiple element type (e.g., linear and quadratic interpolations). Opera was created as an exploratory code, to determine what changes would need to be made to the Framework and to experiment with the Framework in constructing such an application. During this period, the principle developers of Opera were Phil Sackinger, Sam Subia and Matt Hopkins.

On November 20, 2001 Matt Hopkins made the first commit to the Aria code base, using Opera as the starting point. And thus Aria was born.

1.3 Nonlinear Coupling Strategies in Aria

One of the difficulties with writing broadly applicable computational mechanics software is that developers can’t take advantage of specific knowledge of the application domain in order to optimize the algorithm. Thus, in providing generality one sometimes sacrifices efficiency. One place this is evident in multiphysics modeling is in the choice of coupling strategies. While it is well understood that a fully coupled system solved with Newton’s method utilizing analytic sensitivities is formally the most robust and correct approach to solving multiphysics applications it is also computationally expensive and complex to implement. Furthermore, while Newton’s method has the fastest rate of asymptotic convergence it’s domain of convergence is often empirically observed to be smaller than other methods. Lastly, in some applications, certain subsets of the physics may be only weakly coupled so that a loosely coupled approach may be more computationally efficient. To address these concerns while remaining general and flexible Aria offers a number of options for nonlinear solution strategies and physics coupling.

In defining a problem in Aria, users configure one or more Regions. Each Region consists of one or more PDEs to be solved on some or all of the input mesh. All of the PDEs in each Region are solved in a tightly coupled (i.e., single matrix) manner using one of several nonlinear solution strategies available. Users may then define loose couplings between two or more Regions. For example, some or all of a solution from one Region may be transferred to another Region where it is treated as a constant, external field. The aggregate nonlinear problem including the contributions from all of the Regions may be iterated to convergence. The particulars of which physics are solved in each Region and the nonlinear solution strategy used within and between Regions is completely specified through the input file. Furthermore, an Aria user may pick a simple, minimal algorithm without needing to fit it into an overly-generalized worst-case scenario that represents the union of all possible algorithms.

Dynamically-specified loose coupling has many potential advantages that users may leverage. First, the resulting linear system is considerably smaller and contains far fewer off-diagonal contributions which can significantly increase the performance of linear solvers. Also, a resulting linear system may have a more attractive form, such as symmetric positive-definite, that permits the use of tailored iterative solutions techniques. Other extensions to loose coupling include subcycling of transient simulations where each Region may advance in time with its own time step size and in-core coupling to other applications based upon the Sierra framework.
1.4 Constraints Equations within Aria

Aria has a unique capability associated with the specification of global constraint equations. These may be used to specify conserved quantities that are not specifically specified as part of the equations set. For example, in some electrochemistry problems where current is specified as a boundary condition, the global conservation of charge neutrality must be imposed as an additional global condition.

Constraint equations have unique issues associated with their solution.

1.5 Level Set Algorithm

Level set algorithms utilize a signed distance function $F$ such that one material, or phase, is associated with regions of space where $F > 0$ and a different phase is associated with regions of space where $F < 0$. The curve or surface where $F = 0$ defines the interface between the two phases. In Goma and most other level set codes $F$ is used to partition material property models such that the property has the appropriate values in each phase and, typically, transitions smoothly from one phase to the other. In Aria, however, $F$ is used to partition contributions of the residual equations between the two phases.

In both cases the partitioning is done using a Heaviside function to partition the physical space into two phases which we’ll label $A$ and $B$. The Heaviside function $H_A(F)$ is defined such that $H_A(F) = 1$ in phase $A$ and $H_A(F) = 0$ in phase $B$; in the vicinity of $F = 0$ the Heaviside function may be defined to be a smooth function that transitions from 0 to 1. Likewise, $H_B(F) = 1$ in phase $B$ and $H_B(F) = 0$ in phase $A$. In fact $H_B(F)$ is defined as $H_B(F) \equiv 1 - H_A(F)$.

In Goma, this Heaviside function is used to partition the material properties such that a material property $\sigma$ is defined as

\[ \sigma(F, \ldots) = \sigma_A(\ldots)H_A(F) + \sigma_B(\ldots)H_B(F). \tag{1.1} \]

In Aria, however, the integrand of each residual equation is multiplied by the sum of Heaviside function so as to decompose the equation into contributions from each phase,

\[ \int_V (\ldots) \, dV \rightarrow \int_V H_A(F)(\ldots) \, dV + \int_V H_B(F)(\ldots) \, dV. \tag{1.2} \]

This formulation has a number of advantages. Material models are not functions of $F$ so no special models need to be written and the input syntax is the same as well. Secondly, this approach is conservative for conservative governing equations. For example, the MASS and ADVECTION terms of the energy equation (see section 5.3) are proportional to $\rho C_p$ and hence, in Goma’s formulation, proportional to $H^2(F)$ where as the DIFFUSION term is proportional $\kappa$ and hence proportional to $H(F)$. Thus, in the vicinity of the interface $F = 0$ energy is not transported correctly between these transport modes.

In Aria, each assembly kernel has an arbitrary list of coefficients that multiply the integrand of the kernel (see section 42.13). Thus, the formulation depicted in equation 1.2 is accomplished by simply adding the appropriate Heaviside function to the list of coefficients for each kernel associated with the equation.

1.6 “Production” vs. “Experimental” Code and Known Issues

Aria is a unique application code in that it contains both production and experimental code capabilities in the same codebase. This manual makes little effort to differentiate between “production” and “experimental” capabilities, though generally capabilities which are more comprehensively documented can be considered
production-ready. Though the vast majority of features in Aria are well-tested, there may be cases where combinations of “production” features may not function correctly together. When in doubt, one may always refer to sierra-help for guidance.

A compilation of known issues are contained in release notes for each major code release which occurs roughly every six months.

1.7 Outline of the Manual

In chapter 2 we will discuss the overall environment for running Aria applications, including the layout for the Aria input deck. In chapter 5 we will present the general equations that are solved by Aria. These should be read by every user.

In later chapters, we will delve down to discuss individual line commands of the input deck. Chapter 6 discusses equation line cards (i.e., EQ), which serve to add individual equations with coupled independent unknowns to a coupled PDE representation of a region. Chapter 8 discusses how to apply initial conditions to the field variables associated with the equation sets. Chapter 9 presents the line commands associated with specifying boundary conditions. Chapter 10 introduces the concepts associated with distinguishing conditions. Chapter 11 introduces line commands associated with source terms.
Chapter 2

Getting Started

2.1 Setting The Environment-Users External to Sandia Labs

To access Sierra/Aria one will likely need to setup the user environment. This setup will differ upon location and the local system administrator can provide information on setting up your local environment.

2.2 Setting Up Your Environment-Users at Sandia Labs

The environment for using Aria is the same as for individual Sierra applications and can be configured by module files. The modules ensure that the look and feel of running Sierra applications is the same across a multitude of compute platforms. To obtain the proper environment for code execution one simply runs:

$ module load sierra

2.3 Running Aria

This section includes some very simple examples of how to run Aria. For more information on running on some of Sandia's clusters, etc. see [3].

In its simplest form, Aria can be run like this:

$ sierra -np 1 aria -i ariarun.i

In this example, ariarun.i is the Aria input file. The output – nonlinear iterations, time step information, etc. – will be written to a file called ariarun.log. So, you can monitor the progress of the simulation by watching the log file. Alternatively, you can have all of the output sent to the screen by using the -l logfile command line option. If you set the log file to be - (a single “minus” character) all of the output will be sent to the standard output (usually your screen):

$ sierra -np 1 aria -i ariarun.i -l -

If you would like to use aprepro in your input file, add the -a command line option to have your input file automatically processed:

$ sierra -np 1 aria -i ariarun.i -l -a
Oftentimes we want to run Aria remotely or locally in a batch mode, save any standard output and perhaps even log out from a session. Unfortunately, termination of the session through either voluntary (interactive) or involuntary (timeout) log-out may in effect terminate the Aria job. In this case one can prevent the job from terminating by using the Unix nohup command in conjunction with the standard execution command line.

$ nohup sierra -np 1 aria -i ariarun.i -l YourLogFile -a

If one wishes to run the job in a background mode the nohup command should be terminated with & at the end of the command line.

### 2.4 Aria Environment Overview

Aria is a Sierra application implementing the finite element method (FEM) for solving systems of partial differential equations (PDEs). Foremost, Aria’s development targets applications which involve incompressible flow (Navier-Stokes). However, the general design of Aria lends itself to the solution of systems of PDEs describing physical processes including energy transport, species transport with reactions, electrostatics and general transport of scalar, vector and tensor quantities in two and three dimensions both transient and direct to steady state. Moreover, different regions of the physical domain (i.e., the input mesh) may have either different materials and/or different collections of physics (viz., PDEs) defined on them. These systems of equations may be solved alone, in a segregated but coupled algorithm (“loosely coupled”) or as a single, fully-coupled system. Currently, Aria’s loose coupling capabilities are handled by the Arpeggio application which also allows Aria to couple (loosely) to the quasistatic structural mechanics code, Adagio.

Aria’s models and algorithms are integrated into the Sierra framework through the architecture illustrated in Figure 2.1. A Sierra-based application has four layers of code: Domain, Procedure, Region, and Model/Algorithm.

The outermost layer of an application is the Domain, or “main” program of the application. This domain layer is implemented by the Sierra Framework to manage the startup/shutdown of an application, and to orchestrate the execution of an application-proved set of procedures.

Code at the Procedure level is responsible for evolving one or more s loosely coupled sets of physics through a sequence of steps. This sequence may be a set of time steps, nonlinear solver iterations, or some combinations of these or other types of steps.

An application may define multiple procedures to implement hand-off coupling between physics within the same main program. In hand-off coupling the first (or preceding) procedure completes execution, mesh and field data is transferred to a succeeding procedure, and the succeeding procedure continues the simulation with a different set of physics. For example, the first thermal procedure could calculate a temperature distribution inside a differentially heated fluid, and the second procedure could simulate natural convection of the fluid due to the density gradients set up by the resulting temperature field.

Code at the region level is responsible for evolving a tightly coupled set of physics through a single step. Loose coupling of Regions is supported by the advanced transfer services provided by the Sierra framework.

Each region owns (1) a set of models or algorithms that implement its tightly coupled set of physics and solvers and (2) an in-memory parallel distributed mesh and field database. This mesh and field data is fully distributed among parallel processors via domain decomposition.
2.5 Overview of the Input File Structure

An Aria model is described by commands contained in an ASCII input file. The structure of the input file follows a nested hierarchy. The topmost level of this hierarchy is named the domain. Underneath the domain is a level called the procedure, followed by the region level. Figure 1.1 shows this nesting.

The domain level contains one or more procedures. At the domain level, you also find commands associated with describing the finite element mesh, the linear solver set-up, material properties associated with a defined material, and user functions associated with source terms and boundary conditions that are added into Aria’s intrinsic set of functions.

The procedure level contains one or more regions. The procedure level is also used to specify the time stepping parameters, and interactions between regions, such as data transfers. Essentially at the procedure level, loose coupling algorithms are specified. Loose coupling here is defined within the context of Aria’s implicitly full-coupled paradigm. Whenever an independent variable’s interaction with other variables in the solution procedure is not fully represented in the global matrix, the algorithm for loose coupling of that variable and its associated equation will be described at the procedure level. This loose coupling algorithm is given a fancy name called a “solution control description”. The procedure level contains a block specifying the solution control procedure. An analogy to this block in simpler codes would be top level loop. For example in time dependent applications, the solution control description block would involve a block to solve the time dependent problem repeated for each time step until the desired solution time is reached.

The region level is used to specify details about the tightly coupled equation system to be solved. The details include boundary conditions and initial conditions, where materials models are applied, and where surface and volumetric source terms are applied. Essentially, meshes and material properties described at the domain level are tied into the problem statement here via their names.

Global constraints equations are also specified at the region level. At the region level, specification of what gets sent to the output file and at what frequency also is made. Additional post-processing associated with the output is specified. For example, additional volumetric fields which are functions of the independent variables may be specified to be added to the output file.
There are two types of commands in the input file. The first type is referred to as a block command. A block command is a grouping mechanism. A block command contains a set of commands made up of other block commands and line commands. A line command is the second type of command. The domain, procedure, and region levels are all parsed as block commands. A block command is defined in the input file by a matching pair of Begin and End lines. For example,

```
Begin SIERRA myJob
    ... block commands
End SIERRA myJob
```

A set of key words for the block command follows the “Begin” and “End” keywords. In most cases a user-specified name is added to the block commands. In the example above the keywords, SIERRA myJob, are added. Optionally, the keyword may be left off of the end of the block.

The second type of command is the line command. A line command is used to specify parameters within a given block command. In the remaining chapters and sections of this manual, the scope of each block and line command is identified, along with summaries of the meanings. Note that the ordering of any commands within a command block is arbitrary. Thus,

```
Begin Finite Element model fluid
    Database name is pipeflow2d.g
    Use Material water for block1
End Finite Element model fluid
```

will have the same effect as

```
Begin Finite Element model fluid
    Use Material water for block1
    Database name is pipeflow2d.g
End Finite Element model fluid
```

And the ordering of command blocks within the domain/procedure/region blocks are arbitrary—allowing you considerable freedom to collect and arrange commands. Note that the terms “command block” and “block command” are interchangeable.

The sierra command block must contain a block for a procedure containing an aria region:

```
Begin procedure myProcedureName
    .
    Begin Aria region myRegionName
    .
    End Aria region myRegionName
End procedure myProcedure
```

The procedure command block is used to contain all of the Aria commands that are associated with a solution procedure defined for a set of Aria Regions. The `myProcedureName` and `name` keywords of the procedure and region blocks are left up to you. Note that the Aria procedure command block must be present in the input file and must contain at least one Aria region command block. The procedure command block also contains other important command blocks such as the SOLUTION CONTROL block.
2.5.1 Syntax Conventions for Commands

In this section we describe the conventions used in presenting all the command descriptions in the remainder of this manual. There are four basic kinds of tokens, or words, that Aria expects to find as it parses an input file. These are **keywords, names, parameters and delimiters**.

**Keywords**

The words which distinguish one block command, or line command, from another we term keywords. Keywords are denoted in this manual in the monospaced font, for example, `BOUNDARY CONDITION`.

**Names**

The word, or words, that you supply on the same line of the `begin` line of a block command, is the **name**. Many times you may need to supply this name as a character parameter in a separate line command. Names are denoted in italics, `name`, as are parameters.

It is worth noting that the interpreter used to process standard input command lines is also used to process lines defining algebraic operations. This means that a `"-"` appearing within a name would be interpreted as a subtraction operation and as a consequence, the use of `"-"` within a name is not allowed. Thus instead of

```
Begin Aria region name-1
```

one could perhaps use

```
Begin Aria region name_1.
```

**Parameters**

There are three types of input parameters one will need to supply to line commands: character strings, integers, and real numbers. These are denoted in the documentation as `(C)`, `(R)`, and `(I)`, respectively.

In most cases character strings may be specified in a free format. One exception to this paradigm is when a string begins a number. In this case the character string must be specified within quotation marks in order to be properly interpreted.

Real numbers may be entered in decimal form or exponential form. For example `0.0001`, `.1E-3`, `10.0d-5` are all equivalent. Furthermore, if a real(R) is expected, an integer can be used.

Integer values (I) need not include a decimal point in their specification.

**Multiple Parameters**

For the case when a list of one or more parameters is allowed, or required, for a command, `(C,...)` denotes a list of character strings, `(I,...)` a list of integers, and `(R, ...)` a list of real numbers. For a list of character strings, the separator between the strings must be one or more spaces or tab characters. Therefore, phrases with multiple spaces and words in them are tokenized into multiple character parameters before being processed by the application. For a list of real or integer numbers the comma can also be used as a separator.
Enumerated Parameters

Certain commands have predefined parameters, called *enumerations*, which are listed within `{}`. Each parameter in the list is separated using `|`. The default parameter for the list of parameters is enclosed by `<>`.

Delimiters

The keywords of a line command are often required to be separated from the parameters by a delimiter. You have a choice of delimiters to use: the equal sign, `=`, or a word. In this manual, we denote the choices surrounded by `{}`, and separated by `|`. You may use any one of the delimiters from those listed. For example, the line command to specify the density within the Aria Material Block command is

```
Density {=} |IS} (R)
```

Examples of valid forms you could write in the input file are

```
Begin Aria Material water
...
Density = constant rho = 1.0E-3 # kg/m\(^3\) at 20C
...
End
```

and

```
Begin Aria Material water
...
Density is constant rho = 1.0E-3 # kg/m\(^3\) at 20C
...
End
```

White Space

Command keywords, names, and parameters and delimiters must have spaces around them.

Indentation

All leading spaces and/or tab characters are ignored in the input file. Of course, we recommend that you use indentation to improve the readability for yourself and others that may need to see your files.

Including Files

External text files containing input commands can be included at any point in the Aria input file using the `INCLUDEFILE` command. This command can be used in any context in the input file. To use this command, simply use the command `INCLUDEFILE` followed by the name of the file to be included. For example, the command:
INCLUDEFILE extrafile.i

would include the contents of extrafile.i at the locations where it is included in the input file. The included file is contained in the standard echo of the input that is provided at the beginning of the log file.

NOTE: Though this line command works well in many simple circumstances, it is known to cause issues when file names are involved. The most robust method to include files is to use Aprepro's \texttt{include} function and pre-process the input file with aperpro before running. An example is below:

\#{include(extrafile.i)}

Case Sensitivity

None of the command keywords, parameters, or delimiters read from the input file are case sensitive. For example, the following two lines are equivalent:

\begin{verbatim}
Use Material water for block_1
.
\end{verbatim}

and

\begin{verbatim}
USE material wATer for block_1
.
\end{verbatim}

The exception to this rule are file names used for input and output, because the current operating systems on which SIERRA applications are run are based on UNIX, where file names are case sensitive.

Comments and Line Continuation

You may place comments in the input file starting with either the $ or \# character. All further characters on a line following a comment character are ignored.

You can continue a command in the input file to the next line by using the line continuation character, $, or you may optionally following it with a comment, \#. All further characters on the same line following a line continuation character $ are ignored, and the characters on the following line are joined and parsing continues. An example is the line command used to specify the title of a thermal model:

\begin{verbatim}
Begin SIERRA Job_Indentifier
  # This thermal model for Aria simulates a convective heat transfer
  Title The title command is used to set the analysis title \\
  Convective heat transfer to a part. The analysis makes use of conjugate heat transfer to account for \\
  cooling of a part due to flowing water.
  ...
End SIERRA Job_Indentifier
\end{verbatim}
Checking the Syntax

Errors in the input deck can be checked by adding the command, -check-syntax to the aria command line. For example,

$ sierra -np 1 aria -check-syntax -i input.i

This command will print the code echo of the input deck and any syntax errors within it to the screen.

2.6 Fields

Fields are defined as variables which are distributed on mesh objects. For example, if the temperature is defined via Q1 interpolation on a 2D mesh consisting of quadrilaterals, then the vector of nodal temperature coefficients that make up the interpolation would be defined as the Temperature field on that mesh. Fields may be defined on any mesh object type (e.g., elements, faces, edges, nodes, node sets, and side sets), not just at nodes.

The mesh object and field data may be distributed among parallel processors via a domain decomposition algorithm. Both fields and meshes are owned at the region level. A particular field may or may not be part of Aria’s solution vector for the particular region. However, all fields in Aria’s solution vector are fields defined on the mesh for that region.

2.6.1 Field String-Naming Convention

Due to the dynamic nature of fields and variables in Aria a consistent naming convention must be used for sanity sake. This section describes the format of string-names of Aria Expressions. These string forms are used for input and output only; Aria has more efficient internal structures for referencing Expressions.

Briefly, the overall format is described in Figure 2.6.1.

Valid values of the <Operator> field are listed in Table 2.6.1. Valid values of the <Name> field are too numerous to list here; they include things like degrees of freedom (VELOCITY, SPECIES) and material properties (VISCOSITY, ELECTRICAL_CONDUCTIVITY). The <Subindex> field can be used to designate multiple instances of a field. This is typically used for species equations. All integer values are valid subindex values but it’s best to use values ≥ 1. The <Phase> field is used in level set problems. Some fields are present in “all phases” while others, such as material properties, depend on which phase is being referred to. The <Component> field allows the user to specify a particular component of vector and tensor fields; valid values are described in Table 2.6.1.
<table>
<thead>
<tr>
<th>Operator</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>(none)</td>
<td>“No-Op”, no-operator</td>
</tr>
<tr>
<td>DT</td>
<td>Time derivative</td>
</tr>
<tr>
<td>GRAD</td>
<td>Gradient</td>
</tr>
<tr>
<td>DIV</td>
<td>Divergence</td>
</tr>
<tr>
<td>DET</td>
<td>Determinant of a 2-tensor</td>
</tr>
<tr>
<td>DETJ</td>
<td>Determinate of the Jacobian of transformation</td>
</tr>
<tr>
<td>SURFACE_DETJ</td>
<td>Determinate of the Jacobian of transformation</td>
</tr>
<tr>
<td>REF_FRAME</td>
<td>“No-Op” in the undeformed reference frame</td>
</tr>
<tr>
<td>GRAD_REF_FRAME</td>
<td>Gradient in the undeformed reference frame</td>
</tr>
<tr>
<td>DIV_REF_FRAME</td>
<td>Divergence in the undeformed reference frame</td>
</tr>
<tr>
<td>DETJ_REF_FRAME</td>
<td>Determinate of the Jacobian of transformation in the undeformed reference frame</td>
</tr>
<tr>
<td>SURFACE_DETJ_REF_FRAME</td>
<td>Determinate of the Jacobian of transformation in the undeformed reference frame</td>
</tr>
<tr>
<td>OLD</td>
<td>“No-Op” at the previous time step</td>
</tr>
<tr>
<td>GRAD_OLD</td>
<td>Gradient at the previous time step</td>
</tr>
<tr>
<td>DIV_OLD</td>
<td>Divergence at the previous time step</td>
</tr>
</tbody>
</table>

**Table 2.1.** Valid values of the `<Operator>` prefix.

<table>
<thead>
<tr>
<th>Phase</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>(none)</td>
<td>A field present in all phases within a material</td>
</tr>
<tr>
<td>A</td>
<td>Phase A</td>
</tr>
<tr>
<td>B</td>
<td>Phase B</td>
</tr>
<tr>
<td>C</td>
<td>Phase C</td>
</tr>
</tbody>
</table>

**Table 2.2.** Valid values of the `<Phase>` suffix. Phase labels are used in level set calculations only.

<table>
<thead>
<tr>
<th>Component</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>(none)</td>
<td>No specified component</td>
</tr>
<tr>
<td>X</td>
<td>First vector component</td>
</tr>
<tr>
<td>Y</td>
<td>Second vector component</td>
</tr>
<tr>
<td>Z</td>
<td>Third vector component</td>
</tr>
<tr>
<td>XX</td>
<td>(1,1) 2-tensor component</td>
</tr>
<tr>
<td>XY</td>
<td>(1,2) 2-tensor component</td>
</tr>
<tr>
<td>XZ</td>
<td>(1,3) 2-tensor component</td>
</tr>
<tr>
<td>YX</td>
<td>(2,1) 2-tensor component</td>
</tr>
<tr>
<td>YY</td>
<td>(2,2) 2-tensor component</td>
</tr>
<tr>
<td>YZ</td>
<td>(2,3) 2-tensor component</td>
</tr>
<tr>
<td>ZX</td>
<td>(3,1) 2-tensor component</td>
</tr>
<tr>
<td>ZY</td>
<td>(3,2) 2-tensor component</td>
</tr>
<tr>
<td>ZZ</td>
<td>(3,3) 2-tensor component</td>
</tr>
</tbody>
</table>

**Table 2.3.** Valid values of the `<Component>` suffix. In non-Cartesian coordinate systems these may refer to, for example, radial or angular components.
2.7 Equations

Equations are defined within an Aria region to represent a particular continuity equation to be solved. Within the Aria input deck, solution variables are assigned as the independent unknowns to equations. In general, there is a one-to-one correspondence between solution unknowns and equation degrees of freedom.

2.8 Equation String-Naming Convention

Similar to the field string-naming convention, equation names pose a similar requirement. This section describes the format of string-names of Aria equations. These string forms are used for input and output only; Aria has more efficient internal structures for referencing equations.

Briefly, the overall format is described in Figure 2.3.

Valid values of the \(<\text{Equation\_Name}>\) field are numerous and changing in time. Typical values include \text{MOMENTUM}, \text{ENERGY}, \text{SPECIES}, \text{LEVEL\_SET}, \text{MESH}, \text{CURRENT} and \text{VOLTAGE}; see chapter 6 for a complete description of existing equations. All integer values are valid subindex values but it’s best to use values \(\geq 1\) – currently -1 has a special meaning of “no subindex”. The \(<\text{Phase}>\) field is used in level set problems. Some fields are present in “all phases” while others, such as material properties, depend on which phase is being referred to. The \(<\text{Component}>\) field allows the user to specify a particular component of vector and tensor equation; valid values are described in Table 2.6.1.

2.9 Internal Field Definition

Internal to the Aria code Fields mentioned in the previous section 2.6.1 have alternative internal names that facilitate their algorithmic usage. In most cases this naming convention will be “usage”-\text{field\_name} where “usage” categorizes its internal usage. From a user perspective the “usage” category will be “solution” and as an example the voltage solution is internally known as solution-\text{\_voltage}. Application output is managed through Framework services independent of the application thus output requests must correspond to the internal code Field names rather than the Field name stated in the previous section.
2.10 User Fields

Situations often arise where one wishes to provide Field data storage so that data can be transferred into or out of Aria. The mechanism used for this capability is provided by the USER FIELD command line.

2.10.1 User Field

Scope:

User Field REAL_or_INTEGER NODE_or_FACE_or_ELEMENT SCALAR_or_VECTOR_or_TENSOR Variable_name On Mesh_extent [ Value {=|are|is} Magnitude... ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>REAL_or_INTEGER</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>NODE_or_FACE_or_ELEMENT</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>SCALAR_or_VECTOR_or_TENSOR</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Variable_name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Mesh_extent</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary

Defines a user field of standard type data_type (REAL or INTEGER) on the FEM entity (NODE, FACE, or ELEMENT) with the specified name and Sierra field type (SCALAR, VECTOR, or TENSOR). The field will be defined on the specified mesh part or mesh group alias. Normally this field would be supplied to Aria through a transfer.
Chapter 3

Model Definition

3.1 Model Overview

Sierra Framework services provide overall control of input commands, discretization input data and output data, IO. Additionally they provide a directed interaction of Framework services at the so-called Domain level with the application code at the Region level. This controlled interaction is enabled by commands that follow.

The model discretization (mesh) and the mesh components to be used in the model are defined at the Domain level and are later referenced by the application at the Region level. The association of material properties with portions of the mesh are also defined here within the Finite Element Model command block. In coupled analyses the need often arises for different discretizations of the same physical domain. In this case one must supply an independent Finite Element Model command block for each discretization since only a single mesh can be referenced within the command block.

Begin Sierra myJob
.
Begin Finite Element Model my_fem_model
.
End

Begin Global Constants
.
End

Model definition commands
.
Begin Procedure My_Aria_Procedure
.
Begin Aria Region My_Region
.
use Finite Element Model my_fem_model
.
End
.
End
.
End Sierra myJob
3.2 Finite Element Model

Scope: Sierra

```
Begin Finite Element Model Label

    Alias DatabaseName As InternalName
    Component Separator Character Option Separator
    Create GroupType NewSurfaceName Add SurfaceName...
    Coordinate System {=|are|is} CoordinateSystem
    Database Name {=|are|is} StreamName
    Database Type {=|are|is} DatabaseTypes
    Decomposition Method {=|are|is} Method
    Global Id Mapping Backward Compatibility Option1 Option2
    Omit Block BlockList...
    Omit Volume VolumeList...
    Time Scale Factor Option Scale
    Use Generic Names
    Use Material MaterialName For VolumeList...

Begin Parameters For Block Blockname
End

Begin Parameters For Phase Phase Name
End

Begin Parameters For Surface Surface_Name
End

End
```

Summary

Describes the location and type of the input stream used for defining a geometry model for the enclosing region.

### 3.2.1 Alias

Scope: Finite Element Model

```
Alias DatabaseName As InternalName

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>DatabaseName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>InternalName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>
```

Summary

Name the database entity "DatabaseName" as "InternalName"

Description

This "InternalName" may then be referenced in the data file in addition to the original name.
3.2.2 Component Separator Character
Scope: Finite Element Model

**Component Separator Character** Option Separator

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Separator</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary** The separator is the single character used to separate the output variable basename (e.g. "stress") from the suffixes (e.g. "xx", "yy") when displaying the names of the individual variable components. For example, the default separator is "_", which results in names similar to "stress_xx", "stress_yy", ... "stress_zx". To eliminate the separator, specify an empty string ("") or NONE.

3.2.3 Create
Scope: Finite Element Model

Create GroupType NewSurfaceName Add SurfaceName...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>NewSurfaceName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>SurfaceName</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary** Create a new set (node, edge, face, element, side/surface) as the union of two or more existing sets. The sets must exist in the mesh database or have been created by a previous CREATE command.

3.2.4 Coordinate System
Scope: Finite Element Model

Coordinate System {=|are|is} CoordinateSystem

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>CoordinateSystem</td>
<td>axisymmetric</td>
<td>barycentric</td>
</tr>
<tr>
<td></td>
<td>{axisymmetric</td>
<td>barycentric</td>
</tr>
</tbody>
</table>

**Summary** The interpretation of the geometry data stored in this database. Optional. Defaults to Cartesian.

3.2.5 Database Name
Scope: Finite Element Model

Database Name {=|are|is} StreamName

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>StreamName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>
3.2.6 Database Type

Scope: Finite Element Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Database Types</td>
<td>catalyst</td>
<td>dof</td>
</tr>
</tbody>
</table>

Summary: The database type/format used for the mesh.

3.2.7 Decomposition Method

Scope: Finite Element Model

Summary: The decomposition algorithm to be used to partition elements to each processor in a parallel run.

3.2.8 Global Id Mapping Backward Compatibility

Scope: Finite Element Model

Summary: (Unsupported, do not use)

3.2.9 Omit Block

Scope: Finite Element Model

Summary: Specifies that the element blocks named in the blockList be omitted from the analysis.

Description: If an element block is omitted, then it is illegal to refer to it later in the input file e.g an initial condition may not be specified on an omitted element block. The elements, faces, etc are never created and it is as if the omitted element blocks did not exist in the mesh file. If a surface is completely determined by the omitted element block, then it is illegal to specify boundary conditions on that surface. However, if the surface spans multiple element blocks, boundary conditions may be applied on the portion of the surface supported by the element blocks that are not omitted.
### 3.2.10 Omit Volume

**Scope:** Finite Element Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>VolumeList</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specifies that the volumes named in the volumeList be omitted from the analysis.

**Description**
If a volume is omitted, then it is illegal to refer to it later in the input file e.g an initial condition may not be specified on an omitted volume. The elements, faces, etc are never created and it is as if the omitted volumes did not exist in the mesh file. If a surface is completely determined by the omitted volume, then it is illegal to specify boundary conditions on that surface. However, if the surface spans multiple volumes, boundary conditions may be applied on the portion of the surface supported by the volumes that are not omitted.

### 3.2.11 Time Scale Factor

**Scope:** Finite Element Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Scale</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
The scale factor to be applied to the times on the mesh database. If the scale factor is 20 and the times on the mesh database are 0.1, 0.2, 0.3, then the application will see the mesh times as 2, 4, 6.

### 3.2.12 Use Generic Names

**Scope:** Finite Element Model

**Summary**
If this command is present then the name of all blocks and sets in the mesh will be of the form "type_id". For example, an element block with id=42 will be named "block_42"; a sideset with id 314 will be named "surface_314". If there are any names in the mesh file, those names will be aliases for the blocks and sets. If this command is not present, then if a name is in the mesh file, it will be used as the name and the generic generated name will be an alias. This is used a a workaround in codes that do not correctly handle named blocks and sets or as a workaround in meshes which contain non-user-specified names.

### 3.2.13 Use Material

**Scope:** Finite Element Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>MaterialName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>VolumeList</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Summary  
Associate the given volumes with the indicated material name.

### 3.3 Parameters For Block

**Scope:** Finite Element Model

```plaintext
Begin Parameters For Block Blockname
  Include All Blocks
  Local Coordinate System {=|are|is} Mesh Entities
  Material MatName
  Material = MatName
  Minimum Effective Dilatational Moduli Ratio {=|are|is} minEffectiveModuliRatio
  Minimum Effective Shear Moduli Ratio {=|are|is} minEffectiveModuliRatio
  Model {=|are|is} ModelName
  Nonlocal Regularization Kmeans Cell Size {=|are|is} kmeans_cell_size
  Nonlocal Regularization Kmeans Maximum Iterations {=|are|is} kmeans_maximum_iterations
  Nonlocal Regularization Kmeans Tolerance {=|are|is} kmeans_tolerance
  Nonlocal Regularization Partitioning Scheme {=|are|is} PartitioningScheme
  Phase PhaseLabel {=|are|is} MaterialName
  Remove Block {=|are|is} ExcludeBlockList...
End
```

Summary  
Specifies analysis parameters associated with each element block.

#### 3.3.1 Include All Blocks

**Scope:** Parameters For Block

Summary  
Use this parameters definition for all blocks.

When using this option within the FINITE ELEMENT MODEL command block the PARAMETERS FOR BLOCK will not use a Blockname.

#### 3.3.2 Local Coordinate System

**Scope:** Parameters For Block

```plaintext
Local Coordinate System {=|are|is} Mesh Entities
```

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mesh Entities</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  
Associate coordinate system with mesh entity.

Description  
Specify the local coordinate system to be used in conjunction with given element blocks.
3.3.3 Material
Scope: Parameters For Block

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>MatName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary     Associates this element block with its material properties.

3.3.4 Material =
Scope: Parameters For Block

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>MatName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary     Associates this element block with its material properties.

3.3.5 Minimum Effective Dilatational Moduli Ratio
Scope: Parameters For Block

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>minEffectiveModuliRatio</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary     Specifies a minimum effective DILATATIONAL moduli ratio. This value keeps the effective moduli from dropping below minEffectiveModuliRatio*ElasticModulus. This can aid in keeping the corresponding time step and bulk viscosity terms dropping to zero.

3.3.6 Minimum Effective Shear Moduli Ratio
Scope: Parameters For Block

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>minEffectiveModuliRatio</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary     Specifies a minimum effective SHEAR moduli ratio. This value keeps the effective moduli from dropping below minEffectiveModuliRatio*ElasticModulus. This can aid in keeping the corresponding hourglass stiffness terms dropping to zero.

3.3.7 Model
Scope: Parameters For Block
Model \{=|are|is\} ModelName

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>ModelName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Associates a solid mechanics material model with this element block. The material parameters for this block are specified in the material denoted by the MATERIAL command.

### 3.3.8 Nonlocal Regularization Kmeans Cell Size

**Scope:** Parameters For Block

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nonlocal Regularization Kmeans Cell Size {=</td>
<td>are</td>
<td>is} kmeans_cell_size</td>
</tr>
</tbody>
</table>

**Summary** This line command specifies the cell size used to construct the background grid for the computation of the centroidal Voronoi tessellation for the Kmeans partitioning scheme.

### 3.3.9 Nonlocal Regularization Kmeans Maximum Iterations

**Scope:** Parameters For Block

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nonlocal Regularization Kmeans Maximum Iterations {=</td>
<td>are</td>
<td>is} kmeans_maximum_iterations</td>
</tr>
</tbody>
</table>

**Summary** This line command specifies the maximum number of iterations to perform for Lloyd’s algorithm for the computation of the centroidal Voronoi tessellation for the Kmeans partitioning scheme.

### 3.3.10 Nonlocal Regularization Kmeans Tolerance

**Scope:** Parameters For Block

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nonlocal Regularization Kmeans Tolerance {=</td>
<td>are</td>
<td>is} kmeans_tolerance</td>
</tr>
</tbody>
</table>

**Summary** This line command specifies the relative tolerance for Lloyd’s algorithm. Iterations continue until the maximum number is reached or the L2 norm of a vector of all the center steps is less or equal than the tolerance times the cell size of the background grid.

### 3.3.11 Nonlocal Regularization Partitioning Scheme

**Scope:** Parameters For Block
Summary: This line command specifies the type of partitioning algorithm used to perform the domain decomposition for the nonlocal regularization method.

### 3.3.12 Phase

**Scope:** Parameters For Block

| Phase PhaseLabel {=|are|is} MaterialName |
|-----------------------------------------|
| Parameter                               | Value       | Default     |
| PhaseLabel                              | string      | undefined   |
| MaterialName                            | string      | undefined   |

Summary: Associate phase PhaseLabel with material Material_Name on this block.

### 3.3.13 Remove Block

**Scope:** Parameters For Block

| Remove Block {=|are|is} ExcludeBlockList... |
|-----------------------------------------|
| Parameter                               | Value       | Default     |
| ExcludeBlockList                        | string...   | undefined   |

Summary: List of blocks to exclude.

### 3.3.14 Axisymmetry Axis

**Scope:**

| Axisymmetry Axis {=|are|is} AxisType |
|---------------------------------------|
| Parameter                             | Value      | Default |
| AxisType                              | {x|y}      | X       |

Summary: When the COORDINATE SYSTEM = AXISYMMETRIC command appears within the Finite Element Model command block one can select at the Region level the coordinate axis about which the 2D axisymmetric sweep will occur. Note that there can be no overlap between the model and the axis of rotation.

### 3.4 Uniform Mesh Refinement

Produces initial uniform refinement for num iterations. This also precedes initialization of the Region, including setting initial conditions. This line command should be put inside the Aria Region.

**Syntax:**

```
INITIAL UNIFORM REFINEMENT FOR num ITERATIONS
```
3.5 Timing Overview

By default Sierra Framework services provide overall timing information for code execution. Alternatively, the user can increase the granularity of timing information by requesting the output explicitly using the PRINT TIMER command line. Here the timing information is itemized by code functionality and identify what portion of time is spent performing specific operations. Note that this command line must be applied at the Domain Scope (i.e. outside of the Procedure).

3.5.1 Print Timer Information Every

Scope:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Procedure-step-interval</td>
<td>integer</td>
<td>undefined</td>
</tr>
<tr>
<td>Checkpointed</td>
<td>{accumulated</td>
<td>checkpointed}</td>
</tr>
</tbody>
</table>

Summary: Specifies the procedure step count interval to print timer information

3.6 Global Constants

Scope: Sierra

```
Begin Global Constants empty
    Gravity Vector {=|are|is} Gravity1 Gravity2 Gravity3
    Ideal Gas Constant {=|are|is} Sigma
    K-E Turbulence Model Parameter Param {=|are|is} Value
    K-W Turbulence Model Parameter Param {=|are|is} Value
    Les Turbulence Model Parameter Param {=|are|is} Value
    Stefan Boltzmann Constant {=|are|is} Sigma
    Turbulence Model Param Number {=|are|is} Value
End
```

Summary: Set of universal constants for a simulation.

3.6.1 Gravity Vector

Scope: Global Constants

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gravity</td>
<td>real_1 real_2 real_3</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Gravity constant in vector form, acceleration components.
### 3.6.2 Ideal Gas Constant

**Scope:** Global Constants

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sigma</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** Ideal gas constant.

### 3.6.3 K-E Turbulence Model Parameter

**Scope:** Global Constants

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Param</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** $k - \epsilon$ RANS turbulence model parameters.

### 3.6.4 K-W Turbulence Model Parameter

**Scope:** Global Constants

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Param</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** $k - \omega$ RANS turbulence model parameters.

### 3.6.5 Les Turbulence Model Parameter

**Scope:** Global Constants

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Param</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** LES turbulence model parameters.

### 3.6.6 Stefan Boltzmann Constant

**Scope:** Global Constants
Stefan Boltzmann Constant \( \sigma \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \sigma )</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Stefan-Boltzmann constant. Depending on the units involved in the specific problem by the user, this value will differ.

### 3.6.7 Turbulence Model

Scope: Global Constants

<table>
<thead>
<tr>
<th>Param</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \text{Param} )</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>( \text{Value} )</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Turbulence model Schmidt and Prandtl numbers

### 3.7 Function Overview

Sierra Framework services provide an infrastructure that supports definition of user supplied functions. Here the functions can be either tabular or analytical, where the most prevalent use of this capability is tabular data functions. These functions can in turn be tied to different features and models in the applications. Examples of this usage would be to define material data or load profiles, either spatially or in time.

### 3.8 Definition For Function

Scope: Sierra

```
Begin Definition For Function FunctionName

Abscissa \( \text{Name...} \)
Abscissa Offset \( \text{Abscissa_offset} \)
Abscissa Scale \( \text{Abscissa_scale} \)
At Discontinuity Evaluate To \( \text{Option} \)
Column Titles Titles1, Titles2...
Data File = filename [ X From Column xcol Y From Column ycol ]
Debug \( \text{Option} \)
Differentiate Expression \( \text{Expr} \)
Evaluate Expression \( \text{Expr} \)
Evaluate From \( x0 \) To \( x1 \) By \( Dx \)
Expression Variable: \( \text{Expr} = \text{VarType value_var_name...} \)
Expression Variable: \( \text{Expr} \)
```
Ordinate \{:=|\text{are}|\text{is}\} \text{Name...}
Ordinate Offset \{:=|\text{are}|\text{is}\} \text{Ordinate\_offset}
Ordinate Scale \{:=|\text{are}|\text{is}\} \text{Ordinate\_scale}
Scale By $x$
Type \{:=|\text{are}|\text{is}\} \text{Type}
X Offset \{:=|\text{are}|\text{is}\} \text{X\_offset}
X Scale \{:=|\text{are}|\text{is}\} \text{X\_scale}
Y Offset \{:=|\text{are}|\text{is}\} \text{Y\_offset}
Y Scale \{:=|\text{are}|\text{is}\} \text{Y\_scale}
Begin Expressions \text{empty}
End

Begin Values \text{empty}
End

End

Summary
Defines a function in terms of its type and values.

3.8.1 Abscissa
Scope: Definition For Function

Abscissa \{:=|\text{are}|\text{is}\} \text{Name...}

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\text{Name}</td>
<td>\text{string...}</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Specifies a string identifier for the independent variable. Optionally specify a scale and/or offset value which transforms the abscissa values into scaled\_abscissa = scale * (abscissa + abscissa\_offset).

3.8.2 Abscissa Offset
Scope: Definition For Function

Abscissa Offset \{:=|\text{are}|\text{is}\} \text{Abscissa\_offset}

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\text{Abscissa_offset}</td>
<td>\text{real}</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Alias for X OFFSET
3.8.3 Abscissa Scale
Scope: Definition For Function

Abscissa Scale \{=\are\is\} Abscissa\_scale

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abscissa_scale</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Alias for X SCALE

3.8.4 At Discontinuity Evaluate To
Scope: Definition For Function

Summary: Control the behavior of a piecewise constant function when evaluated at a discontinuity (plus or minus a small tolerance). The default behavior is to take the value to the right of the discontinuity. If "Left" is specified, the value to the left of the discontinuity is taken instead.

3.8.5 Column Titles
Scope: Definition For Function

Column Titles Titles\_1 Titles\_2...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Titles</td>
<td>string_1 string_2...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Name the columns (and also defined the expected number of columns) for Multicolumn Piecewise Linear tabular data.

3.8.6 Data File
Scope: Definition For Function

Data File = filename [ X From Column xcol Y From Column ycol ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>filename</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Function will read tabular data from an input file. Compatible with the piecewise linear function type. File must be of form like:

```
# EXAMPLE FILE 1.099 1191 1.101 221 5.9011 133.1
```

Lines headed by a # are considered comments and will be ignored. Data itself must be in tabular columns separated by whitespace or commas.
3.8.7 Debug
Scope: Definition For Function

Summary: Prints functions to the log file.

3.8.8 Differentiate Expression
Scope: Definition For Function

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Expr</td>
<td>(expression)</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Differentiate Expression {=|are|is} Expr

Summary: Specifies the expression of derivative of evaluation expression.

3.8.9 Evaluate Expression
Scope: Definition For Function

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Expr</td>
<td>(expression)</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Evaluate Expression {=|are|is} Expr

Summary: Specifies the expression to evaluate.

Description: This will greatly help with manufactured solutions, and be useful for other purposes as well.

This first implementation goes like this:

```
begin definition for function pressure
  type is analytic
  evaluate expression is "x <= 0.0 ? 0.0 : (x < 0.5 ? x*200.0 : (x < 1.0 ? (x - 0.5) *50.0 + 100.00 : 150.0));"
  # type is piecewise linear
  # begin values
  # 0.0 0.0
  # 0.5 100.0
  # 1.0 150.0
  # end values
end definition for function pressure
```

Also, notice that semicolon at the end. Be sure to put it there for now. You can actually provide multiple expressions to be evaluated, each terminated with a semicolon. This will be handy when multi-dependent variable come into the fold.

The following functions are currently implemented.

**Operators** All C-language operators are supported, e.g. + - * / \ ? : etc

**Parens** ()
Math Functions

\texttt{abs(x)} absolute value of x
\texttt{mod(x, y)} modulus of x/y
\texttt{ipart(x)} integer part of x
\texttt{fpart(x)} fractional part of x
\texttt{min(x0, x1, \ldots)} minimum value of xn
\texttt{max(x0, x1, \ldots)} maximum value of xn

Power functions

\texttt{pow(x, y)} x to the y power
\texttt{sqrt(x)} square root of x

Trig functions

\texttt{sin(x)} sine of x
\texttt{sinh(x)} hyperbolic sine of x
\texttt{asin(x)} arcsine of x
\texttt{cos(x)} cosine of x
\texttt{cosh(x)} hyperbolic cosine of x
\texttt{acos(x)} arccosine of x
\texttt{tan(x)} tangent of x
\texttt{tanh(x)} hyperbolic tangent of x
\texttt{atan(x)} arctangent of x
\texttt{atan2(y, x)} arctangent of y/x, signs of x and y determine quadrant (see atan2 man page)

Logarithm functions

\texttt{log(x)} natural logarithm of x
\texttt{ln(x)} natural logarithm of x
\texttt{exp(x)} e to the x power
\texttt{logn(x, y)} the y base logarithm of x

Rounding functions

\texttt{ceil(x)} smallest integral value not less than x
\texttt{floor(x)} largest integral value not greater than x

Random functions

\texttt{rand(x)} random number between 0.0 and 1.0, not including 1.0
\texttt{srand(x)} seeds the random number generator

Conversion routines

\texttt{deg(x)} converts radians to degrees
\texttt{rad(x)} converts degrees to radians
\texttt{recttopolr(x, y)} magnitude of vector x, y
\texttt{recttopola(x, y)} angle of vector x, y
\texttt{poltorectx(r, theta)} x coordinate of angle theta at distance r
\texttt{poltorecty(r, theta)} y coordinate of angle theta at distance r
3.8.10 Evaluate From
Scope: Definition For Function

Evaluate From \(x_0\) To \(x_1\) By \(Dx\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(x_0)</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>(x_1)</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>(Dx)</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specifies the range and evaluation interval.

3.8.11 Expression Variable:
Scope: Definition For Function

Expression Variable: \(Expr = \text{VarType value}_{\text{var}}\_\text{name...}\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Expr)</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>(value_var_name)</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specifies what the arguments of an expression correspond to. For example:

BEGIN DEFINITION FOR FUNCTION \(dx\_shear\) TYPE = ANALYTIC EXPRESSION variable: \(mx = \text{NODAL model}_\text{coordinates}(x)\) EXPRESSION variable: \(my = \text{NODAL model}_\text{coordinates}(y)\) EXPRESSION variable: \(time = \text{GLOBAL time}\) EVALUATE EXPRESSION = "\((time/termTime)*(stretchx*(mx - 0.0) + ((my-0.25)/0.5)*stretchxy)\)" END

Assuming the above expression is being evaluated on nodes the current values for \(x\) and \(y\) model coordinates would be placed into \(mx\) and \(my\) and current analysis time placed into \(time\)

3.8.12 Expression Variable:
Scope: Definition For Function

Expression Variable: \(Expr\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Expr)</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specifies what the arguments of an expression exists, but does not define it correspond to. For example:

BEGIN DEFINITION FOR FUNCTION \(dx\_shear\) TYPE = ANALYTIC EXPRESSION variable: \(mx\) EXPRESSION variable: \(my\) EXPRESSION variable: \(time\) EVALUATE EXPRESSION = "\((time/termTime)*(stretchx*(mx - 0.0) + ((my-0.25)/0.5)*stretchxy)\)" END

Call function must determine what each variable actually is is based off of the string name
3.8.13 **Ordinate**
Scope: Definition For Function

Ordinate \{=|are|is\} **Name**...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies a string identifier for the dependent variable. Optionally specify a scale and/or offset value which transforms the ordinate values into scaled_ordinate = scale * (ordinate + ordinate_offset).

3.8.14 **Ordinate Offset**
Scope: Definition For Function

Ordinate Offset \{=|are|is\} **Ordinate_offset**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ordinate_offset</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Alias for Y OFFSET

3.8.15 **Ordinate Scale**
Scope: Definition For Function

Ordinate Scale \{=|are|is\} **Ordinate_scale**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ordinate_scale</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Alias for Y SCALE

3.8.16 **Scale By**
Scope: Definition For Function

Scale By \(x\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(x)</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies a scale factor to be applied.

3.8.17 **Type**
Scope: Definition For Function

Summary: Specifies the type of function.
3.8.18 X Offset
Scope: Definition For Function

\[ X \text{ Offset} \{=|\text{are}|\text{is}\} X_{\text{offset}} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(X_{\text{offset}})</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Sets an offset for the x-axis

3.8.19 X Scale
Scope: Definition For Function

\[ X \text{ Scale} \{=|\text{are}|\text{is}\} X_{\text{scale}} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(X_{\text{scale}})</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Sets a scale factor for the x-axis

3.8.20 Y Offset
Scope: Definition For Function

\[ Y \text{ Offset} \{=|\text{are}|\text{is}\} Y_{\text{offset}} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Y_{\text{offset}})</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Sets an offset for the y-axis

3.8.21 Y Scale
Scope: Definition For Function

\[ Y \text{ Scale} \{=|\text{are}|\text{is}\} Y_{\text{scale}} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Y_{\text{scale}})</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Sets a scale factor for the y-axis

3.9 Values
Scope: Definition For Function

Begin Values empty
Summary
Lists the values of the function. The values should be listed one pair per line, independent variable first, with whitespace or comma as a separator.

3.9.1
Scope: Values

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Xyvalues</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
For a piecewise linear function, lists an x-y pair for the nth interpolation point.

3.10 Restarting

Sierra Framework services provide convenient utilities for restarting an analysis from previous results. The most general capability supplements the results of a previous analysis with internal state variables to continue an analysis. In this case the input mesh is supplied from the Input Database Name from the Finite Element Model command block 3.1 and the restart information is obtained from the the Input Database Name from the Restart Data command block 25.6. Continuation of a job using restart data output is invoked using the command line which follows.

Often one wishes to restart an analysis by initializing with results from another analysis. For this case Aria provides another means, IC READ_FILE 8.7, by which to begin the job with variables from existing results.
Chapter 4

Material Properties

4.1 Aria Material Overview

Material property values are prescribed for unique material names using input line commands within an "Aria Material" command block. The association of these material properties with mesh entities is specified within the Finite Element Model command block 3.1.

The general format of the material property line commands as follows

Property Type = Model_Name Value_Name = Value Model_Parameter1 .... Model_ParameterN.

For some material properties, clarification of how the property is to be used must be also be supplied. This clarification usually impacts the computational efficiency of how the property values are being used. As an example, in the case of diffusion coefficients one needs to specify whether it will be used in a scalar or tensor based formulation of flux.

Chemical material models solved using the CHEMEQ library require an entire set of parameters. In this case those parameters are defined within a Parameters for CHEMEQ command block embedded within the Aria Material command block. A description of the CHEMEQ related parameters is contained in a subsequent chapter 36.1.

The material models Encore Function, Global, User Function, Polynomial, User Field and Exponential are termed Generic and are available for any material property evaluation. The Constant material model is also available for all properties but differs syntactically for each material property.

Known Issue: The feature toggling capability is not currently supported for setting material property values, although the line command to do so is listed when the material is specified as a User_Function.

Aria material command blocks appear in the Domain scope of the input file as illustrated below.

Begin Sierra myJob

Begin Aria Material my_mat

material property commands

End Aria Material my_mat

Begin Procedure My_Aria_Procedure


Begin Aria Region MyRegion
.
End Aria Region MyRegion
.
End Procedure My_Aria_Procedure
.
End Sierra myJob

4.1.1 Generic: Encore Function
Scope: Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;Material Property&gt;</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>MaterialPhase</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Mesh Extent Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Data Spec Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>result_name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>eval_type</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Value from an Encore function

4.1.2 Generic: Global
Scope: Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;Material Property&gt;</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>MaterialPhase</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Mesh Extent Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Data Spec Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>global_name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Value from a global variable

4.1.3 Generic: User Function
Scope: Aria Material
**Generic: User Function**

<Material Property> [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@ |at|for|in|on|over} Mesh Extent Name = User_Function [ Using Data Specification Data Spec Name ] [ Name = name |X = x |X_Multiplier = x_multiplier |Multiplier = multiplier |Toggle = toggle ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;Material Property&gt;</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>MaterialPhase</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Mesh Extent Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Data Spec Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>x</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>x_multiplier</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>multiplier</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>toggle</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Value from a user function

---

**4.1.4 Generic: Polynomial**

**Scope:** Aria Material

**Generic: Polynomial**

<Material Property> [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@ |at|for|in|on|over} Mesh Extent Name = Polynomial [ Using Data Specification Data Spec Name ] [ Variable = variable |Order = order |Variable_Offset = variable_offset |C0 = c0 |C1 = c1 |C2 = c2 |C3 = c3 |C4 = c4 |C5 = c5 |C6 = c6 |C7 = c7 |C8 = c8 ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;Material Property&gt;</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>MaterialPhase</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Mesh Extent Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Data Spec Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>variable</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>order</td>
<td>integer</td>
<td>undefined</td>
</tr>
<tr>
<td>variable_offset</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>c0</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>c1</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>c2</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>c3</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>c4</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>c5</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>c6</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>c7</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>c8</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Value from a polynomial function
4.1.5 Generic: User Field
Scope: Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;Material Property&gt;</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>MaterialPhase</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Mesh Extent Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Data Spec Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>scaling</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>global_var</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Value from a user field

4.1.6 Generic: Exponential
Scope: Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;Material Property&gt;</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>MaterialPhase</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Mesh Extent Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Data Spec Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>variable</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>constant</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>multiplier</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>exponent</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Value from an exponential function

4.1.7 Absorption Coefficient
Scope: Aria Material

Absorption Coefficient [ {of|species|subindex} Species ] = Constant [ K = k ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>k</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>
4.1.8 Absorption Cross Section
Scope: Aria Material

Absorption Cross Section \([ \text{Species} ] = \text{Constant} \ [ \text{Abs} = \text{abs} ]\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>abs</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.9 Absorption Cross Section: Linearized
Scope: Aria Material

Absorption Cross Section: Linearized \([ \text{Species} ] = \text{Linearized} \ [ \Sigma_0 = \sigma_0 \mid D_{\Sigma_0} = d_{\sigma_0} \mid D_{\Sigma} = d_{\sigma} \mid T_{\Sigma_0} = t_{\sigma_0} \mid \rho_{\Sigma_0} = \rho_{\sigma_0} ]\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>sigma_0</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>d_sigma_d_rho</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>d_sigma_d_t</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>t_sigma_0</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>rho_sigma_0</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Linearized absorption cross section

4.1.10 Absorption Cross Section: Calore User Sub
Scope: Aria Material

Absorption Cross Section: Calore User Sub \([ \text{Species} ] = \text{Calore_User_Sub} \ [ \text{Name} = \text{name} \mid \text{Type} = \text{type} \mid \text{Multiplier} = \text{multiplier} \mid \text{Material_Data_Block} = \text{material_data_block} \mid \text{Data} = \text{data} \mid \text{Scaling_Field} = \text{scaling_field} ]\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>type</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>multiplier</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>material_data_block</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>data</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>scaling_field</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Calore user subroutine absorption cross section
4.1.11 Advected Bubble
Scope: Aria Material

Advected Bubble [ {of|species|subindex} Species ]= Constant [ Value = value ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.12 Advection Velocity
Scope: Aria Material

Advection Velocity [ {of|species|subindex} Species ]= Constant [ Valuex = valuex | Valuey = valuey | Valuez = valuez ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>valuex</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>valuey</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>valuez</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.13 Air Water Mass Flux: Fickian
Scope: Aria Material

Air Water Mass Flux: Fickian [ {of|species|subindex} Species ]= Fickian [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Fickian diffusion for the air-water mass flux

4.1.14 Altitude
Scope: Aria Material

Altitude [ {of|species|subindex} Species ]= Constant [ A = a ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>a</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>
4.1.15 Ambient Pressure
Scope: Aria Material

Ambient Pressure \([\{\text{of|species|subindex}\} \text{Species}] = \text{Constant} \ [ \text{Value} = \text{value} ]\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.16 Annulus Diameter Ratio
Scope: Aria Material

Annulus Diameter Ratio \([\{\text{of|species|subindex}\} \text{Species}] = \text{Constant} \ [ R = r ]\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>r</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.17 Annulus Diameter Ratio: Spatial User Function
Scope: Aria Material

Annulus Diameter Ratio: Spatial User Function \([\{\text{of|species|subindex}\} \text{Species}] = \text{Spatial User Function} [ \text{Threshold} = \text{threshold} | \text{Multiplier} = \text{multiplier} | \text{Name} = \text{name} | \text{Bar} = \text{bar} | X = x ]\)

<table>
<thead>
<tr>
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<tbody>
<tr>
<td>Species</td>
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<tr>
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</tr>
<tr>
<td>x</td>
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</tbody>
</table>

Summary: Spatially varying annulus diameter ratio based on a coordinate dependent user function

4.1.18 Annulus Diameter Ratio: Correlation
Scope: Aria Material

Annulus Diameter Ratio: Correlation \([\{\text{of|species|subindex}\} \text{Species}] = \text{Correlation} [ R = r | \text{Threshold} = \text{threshold} | \text{Bulk Node Interface} = \text{bulk_node_interface} ]\)
### 4.1.19 Annulus Diameter Ratio: Correlation Spatial User Function

**Scope:** Aria Material

Annulus Diameter Ratio: Correlation Spatial User Function \( \{ \text{Species} \} = \text{Correlation}_\text{Spatial}_{\text{User Function}} \ [ \text{Threshold} = \text{threshold} \ | \text{Multiplier} = \text{multiplier} \ | \text{Name} = \text{name} \ | \text{Bar} = \text{bar} \ | \text{X} = \text{x} \ | \text{Bulk Node Interface} = \text{bulk_node_interface} ] \)

<table>
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<th>Value</th>
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<tr>
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</table>

**Summary**
Correlation annulus diameter ratio based on hydraulic diameter and wetted perimeter

### 4.1.20 Auxiliary Mesh Field: Scalar

**Scope:** Aria Material

Auxiliary Mesh Field: Scalar \( \{ \text{Species} \} = \text{Scalar} [ ] \)

<table>
<thead>
<tr>
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<th>Value</th>
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</tr>
</tbody>
</table>

**Summary**
Scalar auxiliary mesh field

### 4.1.21 Bar Area

**Scope:** Aria Material

Bar Area \( \{ \text{Species} \} = \text{Constant} [ \ A = \ a \ ] \)

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<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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<tbody>
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<tr>
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<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Constant value
4.1.22 Bar Area: Element Attribute
Scope: Aria Material

Bar Area: Element Attribute [ {of|species|subindex} Species ]= Element_Attribute [ Multiplier = multiplier | Attribute_Name = attribute_name ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
<td>multiplier</td>
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<tr>
<td>attribute_name</td>
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<td>undefined</td>
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</table>

Summary Distribution Factor for the bar area

4.1.23 Bar Perimeter
Scope: Aria Material

Bar Perimeter [ {of|species|subindex} Species ]= Constant [ P = p ]

<table>
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<tr>
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</table>

Summary Constant value

4.1.24 Bar Perimeter: Element Attribute
Scope: Aria Material

Bar Perimeter: Element Attribute [ {of|species|subindex} Species ]= Element_Attribute [ Multiplier = multiplier | Attribute_Name = attribute_name ]

<table>
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<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
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Summary Distribution Factor for the bar perimeter

4.1.25 Bc Rad Reference Temperature
Scope: Aria Material

Bc Rad Reference Temperature [ {of|species|subindex} Species ]= Constant [ T_Ref = t_ref ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
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</tr>
</tbody>
</table>

Summary Constant value
4.1.26 Bc Rad Reference Temperature: Bulk Node
Scope: Aria Material

\[
\text{Bc Rad Reference Temperature: Bulk Node } \{ \text{of } \text{species } \text{subindex} \} \text{ Species } = \text{Bulk_Node } \{ \text{Name } = \text{name } \}
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
<tr>
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</tr>
<tr>
<td>name</td>
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</tbody>
</table>

Summary Use a bulk node temperature as the reference temperature for radiation boundary conditions

4.1.27 Bc Rad Reference Temperature: Calore User Sub
Scope: Aria Material

\[
\text{Bc Rad Reference Temperature: Calore User Sub } \{ \text{of } \text{species } \text{subindex} \} \text{ Species } = \text{Calore_User_Sub } \{ \text{Name } = \text{name } | \text{Type } = \text{type } | \text{Multiplier } = \text{multiplier } | \text{Material_Data_Block } = \text{material_data_block } | \text{Data } = \text{data } | \text{Scaling_Field } = \text{scaling_field } \}
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
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</table>

Summary Values from a Calore user subroutine

4.1.28 Bc Reference Temperature
Scope: Aria Material

\[
\text{Bc Reference Temperature } \{ \text{of } \text{species } \text{subindex} \} \text{ Species } = \text{Constant } \{ \text{T_Ref } = \text{t_ref } \}
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
<td>t_ref</td>
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<td>undefined</td>
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</table>

Summary Constant value

4.1.29 Bc Reference Temperature: Bulk Node
Scope: Aria Material

\[
\text{Bc Reference Temperature: Bulk Node } \{ \text{of } \text{species } \text{subindex} \} \text{ Species } = \text{Bulk_Node } \{ \text{Name } = \text{name } \}
\]
**Parameter** 

<table>
<thead>
<tr>
<th>Value</th>
<th>Default</th>
</tr>
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<tbody>
<tr>
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<td>string undefined</td>
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<tr>
<td>name</td>
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</tbody>
</table>

**Summary**

Use a bulk node temperature as the reference temperature for convection boundary conditions

### 4.1.30 Bc Reference Temperature: Calore User Sub

**Scope:** Aria Material

Bc Reference Temperature: Calore User Sub [ {of|species|subindex} Species ] = Calore_User_Sub [ Name = name | Type = type | Multiplier = multiplier | Material_Data_Block = material_data_block | Data = data | Scaling_Field = scaling_field ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
<td>name</td>
<td>&quot;string&quot;</td>
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<tr>
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<td>undefined</td>
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<tr>
<td>material_data_block</td>
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<td>data</td>
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<td>scaling_field</td>
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</table>

**Summary**

Values from a Calore user subroutine

### 4.1.31 Bc Reference Temperature: Bar Segment

**Scope:** Aria Material

Bc Reference Temperature: Bar Segment [ {of|species|subindex} Species ] = Bar_Segment [ Name = name ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>name</td>
<td>&quot;string&quot;</td>
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</tbody>
</table>

**Summary**

Use a bar segment average temperature as the reference temperature for convection boundary conditions

### 4.1.32 Bc Reference Temperature: Computed Adiabatic Wall

**Scope:** Aria Material

Bc Reference Temperature: Computed Adiabatic Wall [ {of|species|subindex} Species ] = Computed_Adiabatic_Wall [ T_On = t_on | T_Off = t_off | Specific_Heat_Function = specific_heat_function | Conductivity_Function = conductivity_function | Viscosity_Function = viscosity_function | Offset_Direction = offset_direction | Coord_Offset = coord_offset ]
### 4.1.33 Beta

**Scope:** Aria Material

Beta \[ \{ \text{of|species|subindex} \} \text{Species} \] = Constant \[ \text{Beta} = \text{beta} \]

<table>
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<tr>
<th>Parameter</th>
<th>Value</th>
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<tr>
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<td>undefined</td>
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</table>

**Summary**
Constant value

### 4.1.34 Beta: Converted

**Scope:** Aria Material

Beta: Converted \[ \{ \text{of|species|subindex} \} \text{Species} \] = Converted \[ \]  

<table>
<thead>
<tr>
<th>Parameter</th>
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</tbody>
</table>

**Summary**
Coefficient of thermal stress computed from Poisson’s ratio, thermal expansion coefficient and elastic modulus

### 4.1.35 Beta: Linear

**Scope:** Aria Material

Beta: Linear \[ \{ \text{of|species|subindex} \} \text{Species} \] = Linear \[ A = a \ | B = b \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>a</td>
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<td>undefined</td>
</tr>
<tr>
<td>b</td>
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<td>undefined</td>
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</table>

**Summary**
Coefficient of thermal stress as a linear function of temperature
4.1.36  Body Acceleration
Scope:  Aria Material

Body Acceleration [ {of|species|subindex} Species ]= Constant [ Valuex = valuex |Valuey = valuey |Valuez = valuez ]

<table>
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<tr>
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<tr>
<td>valuey</td>
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<tr>
<td>valuez</td>
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Summary  Constant value

4.1.37  Bulk Conductivity
Scope:  Aria Material

Bulk Conductivity [ {of|species|subindex} Species ]= Constant [ Kbulk = Kbulk ]

<table>
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<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
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</table>

Summary  Constant value

4.1.38  Bulk Conductivity: T Exponent
Scope:  Aria Material

Bulk Conductivity: T Exponent [ {of|species|subindex} Species ]= T_Exponent [ Kbulk_Ref = kbulk_ref |T_Ref = t_ref |N = n ]

<table>
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<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
<tr>
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<tr>
<td>kbulk_ref</td>
<td>real</td>
<td>undefined</td>
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<tr>
<td>t_ref</td>
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<td>undefined</td>
</tr>
<tr>
<td>n</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Thermal conductivity for bulk material, \( \kappa_{bulk} = \kappa_{ref} \frac{\rho_{solid}}{\rho_{bulk}} \left( \frac{T}{T_{ref}} \right)^n \).

4.1.39  Bulk Conductivity: Volume Average
Scope:  Aria Material

Bulk Conductivity: Volume Average [ {of|species|subindex} Species ]= Volume_Average [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
</tbody>
</table>

69
Summary  Bulk conductivity computed from volume average of species bulk conductivities

4.1.40  Bulk Conductivity: Linear Temperature And Density
Scope:  Aria Material

Bulk Conductivity:  Linear Temperature And Density [ of {species|subindex} Species ] = Linear_Temperature_And_Density
[ C_0 = c_0 | C_Rho = C_rho | C_T = c_t ]

<table>
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<th>Default</th>
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<tr>
<td>c_t</td>
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</table>

Summary  Bulk conductivity that is a linear function of both temperature and density.  \( \kappa_{bulk} = C_0 + C_{rho} \rho + C_T T \)

4.1.41  Bulk Density
Scope:  Aria Material

Bulk Density [ of {species|subindex} Species ] = Constant [ Rho_B = rho_b ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
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Summary  Constant value

4.1.42  Bulk Density: T Exponent
Scope:  Aria Material

Bulk Density:  T Exponent [ of {species|subindex} Species ] = T_Exponent [ Rho_Ref = rho_ref | T_Ref = t_ref | N = n ]

<table>
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<tr>
<th>Parameter</th>
<th>Value</th>
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<tr>
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</tr>
<tr>
<td>n</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Bulk density as a power of normalized temperature \( \rho_{bulk} = \rho_{ref} \left( \frac{T}{T_{ref}} \right)^n \).

4.1.43  Bulk Density: Single Component Ideal Gas
Scope:  Aria Material
4.1.44 Bulk Element Pressure

Scope: Aria Material

Summary: Constant value for the bulk element pressure

4.1.45 Bulk Element Pressure: Integrated

Scope: Aria Material

Summary: Bulk element pressure computed from equation of state

4.1.46 Bulk Element Pressure: Calore User Sub

Scope: Aria Material

Summary: Bulk element pressure from Calore User Subroutine
4.1.47 Bulk Element Volume
Scope: Aria Material

\[
\text{Bulk Element Volume} \ [ \ \{\text{of|species|subindex}\} \ \text{Species} \ ] = \text{Constant} \ [ \ V = v \ | \text{Bulk_Node} = \text{bulk_node}\ ]
\]

<table>
<thead>
<tr>
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<th>Value</th>
<th>Default</th>
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<tr>
<td>\text{v}</td>
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<tr>
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</tr>
</tbody>
</table>

Summary: Constant value for the bulk element volume

4.1.48 Bulk Element Volume: Integrated
Scope: Aria Material

\[
\text{Bulk Element Volume: Integrated} \ [ \ \{\text{of|species|subindex}\} \ \text{Species} \ ] = \text{Integrated} \ [ \ \text{Name} = \text{name} \ ]
\]

<table>
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<tbody>
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Summary: Bulk element volume computed from bounding surfaces

4.1.49 Bulk Element Volume: Calore User Sub
Scope: Aria Material

\[
\text{Bulk Element Volume: Calore User Sub} \ [ \ \{\text{of|species|subindex}\} \ \text{Species} \ ] = \text{Calore_User_Sub} \ [ \ \text{Bulk_Node} = \text{bulk_node} \ ]
\]

<table>
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<th>Value</th>
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Summary: Bulk element volume from Calore User Subroutine

4.1.50 Bulk Mass Density: Porous Density
Scope: Aria Material

\[
\text{Bulk Mass Density: Porous Density} \ [ \ \{\text{of|species|subindex}\} \ \text{Species} \ ] = \text{Porous_Density} \ [ \ ]
\]

<table>
<thead>
<tr>
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<tbody>
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</table>

Summary: Bulk mass density computed from density and porosity.
4.1.51  Bulk Mass Density: Density  
**Scope:** Aria Material  

<table>
<thead>
<tr>
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<th>Value</th>
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</tr>
</thead>
<tbody>
<tr>
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</tbody>
</table>

**Summary**  
Bulk mass density copied from density.

4.1.52  Bulk Mass Density: Mass Fraction Density  
**Scope:** Aria Material  

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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<tbody>
<tr>
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</table>

**Summary**  
Bulk mass density computed from mass fraction and density.

4.1.53  Bulk Mass Density: Mass Fraction Porous Density  
**Scope:** Aria Material  

<table>
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</thead>
<tbody>
<tr>
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</tr>
</tbody>
</table>

**Summary**  
Bulk mass density computed from mass fraction, porosity and density.

4.1.54  Bulk Mass Density: Multiphase Porous Density  
**Scope:** Aria Material  

<table>
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<tr>
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</thead>
<tbody>
<tr>
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</tbody>
</table>

**Summary**  
Bulk mass density computed from density, saturation and porosity for a multiphase material.

4.1.55  Bulk Mass Density  
**Scope:** Aria Material  

---

73
Bulk Mass Density \( \{\text{of|species|subindex}\} \) Species \( \Rightarrow \) Constant \( \{\text{Value = value}\} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
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</tbody>
</table>

Summary: Constant value

4.1.56 Bulk Viscosity
Scope: Aria Material

Bulk Viscosity \( \{\text{of|species|subindex}\} \) Species \( \Rightarrow \) Constant \( \{\text{Kappa = kappa}\} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
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</table>

Summary: Constant value

4.1.57 Bulk Viscosity: Curing Foam
Scope: Aria Material

Bulk Viscosity: Curing Foam \( \{\text{of|species|subindex}\} \) Species \( \Rightarrow \) Curing_Foam \( \{\text{Extent_Subindex = extent_subindex |Extent_Species_Name = extent_species_name |Mu_A = mu_a |T_Mu = T_mu |N_Mu = n_mu |Ksi_C = ksi_c |M_Ksi_C = m_ksi_c |E_Mu = E_mu |R = r}\} \)

<table>
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</tr>
<tr>
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<td>undefined</td>
</tr>
<tr>
<td>ksi_c</td>
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<td>undefined</td>
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<tr>
<td>m_ksi_c</td>
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<tr>
<td>E_mu</td>
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<tr>
<td>r</td>
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</tbody>
</table>

4.1.58 Bulk Viscosity: Phase Average
Scope: Aria Material

Bulk Viscosity: Phase Average \( \{\text{of|species|subindex}\} \) Species \( \Rightarrow \) Phase_Average \( \{\text{Subindex_A = subindex_a |Subindex_B = subindex_b}\} \)

<table>
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<tr>
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<th>Value</th>
<th>Default</th>
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</tr>
<tr>
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<td>undefined</td>
</tr>
<tr>
<td>subindex_b</td>
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</tbody>
</table>

74
**4.1.59 Bulk Viscosity: Interpolated Phase Average**

**Scope:** Aria Material

Bulk Viscosity: Interpolated Phase Average \(\{\text{of|species|subindex}\}\) \(\text{Species}\)= Interpolated\_Phase\_Average \(\{\text{Subindex}_A = \text{subindex}_a \mid \text{Subindex}_B = \text{subindex}_b\}\)

<table>
<thead>
<tr>
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<th>Default</th>
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<tbody>
<tr>
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<tr>
<td>(\text{subindex}_b)</td>
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</tbody>
</table>

**Summary**  
Phase-averaged values

**4.1.60 Capillary Pressure**

**Scope:** Aria Material

Capillary Pressure \(\{\text{of|species|subindex}\}\) \(\text{Species}\)= Constant \(\{\text{Value} = \text{value}\}\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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<tr>
<td>(\text{value})</td>
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</tr>
</tbody>
</table>

**Summary**  
Constant value

**4.1.61 Capillary Pressure: Van Genuchten**

**Scope:** Aria Material

Capillary Pressure: Van Genuchten \(\{\text{of|species|subindex}\}\) \(\text{Species}\)= Van\_Genuchten \(\{\text{Residual\_Wetting\_Phase\_Saturation} = \text{residual\_wetting\_phase\_saturation} \mid \text{Residual\_Nonwetting\_Phase\_Saturation} = \text{residual\_nonwetting\_phase\_saturation} \mid \text{Reference\_Capillary\_Pressure} = \text{reference\_capillary\_pressure} \mid \text{Beta} = \text{beta} \mid \text{Wetting\_Phase} = \text{wetting\_phase}\}\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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<tr>
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<tr>
<td>(\text{residual_nonwetting_phase_saturation})</td>
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<td>undefined</td>
</tr>
<tr>
<td>(\text{reference_capillary_pressure})</td>
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<tr>
<td>(\text{beta})</td>
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</table>

**Summary**  
A Van Genuchten model for air/water capillary pressure

**4.1.62 Capillary Pressure: Encore Saturation**

**Scope:** Aria Material
Capillary Pressure: Encore Saturation

\[
\text{Species} = \text{Encore Saturation} \quad \text{[Name} = \text{name} \quad \text{[Result}_\text{Name} = \text{result}_\text{name} \quad \text{[Eval}_\text{Type} = \text{eval}_\text{type}]
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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<tr>
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</tr>
<tr>
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</table>

Summary: A Van Genuchten model for air/water capillary pressure based on an encore function for saturation.

### 4.1.63 Capillary Pressure: Udell Cubic

Scope: Aria Material

\[
\text{Species} = \text{Udell Cubic} \quad \text{[Residual}_\text{Wetting}_\text{Phase}_\text{Saturation} = \text{residual}_\text{wetting}_\text{phase}_\text{saturation} \quad \text{[Residual}_\text{Nonwetting}_\text{Phase}_\text{Saturation} = \text{residual}_\text{nonwetting}_\text{phase}_\text{saturation} \quad \text{[Sigma} = \text{sigma} \quad \text{[C1} = \text{c1} \quad \text{[C2} = \text{c2} \quad \text{[C3} = \text{c3} \quad \text{[Wetting}_\text{Phase} = \text{wetting}_\text{phase}]
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
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<tr>
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</tr>
<tr>
<td>c2</td>
<td>real</td>
<td>undefined</td>
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<tr>
<td>c3</td>
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</tbody>
</table>

Summary: A Udell cubic model for air/water capillary pressure.

### 4.1.64 Capillary Pressure: Computed

Scope: Aria Material

\[
\text{Species} = \text{Computed} \quad \text{[Wetting}_\text{Phase} = \text{wetting}_\text{phase} \quad \text{[Nonwetting}_\text{Phase} = \text{nonwetting}_\text{phase}]
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>nonwetting_phase</td>
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</tbody>
</table>

Summary: Capillary pressure computed based on the pressure difference between the non-wetting and wetting phases, each of which may be specified as parameters.

### 4.1.65 Characteristic Length: Correlation

Scope: Aria Material

76
Characteristic Length: Correlation [ {of|species|subindex} Species ]= Correlation [ Dh = Dh |Bulk_Node_Interface = bulk_node_interface ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</table>

Summary Correlation characteristic length with constant value

**4.1.66 Characteristic Length: Correlation Bar Perimeter**

Scope: Aria Material

Characteristic Length: Correlation Bar Perimeter [ {of|species|subindex} Species ]= Correlation_Bar_Perimeter [ Bulk_Node_Interface = bulk_node Interface ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
<tr>
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Summary Correlation characteristic length with bar perimeter model

**4.1.67 Characteristic Length: Correlation Hydraulic Diameter**

Scope: Aria Material

Characteristic Length: Correlation Hydraulic Diameter [ {of|species|subindex} Species ]= Correlation_Hydraulic_Diameter [ Bulk_Node_Interface = bulk_node Interface ]

<table>
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<tr>
<th>Parameter</th>
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Summary Correlation characteristic length with hydraulic diameter model

**4.1.68 Characteristic Length: Correlation Wetted Perimeter**

Scope: Aria Material

Characteristic Length: Correlation Wetted Perimeter [ {of|species|subindex} Species ]= Correlation_Wetted_Perimeter [ Bulk_Node_Interface = bulk_node Interface ]

<table>
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<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>bulk_node_interface</td>
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</tbody>
</table>

Summary Correlation characteristic length with wetted perimeter model
### 4.1.69 Charge Density

**Scope:** Aria Material

Charge Density \( \{ \text{of|species|subindex} \} \) \( \text{Species} \) = Constant \( \{ \text{Value} = \text{value} \} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</tbody>
</table>

**Summary**

Constant value

### 4.1.70 Chemical Potential: Ideal Solution

**Scope:** Aria Material

Chemical Potential: Ideal Solution \( \{ \text{of|species|subindex} \} \) \( \text{Species} \) = Ideal_Solution \( \{ \text{R} = r \} \) \( \text{Temperature_Material_Phase} = \text{temperature_material_phase} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
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<td>( r )</td>
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<td>undefined</td>
</tr>
<tr>
<td>( \text{temperature_material_phase} )</td>
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</tr>
</tbody>
</table>

**Summary**

Ideal solution chemical potential, \( \mu = RT \log(C) \).

### 4.1.71 Chemical Potential: From Equilibrium Potential

**Scope:** Aria Material

Chemical Potential: From Equilibrium Potential \( \{ \text{of|species|subindex} \} \) \( \text{Species} \) = From_Equilibrium \( \{ \text{F} = f \} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
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</tr>
<tr>
<td>( f )</td>
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<td>undefined</td>
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</tbody>
</table>

**Summary**

Chemical potential from the equilibrium voltage of an electrochemical reaction. This model assumes that it is a single electron intercalation reaction and that only differences in the chemical potential are relevant, not the absolute magnitude. Therefore \( \mu = -F\phi_{eq} \).

### 4.1.72 Chemical Potential: Isotropic Mesh Stress

**Scope:** Aria Material

Chemical Potential: Isotropic Mesh Stress \( \{ \text{of|species|subindex} \} \) \( \text{Species} \) = Isotropic_Mesh_Stress \( \{ \text{Strain_Coeff} = \text{strain_coeff} \} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
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</tr>
<tr>
<td>( \text{strain_coeff} )</td>
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<td>undefined</td>
</tr>
</tbody>
</table>
Summary: Contribution to the chemical potential from species strain. Assumes that the strain is isotropic so \( \mu = \Omega \sigma_{kk}/3 \) where \( \Omega \) is the volumetric expansion coefficient and \( \sigma \) is the stress tensor. This model uses the mesh stress provided by a projection equation in order to recover stress gradients.

4.1.73 Contact Electrical Conductance Coefficient
Scope: Aria Material

Contact Electrical Conductance Coefficient [ {of species subindex} Species ] = Constant [ \( H = h \) ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>( h )</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.74 Contact Electrical Conductance Coefficient: Calore User Sub
Scope: Aria Material

Contact Electrical Conductance Coefficient: Calore User Sub [ {of species subindex} Species ] = Calore_User_Sub [ Name = name | Type = type | Multiplier = multiplier | Material_Data_Block = material_data_block | Data = data | Scaling_Field = scaling_field ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
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</tr>
<tr>
<td>material_data_block</td>
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<td>undefined</td>
</tr>
<tr>
<td>data</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>scaling_field</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Values from a Calore user subroutine

4.1.75 Contact Heat Transfer Coefficient
Scope: Aria Material

Contact Heat Transfer Coefficient [ {of species subindex} Species ] = Constant [ \( H = h \) ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
<td>( h )</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value
4.1.76 Contact Heat Transfer Coefficient: Calore User Sub
Scope: Aria Material

Contact Heat Transfer Coefficient: Calore User Sub [ {of|species|subindex} Species ]= Calore_User_Sub [ Name = name |Type = type |Multiplier = multiplier |Material_Data_Block = material_data_block |Data = data |Scaling_Field = scaling_field ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>name</td>
<td>&quot;string&quot;</td>
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<tr>
<td>type</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
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<tr>
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<td>data</td>
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</tr>
<tr>
<td>scaling_field</td>
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</table>

Summary Values from a Calore user subroutine

4.1.77 Continuity Diffusion Coefficient
Scope: Aria Material

Continuity Diffusion Coefficient [ {of|species|subindex} Species ]= Constant [ Value = value ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>value</td>
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</tr>
</tbody>
</table>

Summary Constant value

4.1.78 Continuity Diffusion Coefficient: Shakib
Scope: Aria Material

Continuity Diffusion Coefficient: Shakib [ {of|species|subindex} Species ]= Shakib [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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</tr>
</tbody>
</table>

Summary Pressure POISSON with diffusion coefficient given by Shakib tau.

4.1.79 Continuity Diffusive Flux: Basic
Scope: Aria Material

Continuity Diffusive Flux: Basic [ {of|species|subindex} Species ]= Basic [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>
4.1.80 Continuity Diffusive Flux: Pressure Poisson

**Scope:** Aria Material

Continuity Diffusive Flux: Pressure Poisson [ {of|species|subindex} Species ]= Pressure_Poisson [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Pressure poisson model of continuity diffusive flux

4.1.81 Cte

**Scope:** Aria Material

Cte [ {of|species|subindex} Species ]= Constant [ Cte = cte ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>cte</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Constant value

4.1.82 Current Density

**Scope:** Aria Material

Current Density [ {of|species|subindex} Species ]= Constant [ Valuex = valuex |Valuey = valuey |Valuez = valuez ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>valuex</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>valuey</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>valuez</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Constant value

4.1.83 Current Density: Ohms Law

**Scope:** Aria Material

Current Density: Ohms Law [ {of|species|subindex} Species ]= Ohms_Law [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tbody>
</table>

Summary Ohms law current density
4.1.84 Current Density: Basic
Scope: Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
</tbody>
</table>

Summary: Basic current density

4.1.85 Current Density: Electrolyte
Scope: Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
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<td>undefined</td>
</tr>
<tr>
<td>f</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Electrolyte current density

4.1.86 Current Density: Dielectric Displacement
Scope: Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
</tbody>
</table>

Summary: Dielectric displacement current density

4.1.87 Current Density: Thermoelectric
Scope: Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Thermoelectric current density

4.1.88 Current Exchange Density
Scope: Aria Material
Current Exchange Density \[ \{ \text{of|species|subindex} \} \text{Species} \] = Constant \[ I_0 = i_0 \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>(i_0)</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Constant value

### 4.1.89 Cantera Xml File

**Scope:** Aria Material

**Cantera Xml File \{=|are|is\} XML Path**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>XML Path</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Description** This specifies the name of the XML file used by Cantera to import the properties of the chemical species mixture. The order of the species in the simulation will be set by the order in this file.

### 4.1.90 Damage Capacitance

**Scope:** Aria Material

**Damage Capacitance \{=|are|is\} Species \] = Constant \[ \text{Value} = \text{value} \] **

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>(\text{value})</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Constant value

### 4.1.91 Damage Diffusive Flux: Basic

**Scope:** Aria Material

**Damage Diffusive Flux: Basic \{=|are|is\} Species \] = Basic \[ \] **

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Basic model of damage diffusive flux

### 4.1.92 Damage Diffusivity

**Scope:** Aria Material

**Damage Diffusivity \{=|are|is\} Species \] = Constant \[ \text{Value} = \text{value} \] **
4.1.93  Density
Scope:  Aria Material

Density [ {of|species|subindex} Species ] = Constant [ Rho = rho ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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<td>undefined</td>
</tr>
<tr>
<td>rho</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Constant value

4.1.94  Density: Calore User Sub
Scope:  Aria Material

Density:  Calore User Sub [ {of|species|subindex} Species ] = Calore_User_Sub [ Name = name |Type = type |Multiplier = multiplier |Material_Data_Block = material_data_block |Data = data |Scaling_Field = scaling_field ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>type</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>multiplier</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>material_data_block</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>data</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>scaling_field</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Values from a Calore user subroutine

4.1.95  Density: Cantera
Scope:  Aria Material

Density:  Cantera [ {of|species|subindex} Species ] = Cantera [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Cantera density summary
4.1.96 Density: Cantera Molten Salt
Scope: Aria Material

Density: Cantera Molten Salt [ {of|species|subindex} Species ] = Cantera_Molten_Salt [ DensityConversion = densityConversion | CanteraXMLFile = canteraXMLFile ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>densityConversion</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>canteraXMLFile</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Cantera density summary

4.1.97 Density: Clsm
Scope: Aria Material

Density: Clsm [ {of|species|subindex} Species ] = Clsm [ PrimaryProperty = primaryProperty ]
| SecondaryProperty       | real     | undefined   |

Summary CLSM density

4.1.98 Density: Compressible Boussinesq
Scope: Aria Material

Density: Compressible Boussinesq [ {of|species|subindex} Species ] = Compressible_Boussinesq [ Ref_Density = ref_density | Alpha = alpha | Ref_Pressure = ref_pressure | Ref_Temperature = ref_temperature ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>ref_density</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>alpha</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>ref_pressure</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>ref_temperature</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary A compressible Boussinesq model for the density

4.1.99 Density: Curing Foam
Scope: Aria Material

Density: Curing Foam [ {of|species|subindex} Species ] = Curing_Foam [ Vfrac_Subindex = vfrac_subindex | Rho_E = rho_e | Rho_F = rho_f | R = r | Phi_Zero = phi_zero ]
4.1.100  Density: Exp Decay
Scope:  Aria Material

Density:  Exp Decay [ {of|species|subindex} Species ] = Exp_Decay [ Rho_Initial = rho_initial |Rho_Final = rho_final |K = k ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>rho_initial</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>rho_final</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>k</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Density as an exponential decay function in time

4.1.101  Density: Foam Time Temp
Scope:  Aria Material


<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>rho_initial</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>rho_final</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>c_param</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>d_param</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>t_max</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>time_naught</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Foam time temp density

4.1.102  Density: From Volume Fraction Gas
Scope:  Aria Material

Density:  From Volume Fraction Gas [ {of|species|subindex} Species ] = From_Volume_Fraction_Gas [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>
4.1.103 Density: General Ideal Gas
Scope: Aria Material

Density: General Ideal Gas [ {of|species|subindex} Species ]= General_Ideal_Gas [ R = r |Pref = pref ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>r</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>pref</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary General ideal gas density

4.1.104 Density: General Ideal Gas Extract Average Pressure
Scope: Aria Material

Density: General Ideal Gas Extract Average Pressure [ {of|species|subindex} Species ]= General_Ideal_Gas_Extract_Average_Pressure [ R = r |Pref = pref ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>r</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>pref</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary General ideal gas density that subtracts off the average pressure value as a constant every time step so that the pressure dof stays O(1) throughout the simulation.

4.1.105 Density: Ideal Gas
Scope: Aria Material

Density: Ideal Gas [ {of|species|subindex} Species ]= Ideal_Gas [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tbody>
</table>

Summary Ideal gas density

4.1.106 Density: Incompressible Ideal Gas
Scope: Aria Material

Density: Incompressible Ideal Gas [ {of|species|subindex} Species ]= Incompressible_Ideal_Gas [ P_Ref = p_ref |T_Ref = t_ref |R = r ]
Summary  
Incompressible ideal gas density

4.1.107 Density: Mixture Fraction
Scope: Aria Material

\[
\text{Density: Mixture Fraction} \ [\{\text{of|species|subindex}\} \text{Species}] = \text{Mixture_Fraction} [\text{Primaryproperty} = \text{primaryProperty} |\text{Secondaryproperty} = \text{secondaryProperty} ]
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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<tr>
<td>primaryProperty</td>
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<td>undefined</td>
</tr>
<tr>
<td>secondaryProperty</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.108 Density: Phase Average
Scope: Aria Material

\[
\text{Density: Phase Average} \ [\{\text{of|species|subindex}\} \text{Species}] = \text{Phase_Average} [\text{Subindex}_A = \text{subindex}_a |\text{Subindex}_B = \text{subindex}_b ]
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>subindex_a</td>
<td>integer</td>
<td>undefined</td>
</tr>
<tr>
<td>subindex_b</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  
Phase-averaged values

4.1.109 Density: Interpolated Phase Average
Scope: Aria Material

\[
\text{Density: Interpolated Phase Average} \ [\{\text{of|species|subindex}\} \text{Species}] = \text{Interpolated_Phase_Average} [\text{Subindex}_A = \text{subindex}_a |\text{Subindex}_B = \text{subindex}_b ]
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>subindex_a</td>
<td>integer</td>
<td>undefined</td>
</tr>
<tr>
<td>subindex_b</td>
<td>integer</td>
<td>undefined</td>
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</tbody>
</table>

Summary  
Interpolated Phase-averaged values

4.1.110 Density: Single Component Ideal Gas
Scope: Aria Material
4.1.111 Density: Single Component Adiabatic Ideal Gas

Scope: Aria Material

Density: Single Component Adiabatic Ideal Gas \[ \{ \text{of|species|subindex} \} \text{Species} \] = Single_Component_Adiabatic_Ideal_Gas
\[ P_{\text{Ref}} = p_{\text{ref}} \ | P_0 = p_0 \ | \text{Density}_0 = \text{density}_0 \ | \Gamma = \gamma \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>p_ref</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>p_0</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>density_0</td>
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<td>undefined</td>
</tr>
<tr>
<td>gamma</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.112 Density: Stanford

Scope: Aria Material

Density: Stanford \[ \{ \text{of|species|subindex} \} \text{Species} \] = Stanford \[ N = n \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>n</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.113 Density: Thermal

Scope: Aria Material

Density: Thermal \[ \{ \text{of|species|subindex} \} \text{Species} \] = Thermal \[ A = a \ | B = b \ | C = c \ | D = d \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>a</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>b</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>c</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>d</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Density as a cubic polynomial in temperature
4.1.114  Density: Mass Average
Scope: Aria Material

Density: Mass Average [{of|species|subindex} Species ]= Mass_Average [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Density as a mass average of the species bulk densities.

4.1.115  Density: Mass Preserving
Scope: Aria Material


<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>rho</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Density applied as a ratio of original to current element volume.

4.1.116  Density: Concentration Average
Scope: Aria Material

Density: Concentration Average [{of|species|subindex} Species ]= Concentration_Average [ Fluid_Density = fluid_density |Lambda = lambda ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
<td>fluid_density</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>lambda</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Density as a concentration (phi) average.

4.1.117  Density: From Mass Fraction
Scope: Aria Material

Density: From Mass Fraction [{of|species|subindex} Species ]= From_Mass_Fraction [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Species density computed from mass fraction.
4.1.118 Density: Adaptive Table Lookup
Scope: Aria Material

Density: Adaptive Table Lookup [ {of|species|subindex} Species ] = Adaptive_Table_Lookup
[ File_Name = file_name ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>file_name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: An adaptive table lookup of density

4.1.119 Density: Porous Phase Average
Scope: Aria Material

Density: Porous Phase Average [ {of|species|subindex} Species ] = Porous_Phase_Average
[ Liquid_Phase_Name = Liquid_Phase_Name | Gas_Phase_Name = Gas_Phase_Name ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Liquid_Phase_Name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>Gas_Phase_Name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Density for a multiphase material of two or three phases

4.1.120 Density: From Chemeq
Scope: Aria Material

Density: From Chemeq [ {of|species|subindex} Species ] = From_Chemeq [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Density of one species extracted from ChemEq

4.1.121 Density: From Chemeq Solids
Scope: Aria Material

Density: From Chemeq Solids [ {of|species|subindex} Species ] = From_Chemeq_Solids [ Rhob0 = rhob0 ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>rhob0</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Condensed phase density from ChemEq species concentrations for solid species
4.1.122  Density: Sum All Species
Scope:  Aria Material

\[
\text{Density: Sum All Species [ \{of \text{species} \text{subindex}\} Species ] = Sum\_All\_Species [ ]}
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.123  Density: Correlation Fluid
Scope:  Aria Material

\[
\text{Density: Correlation Fluid [ \{of \text{species} \text{subindex}\} Species ] = Correlation\_Fluid [ Rho = rho | Bulk\_Node\_Interface = bulk\_node\_interface ]}
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>rho</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>bulk_node_interface</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Correlation fluid temperature

4.1.124  Dispersed Phase Density
Scope:  Aria Material

\[
\text{Dispersed Phase Density [ \{of \text{species} \text{subindex}\} Species ] = Constant [ Value = value ]}
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Constant value

4.1.125  Dispersed Phase Momentum Stress: Newtonian
Scope:  Aria Material

\[
\text{Dispersed Phase Momentum Stress: Newtonian [ \{of \text{species} \text{subindex}\} Species ] = Newtonian [ Cts = cts ]}
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>cts</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.126  Dispersed Phase Velocity
Scope:  Aria Material

\[
\text{Dispersed Phase Velocity [ \{of \text{species} \text{subindex}\} Species ] = Constant [ Velocity\_X = Velocity\_X | Velocity\_Y = Velocity\_Y | Velocity\_Z = Velocity\_Z ]}
\]
Parameter | Value | Default
---|---|---
Species | string | undefined
Velocity X | real | undefined
Velocity Y | real | undefined
Velocity Z | real | undefined

Summary | Constant value

### 4.1.127 Dispersed Phase Volume Fraction

**Scope:** Aria Material

Dispersed Phase Volume Fraction [ {of|species|subindex} Species ] = Constant [ Value = value ]

Parameter | Value | Default
---|---|---
Species | string | undefined
value | real | undefined

Summary | Constant value

### 4.1.128 Distribution Coefficient

**Scope:** Aria Material

Distribution Coefficient [ {of|species|subindex} Species ] = Constant [ Kd = kd ]

Parameter | Value | Default
---|---|---
Species | string | undefined
kd | real | undefined

Summary | Constant value

### 4.1.129 Distribution Coefficient: Langmuir

**Scope:** Aria Material

Distribution Coefficient: Langmuir [ {of|species|subindex} Species ] = Langmuir [ K = k | K I = k I ]

Parameter | Value | Default
---|---|---
Species | string | undefined
k | real | undefined
k I | real | undefined

Summary | Langmuir isotherm distribution coefficient for species equation

### 4.1.130 Distribution Coefficient: Competitive Langmuir

**Scope:** Aria Material
Distribution Coefficient: Competitive Langmuir \[ \text{of|species|subindex} \text{Species} = \text{Competitive}_\text{Langmuir} \]
\[ K_I = k_i \mid K_J = k_j \mid K_{ie} = k_{ie} \mid \text{Species}_J = \text{species}_j \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>$k_i$</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>$k_j$</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>$k_{ie}$</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>$\text{species}_j$</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Competitive Langmuir isotherm distribution coefficient for species equation

4.1.131 Effective Diffusivity: Hdiff
Scope: Aria Material

Effective Diffusivity: Hdiff \[ \text{of|species|subindex} \text{Species} = \text{Hdiff} \]
\[ K_T = k_t \mid N_L = n_l \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>$k_t$</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>$n_l$</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: ?

4.1.132 Electric Displacement: Linear
Scope: Aria Material

Electric Displacement: Linear \[ \text{of|species|subindex} \text{Species} = \text{Linear} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
</tbody>
</table>

Summary: Linear electric displacement

4.1.133 Electric Displacement: Basic
Scope: Aria Material

Electric Displacement: Basic \[ \text{of|species|subindex} \text{Species} = \text{Basic} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Basic electric displacement
### 4.1.134 Electric Displacement: Generalized

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tbody>
</table>

**Summary**

Generalized electric displacement

### 4.1.135 Electrical Conductivity

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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<td>sigma</td>
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</tr>
</tbody>
</table>

**Summary**

Constant value

### 4.1.136 Electrical Conductivity: From Resistivity

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
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</tr>
</tbody>
</table>

**Summary**

Electrical conductivity calculated from the electrical resistance

### 4.1.137 Electrical Conductivity: Tbc

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
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</tr>
<tr>
<td>ki</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>ti</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>e</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>r</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

TBC model for electrical conductivity
4.1.138 Electrical Conductivity: Thermal
Scope: Aria Material

Electrical Conductivity: Thermal [ of|species|subindex] Species ] = Thermal [ A = a | B = b | C = c | D = d ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>a</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>b</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>c</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>d</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Electrical conductivity as a cubic polynomial in temperature

4.1.139 Electrical Conductivity: Bruggeman Volume Averaged
Scope: Aria Material


<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>Exponent</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary This model averages the properties of each species in the same material phase based on their volume fraction assuming a Bruggeman tortuosity model.

4.1.140 Electrical Conductivity: Arrhenius
Scope: Aria Material

Electrical Conductivity: Arrhenius [ of|species|subindex] Species ] = Arrhenius [ R = r | Sigma0 = sigma0 | A = a | Ea = Ea ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>r</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>sigma0</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>a</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>Ea</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Arrhenius electrical conductivity, \( \sigma = \sigma_0 + A \exp(-E_a/(R*T)) \)

4.1.141 Electrical Permittivity
Scope: Aria Material

Electrical Permittivity [ of|species|subindex] Species ] = Constant [ Kappa = kappa ]
Summary: Constant value

4.1.142 Electrical Resistance
Scope: Aria Material

Electrical Resistance [ {of|species|subindex} Species ] = Constant [ Value = value ]

Parameter Value Default
Species string undefined
kappa real undefined

Summary: Constant value

4.1.143 Electrical Resistivity
Scope: Aria Material

Electrical Resistivity [ {of|species|subindex} Species ] = Constant [ Rho = rho ]

Parameter Value Default
Species string undefined
rho real undefined

Summary: Constant value

4.1.144 Electrical Resistivity: From Conductivity
Scope: Aria Material

Electrical Resistivity: From Conductivity [ {of|species|subindex} Species ] = From_Conductivity [ ]

Parameter Value Default
Species string undefined

Summary: Electrical resistivity calculated from the electrical conductivity

4.1.145 Electromigration Coefficient
Scope: Aria Material

Electromigration Coefficient [ {of|species|subindex} Species ] = Constant [ M = m ]

Parameter Value Default
Species string undefined
m real undefined
4.1.146  Emissivity
Scope:  Aria Material

Emissivity [ {of|species|subindex} Species ]= Constant [ \( E = e \) ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>( e )</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.147  Emissivity: Calore User Sub
Scope:  Aria Material

Emissivity:  Calore User Sub [ {of|species|subindex} Species ]= Calore_User_Sub [ Name = name | Type = type | Multiplier = multiplier | Material_Data_Block = material_data_block | Data = data | Scaling_Field = scaling_field ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>type</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>multiplier</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>material_data_block</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>data</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>scaling_field</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Values from a Calore user subroutine

4.1.148  Enclosure Mk
Scope:  Aria Material

Enclosure Mk [ {of|species|subindex} Species ]= Constant [ \( K = k \) ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>( k )</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Constant value

4.1.149  Enclosure Mk: Calore User Sub
Scope:  Aria Material
Enclosure Mbk: Calore User Sub [ {of|species|subindex} Species ] = Calore_User_Sub [ Name = name | Type = type | Multiplier = multiplier | Material_Data_Block = material_data_block | Data = data | Scaling_Field = scaling_field ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>type</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>multiplier</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>material_data_block</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>data</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>scaling_field</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Values from a Calore user subroutine

4.1.150 Enclosure Mbl
Scope: Aria Material

Enclosure Mbl [ {of|species|subindex} Species ] = Constant [ L = l ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>l</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.151 Enclosure Mbl: Calore User Sub
Scope: Aria Material

Enclosure Mbl: Calore User Sub [ {of|species|subindex} Species ] = Calore_User_Sub [ Name = name | Type = type | Multiplier = multiplier | Material_Data_Block = material_data_block | Data = data | Scaling_Field = scaling_field ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>type</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>multiplier</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>material_data_block</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>data</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>scaling_field</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Values from a Calore user subroutine

4.1.152 Enclosure Mbl: From Geometry
Scope: Aria Material
Enclosure Mbl: From Geometry \{of|species|subindex\} Species = From_Geometry \{ Prefactor = \textit{prefactor} \ | Enclosure = \textit{enclosure} \} 

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>prefactor</td>
<td>real</td>
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</tr>
<tr>
<td>enclosure</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Values from geometry with a user selectable pre-factor

4.1.153 Energy Diffusive Flux: Basic
Scope: Aria Material

Energy Diffusive Flux: Basic \{of|species|subindex\} Species = Basic \[

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Basic model of energy diffusive flux

4.1.154 Energy Diffusive Flux: From Mixture Mass Diffusivity
Scope: Aria Material

Energy Diffusive Flux: From Mixture Mass Diffusivity \{of|species|subindex\} Species = From_Mixture_Mass_Diffusivity \[

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Energy diffusive flux = -density * mixture_mass_diffusivity * grad(enthalpy)

4.1.155 Energy Diffusive Flux: Energy Mass
Scope: Aria Material

Energy Diffusive Flux: Energy Mass \{of|species|subindex\} Species = Energy_Mass \[

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Energy mass model of energy diffusive flux

4.1.156 Energy Face Stabilization Scaling: Default
Scope: Aria Material

Summary: Default face stabilization scaling for energy equation
4.1.157 Enthalpy
Scope: Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
<tr>
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<tr>
<td>h</td>
<td>real</td>
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</table>

Summary: Constant value

4.1.158 Enthalpy: Cantera
Scope: Aria Material

<table>
<thead>
<tr>
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<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td></td>
</tr>
</tbody>
</table>

Summary: Cantera enthalpy computed from temperature

4.1.159 Enthalpy: From Temperature
Scope: Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
<td>t</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Cantera enthalpy computed from temperature

4.1.160 Enthalpy: Mass Average
Scope: Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td></td>
</tr>
</tbody>
</table>

Summary: Enthalpy computed from mass average of species enthalpies

4.1.161 Enthalpy Advection: Porous
Scope: Aria Material
### 4.1.162 Enthalpy Advection: Porous Upwind

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Upwinded advected quantity for porous enthalpy equation. Only valid for CVFEM.

### 4.1.163 Entrance Distance: Correlation Bar

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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</tr>
<tr>
<td>phase</td>
<td>&quot;string&quot;</td>
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</tr>
<tr>
<td>gas_exponent</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>bulk_node_interface</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Correction factor for difference between near-wall and bulk fluid state for Gnielinski pipe and annulus correlations.

### 4.1.164 Equilibrium Constant: Arrhenius

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>c</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>m</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>w_b</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>r</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

102
4.1.165 Equilibrium Potential
Scope: Aria Material

Equilibrium Potential \[ \{ \text{of|species|subindex} \} \text{Species } = \text{Constant } [ \text{U} = \text{u} ] \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\text{Species}</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{u}</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Constant value

4.1.166 Equilibrium Potential: Nernst
Scope: Aria Material

Equilibrium Potential: Nernst \[ \{ \text{of|species|subindex} \} \text{Species } = \text{Nernst } [ \text{Species}_A = \text{species}_a | \text{Species}_B = \text{species}_b | \text{R} = \text{r} | \text{F} = \text{f} | \text{U} = \text{u} ] \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\text{Species}</td>
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<td>undefined</td>
</tr>
<tr>
<td>\text{species}_a</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{species}_b</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{r}</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{f}</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{u}</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Nernst-Einstein form of the equilibrium potential

4.1.167 Equilibrium Potential: From Chemical Potential
Scope: Aria Material

Equilibrium Potential: From Chemical Potential \[ \{ \text{of|species|subindex} \} \text{Species } = \text{From_Chemical_Potential } [ \text{F} = \text{f} | \text{Species}_Name = \text{species}_name | \text{Species}_Phase = \text{species}_phase ] \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\text{Species}</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{f}</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{species}_name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{species}_phase</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Determine the equilibrium voltage based on the chemical potential of a selected species where \( \varphi_{\text{equil}} = -\mu/F \).

4.1.168 Equilibrium Potential: Two Phase
Scope: Aria Material

Equilibrium Potential: Two Phase \[ \{ \text{of|species|subindex} \} \text{Species } = \text{Two_Phase } [ \text{Cmin} = \text{Cmin} | \text{Cmax} = \text{Cmax} | \text{R} = \text{r} | \text{Species}_Name = \text{species}_name ] \]
### 4.1.169 Equilibrium Pressure

**Scope:** Aria Material

 Equilibrium Pressure \([ \{ \text{of|species|subindex} \} \text{Species } ] = \text{Constant } [ \text{P } = \text{p} ]\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>(p)</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Constant value

### 4.1.170 Exchange Current Density

**Scope:** Aria Material

 Exchange Current Density \([ \{ \text{of|species|subindex} \} \text{Species } ] = \text{Constant } [ \text{Value } = \text{value} ]\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>(\text{value})</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Constant value

### 4.1.171 Exchange Current Density: Rate Constant Species

**Scope:** Aria Material

 Exchange Current Density: Rate Constant Species \([ \{ \text{of|species|subindex} \} \text{Species } ] = \text{RateConstantSpecies } [ \text{F } = \text{f} | \text{K } = \text{k} | \text{Liquid_Exponent } = \text{liquid_exponent} | \text{Solid_Exponent } = \text{solid_exponent} | \text{Liquid_Species } = \text{liquid_species} | \text{Solid_Species } = \text{solid_species} ]\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\text{Species})</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>(f)</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>(k)</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>(\text{liquid_exponent})</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>(\text{solid_exponent})</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>(\text{liquid_species})</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>(\text{solid_species})</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Computes an exchange current density of the form
\[ i_0 = kF \prod_{\text{liquid\_species}}^{\text{liquid\_exponent}} \times \prod_{\text{solid\_species}}^{\text{solid\_exponent}} \] where the species whose concentrations are included are user-specified parameters.

### 4.1.172 Extension Speed

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>value</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

### 4.1.173 Extension Speed Diffusive Flux: Definition

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: The definition of the extension speed diffusive flux

### 4.1.174 Extension Speed Multiplier

**Scope:** Aria Material

<table>
<thead>
<tr>
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<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

### 4.1.175 Fission Cross Section

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>fiss</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>
4.1.176 Fission Cross Section: Linearized

Scope: Aria Material

Fission Cross Section: Linearized [{of|species|subindex} Species ]= Linearized [{ Sigma_0 = sigma_0 | D_Sigma_D_Rho = d_sigma_d_rho | D_Sigma_D_T = d_sigma_d_t | T_Sigma_0 = t_sigma_0 | Rho_Sigma_0 = rho_sigma_0 }]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>sigma_0</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>d_sigma_d_rho</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>d_sigma_d_t</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>t_sigma_0</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>rho_sigma_0</td>
<td>real</td>
<td>undefined</td>
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</tbody>
</table>

Summary Linearized fission cross section

4.1.177 Flow Area

Scope: Aria Material

Flow Area [{of|species|subindex} Species ]= Constant [{ A = a }]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>a</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary Constant value

4.1.178 Flow Area: Spatial User Function

Scope: Aria Material

Flow Area: Spatial User Function [{of|species|subindex} Species ]= Spatial_User_Function [{ Threshold = threshold | Multiplier = multiplier | Name = name | Bar = bar | X = x }]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>threshold</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>multiplier</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>bar</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>x</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Spatially varying flow area based on a coordinate dependent user function
4.1.179  Flow Area: Correlation Spatial User Function
Scope:  Aria Material

Flow Area: Correlation Spatial User Function [ {of|species|subindex} Species ]= Correlation_Spatial finanzi [ Threshold = threshold |Multiplier = multiplier |Name = name |Bar = bar |X = x |Bulk_Node_Interface = bulk_node_interface ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
<td>threshold</td>
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<td>undefined</td>
</tr>
<tr>
<td>multiplier</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>bar</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>x</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>bulk_node_interface</td>
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<td>undefined</td>
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</tbody>
</table>

Summary  Correlation spatially varying flow area based on a coordinate dependent user function

4.1.180  Flowing Liquid Viscosity
Scope:  Aria Material

Flowing Liquid Viscosity [ {of|species|subindex} Species ]= Constant [ Mu = mu ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Constant value

4.1.181  Fluid Temperature
Scope:  Aria Material

Fluid Temperature [ {of|species|subindex} Species ]= Constant [ T = t ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>t</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Constant value

4.1.182  Forchheimer Drag Coeff
Scope:  Aria Material

Forchheimer Drag Coeff [ {of|species|subindex} Species ]= Constant [ C_F = c_f ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>c_f</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>
### 4.1.183 Freestream Density

**Scope:** Aria Material

Freestream Density \([ \{ \text{of|species|subindex} \} \text{Species } \]= \text{Constant } [ \text{Fsdens} = \text{fsdens} ]\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>fsdens</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

### 4.1.184 Freestream Gamma

**Scope:** Aria Material

Freestream Gamma \([ \{ \text{of|species|subindex} \} \text{Species } \]= \text{Constant } [ \text{G} = \text{g} ]\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>g</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

### 4.1.185 Freestream Pressure

**Scope:** Aria Material

Freestream Pressure \([ \{ \text{of|species|subindex} \} \text{Species } \]= \text{Constant } [ \text{Fspres} = \text{fspres} ]\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>fspres</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

### 4.1.186 Freestream Temperature

**Scope:** Aria Material

Freestream Temperature \([ \{ \text{of|species|subindex} \} \text{Species } \]= \text{Constant } [ \text{Ftemp} = \text{fstemp} ]\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>fstemp</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value
4.1.187 Friction Factor
Scope: Aria Material

Friction Factor \{ \text{of|species|subindex} \ Species \} = \text{Constant} [ \ F = f \ ]

<table>
<thead>
<tr>
<th>Parameter \ Species</th>
<th>Value \ f</th>
<th>Default \ undefined</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>string</td>
<td></td>
</tr>
<tr>
<td></td>
<td>real</td>
<td></td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.188 Friction Factor: Smooth Tube
Scope: Aria Material

Friction Factor: Smooth Tube \{ \text{of|species|subindex} \ Species \} = \text{Smooth\_Tube} [ ]

<table>
<thead>
<tr>
<th>Parameter \ Species</th>
<th>Value \ string</th>
<th>Default \ undefined</th>
</tr>
</thead>
</table>

Summary: Friction factor computed from smooth tube model

4.1.189 Friction Factor: Smooth Annulus
Scope: Aria Material

Friction Factor: Smooth Annulus \{ \text{of|species|subindex} \ Species \} = \text{Smooth\_Annulus} [ R = r ]

<table>
<thead>
<tr>
<th>Parameter \ Species</th>
<th>Value \ string</th>
<th>Default \ undefined</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>&quot;string&quot;</td>
<td></td>
</tr>
</tbody>
</table>

Summary: Friction factor computed from smooth annulus model

4.1.190 Friction Factor: Correlation
Scope: Aria Material

Friction Factor: Correlation \{ \text{of|species|subindex} \ Species \} = \text{Correlation} [ \ F = f \ | \ Bulk\_Node\_Interface = \text{bulk\_node\_interface} ]

<table>
<thead>
<tr>
<th>Parameter \ Species</th>
<th>Value \ string</th>
<th>Default \ undefined</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>&quot;string&quot;</td>
<td></td>
</tr>
</tbody>
</table>

Summary: Correlation friction factor with constant value
4.1.191 Friction Factor: Correlation Smooth Tube
Scope: Aria Material

Friction Factor: Correlation Smooth Tube [{of|species|subindex} Species ] = Correlation_Smooth_Tube [{ Bulk_Node_Interface = bulk_node_interface }]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>bulk_node_interface</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
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</tbody>
</table>

Summary: Correlation friction factor with smooth tube model

4.1.192 Friction Factor: Correlation Smooth Annulus
Scope: Aria Material

Friction Factor: Correlation Smooth Annulus [{of|species|subindex} Species ] = Correlation_Smooth_Annulus [{ Bulk_Node_Interface = bulk_node_interface }]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>bulk_node_interface</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Correlation friction factor with smooth annulus model

4.1.193 Gamma
Scope: Aria Material

Gamma [{of|species|subindex} Species ] = Constant [ G = g ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>g</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.194 Gamma Dot
Scope: Aria Material

Gamma Dot [{of|species|subindex} Species ] = Constant [ Value = value ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
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<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.195 Gap Conductance Coefficient
Scope: Aria Material
Gap Conductance Coefficient \[ \{ \text{of|species|subindex} \} \text{Species } \]= Constant \[ H = h \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\text{Species}</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{h}</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.196 Gap Conductance Coefficient: Calore User Sub
Scope: Aria Material

Gap Conductance Coefficient: Calore User Sub \[ \{ \text{of|species|subindex} \} \text{Species } \]= Calore\_User\_Sub \[ \text{Name } = \text{name } | \text{Type } = \text{type } | \text{Multiplier } = \text{multiplier } | \text{Material\_Data\_Block } = \text{material\_data\_block } | \text{Data } = \text{data } | \text{Scaling\_Field } = \text{scaling\_field } \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\text{Species}</td>
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<td>undefined</td>
</tr>
<tr>
<td>\text{name}</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{type}</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{multiplier}</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{material_data_block}</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{data}</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{scaling_field}</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Values from a Calore user subroutine

4.1.197 Gap Conductance Coefficient: Fortran
Scope: Aria Material

Gap Conductance Coefficient: Fortran \[ \{ \text{of|species|subindex} \} \text{Species } \]= Fortran \[ \text{Multiplier } = \text{multiplier } | \text{Sub\_Name } = \text{sub\_name } | \text{Real\_Data } = \text{real\_data } | \text{Int\_Data } = \text{int\_data } | \text{Resource\_Name } = \text{resource\_name } | \text{Data } = \text{data } \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\text{Species}</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{multiplier}</td>
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<td>undefined</td>
</tr>
<tr>
<td>\text{sub_name}</td>
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<td>undefined</td>
</tr>
<tr>
<td>\text{real_data}</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{int_data}</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{resource_name}</td>
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<td>undefined</td>
</tr>
<tr>
<td>\text{data}</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Fortran subroutine energy source

4.1.198 Gap Height
Scope: Aria Material

Gap Height \[ \{ \text{of|species|subindex} \} \text{Species } \]= Constant \[ H = h \]
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>h</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

### 4.1.199 Gas Phase Retardation

**Scope:** Aria Material

Gas Phase Retardation [ of species subindex Species ] = Constant [ R = r ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>r</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

### 4.1.200 Glass Transition Temperature

**Scope:** Aria Material

Glass Transition Temperature [ of species subindex Species ] = Constant [ Value = value ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

### 4.1.201 Glass Transition Temperature: DeBenedeto

**Scope:** Aria Material

Glass Transition Temperature: DeBenedeto [ of species subindex Species ] = DeBenedeto [ Extent_Species_Name = extent_species_name | A = a | Tg0 = Tg0 | TgInfinity = TgInfinity ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>extent_species_name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>a</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>Tg0</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>TgInfinity</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: deBenedeto model for the glass transition temperature based on the extent of polymerization, x, and material parameters A, Tg0, TgInfinity: 

\[ T_g = \frac{T_{g0}(1-x)+AxT_{g\infty}}{1-x+Ax} \]
4.1.202 Gnielinski Entrance Effect Correction: Correlation
Scope: Aria Material

Gnielinski Entrance Effect Correction: Correlation [ {of|species|subindex} Species ] = Correlation [ Max_Correction = max_correction | Bulk_Node_Interface = bulk_node_interface ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>max_correction</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>bulk_node_interface</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Correction factor for entrance effects for Gnielinski pipe and annulus correlations.

4.1.203 Gnielinski Film Gradient Correction: Correlation
Scope: Aria Material

Gnielinski Film Gradient Correction: Correlation [ {of|species|subindex} Species ] = Correlation [ Phase = phase | Gas_Exponent = gas_exponent | Bulk_Node_Interface = bulk_node_interface ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>phase</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>gas_exponent</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>bulk_node_interface</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Correction factor for difference between near-wall and bulk fluid state for Gnielinski pipe and annulus correlations.

4.1.204 Hdiff Flux: Constant Cl
Scope: Aria Material

Hdiff Flux: Constant Cl [ {of|species|subindex} Species ] = Constant_Cl [ Jx = jx | Jy = jy | Jz = jz ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>jx</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>jy</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>jz</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: ?

4.1.205 Heat Conduction: Generalized
Scope: Aria Material

Heat Conduction: Generalized [ {of|species|subindex} Species ] = Generalized [ ]
Generalized model for heat conduction - can use either a scalar or tensor thermal conductivity.

### 4.1.206 Heat Conduction: Convected Enthalpy

**Scope:** Aria Material

```plaintext
Heat Conduction: Convected Enthalpy [ {of|species|subindex} Species ] = Convected_Enthalpy [ ]
```

**Parameter** | **Value** | **Default**
--- | --- | ---
Species | string | undefined

Summary: Convected enthalpy model for heat conduction

### 4.1.207 Heat Conduction: Phase Average

**Scope:** Aria Material

```plaintext
Heat Conduction: Phase Average [ {of|species|subindex} Species ] = Phase_Average [ ]
```

**Parameter** | **Value** | **Default**
--- | --- | ---
Species | string | undefined

Summary: Phase average model for heat conduction

### 4.1.208 Heat Conduction: Thermoelectric

**Scope:** Aria Material

```plaintext
Heat Conduction: Thermoelectric [ {of|species|subindex} Species ] = Thermoelectric [ ]
```

**Parameter** | **Value** | **Default**
--- | --- | ---
Species | string | undefined

Summary: Thermoelectric model for heat conduction

### 4.1.209 Heat Conduction: Fouriers Law

**Scope:** Aria Material

```plaintext
Heat Conduction: Fouriers Law [ {of|species|subindex} Species ] = Fouriers_Law [ ]
```

**Parameter** | **Value** | **Default**
--- | --- | ---
Species | string | undefined

Summary: Fourier’s Law heat conduction
4.1.210  Heat Conduction: Basic
Scope:  Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{Species}$</td>
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</tr>
</tbody>
</table>

Summary: Basic heat conduction

4.1.211  Heat Conduction: Mass Diffusion Energy Transport
Scope:  Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{Species}$</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Mass diffusion model for energy transport

4.1.212  Heat Conduction: Porous Diffusive Enthalpy Flux
Scope:  Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{Species}$</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Porous diffusive enthalpy flux model

4.1.213  Heat Conduction: Porous Simplified Diffusive Enthalpy Flux
Scope:  Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{Species}$</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Porous simplified diffusive enthalpy flux model

4.1.214  Heat Of Vaporization
Scope:  Aria Material
Heat Of Vaporization [ {of|species|subindex} Species ] = Constant [ Hv = hv ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>hv</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.215 Heat Transfer Coefficient
Scope: Aria Material

Heat Transfer Coefficient [ {of|species|subindex} Species ] = Constant [ H = h ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>h</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.216 Heat Transfer Coefficient: Calore User Sub
Scope: Aria Material

Heat Transfer Coefficient: Calore User Sub [ {of|species|subindex} Species ] = Calore_User_Sub [ Name = name | Type = type | Multiplier = multiplier | Material_Data_Block = material_data_block | Data = data | Scaling_Field = scaling_field ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>type</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>multiplier</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>material_data_block</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>data</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>scaling_field</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Values from a Calore user subroutine

4.1.217 Heat Transfer Coefficient: Aero
Scope: Aria Material

### 4.1.218 Heat Transfer Coefficient: Correlation

**Scope:** Aria Material

Heat Transfer Coefficient: Correlation \[ \{ \text{of } \text{species } \text{subindex} \} \text{Species} \] = Correlation
\[ \text{Re}_x = \text{Re}_x \quad \text{Laminar} = \text{laminar} \quad \text{Turbulent} = \text{turbulent} \quad \text{Bulk Node Interface} = \text{bulk_node_interface} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{Re}_x</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>laminar</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>turbulent</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>bulk_node_interface</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Heat transfer coefficient for external flow about conical tipped body accounting for flow conditions including Mach number, altitude, fluid properties and local flow regime.

### 4.1.219 Heat Transfer Coefficient: Correlation1

**Scope:** Aria Material

Heat Transfer Coefficient: Correlation1 \[ \{ \text{of } \text{species } \text{subindex} \} \text{Species} \] = Correlation1
\[ \text{Laminar} = \text{laminar} \quad \text{Turbulent} = \text{turbulent} \quad \text{Bulk Node Interface} = \text{bulk_node_interface} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>laminar</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>turbulent</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>bulk_node_interface</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Heat transfer coefficient based on type 1 correlation

### 4.1.220 Hydraulic Diameter

**Scope:** Aria Material

Hydraulic Diameter \[ \{ \text{of } \text{species } \text{subindex} \} \text{Species} \] = Constant \[ D = d \]
### 4.1.221 Hydraulic Diameter: Spatial User Function

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
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</tr>
<tr>
<td>(d)</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Spatially varying hydraulic diameter based on a coordinate dependent user function

### 4.1.222 Hydraulic Diameter: Correlation

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>(d)</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>bulk_node_interface</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Correlation hydraulic diameter based on sectional area and wetted perimeter

### 4.1.223 Hydraulic Diameter: Correlation Spatial User Function

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>(d)</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>bulk_node_interface</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Correlation hydraulic diameter based on sectional area and wetted perimeter
Summary: Correlation spatially varying hydraulic diameter based on a coordinate dependent user function

### 4.1.224 Hydrogen Diffusion Supg Tau: Shakib

**Scope:** Aria Material

\[
\text{Hydrogen Diffusion Supg Tau: Shakib} \quad \text{Species} = \text{Shakib} \quad [H = h] \quad \text{Use_Advection} = \text{use_advection} \quad \text{Use_Diffusion} = \text{use_diffusion} \quad \text{Use_Time} = \text{use_time} \quad \text{Multiplier} = \text{multiplier}
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>( h )</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>( \text{use_advection} )</td>
<td>integer</td>
<td>undefined</td>
</tr>
<tr>
<td>( \text{use_diffusion} )</td>
<td>integer</td>
<td>undefined</td>
</tr>
<tr>
<td>( \text{use_time} )</td>
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<td>undefined</td>
</tr>
<tr>
<td>( \text{multiplier} )</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** ?

### 4.1.225 Initial Porosity

**Scope:** Aria Material

\[
\text{Initial Porosity} \quad \text{Species} = \text{Constant} \quad [\Phi = \phi]
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>( \phi )</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** Constant value

### 4.1.226 Intensity Diffusion: Spherical Harmonic

**Scope:** Aria Material

\[
\text{Intensity Diffusion: Spherical Harmonic} \quad \text{Species} = \text{Spherical_Harmonic} \quad [\text{Spn_Order} = \text{SPn_order} \quad \text{Scaled} = \text{scaled}]
\]
Parameter | Value | Default
---|---|---
Species | string | undefined
$SPn\_order$ | integer | undefined
scaled | "string" | undefined

Summary: Spherical harmonic intensity diffusion

### 4.1.227 Internal Energy
**Scope:** Aria Material

Internal Energy $[\{of|species|subindex\} \text{Species}]=\text{Constant} \ [\text{Value} = \text{value}]$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

### 4.1.228 Internal Energy: Gas Phase
**Scope:** Aria Material

Internal Energy: Gas Phase $[\{of|species|subindex\} \text{Species}]=\text{Gas\_Phase} [\ ]$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Gas phase model for internal energy

### 4.1.229 Interphase Friction Coefficient: Wen Yu
**Scope:** Aria Material

Interphase Friction Coefficient: Wen Yu $[\{of|species|subindex\} \text{Species}]=\text{Wen\_Yu} [\ \text{Diameter} = \text{diameter}]$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>diameter</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

### 4.1.230 Intrinsic Permeability
**Scope:** Aria Material

Intrinsic Permeability $[\{of|species|subindex\} \text{Species}]=\text{Constant} \ [\text{Xx} = \text{zx}\ |\ T11 = \text{t11}\ |\ XY = \text{yz}\ |\ T12 = \text{t12}\ |\ XZ = \text{zx}\ |\ T13 = \text{t13}\ |\ YX = \text{yx}\ |\ T21 = \text{t21}\ |\ YY = \text{yy}\ |\ T22 = \text{t22}\ |\ YZ = \text{yz}\ |\ T23 = \text{t23}\ |\ ZX = \text{zx}\ |\ T31 = \text{t31}\ |\ ZZ = \text{zz}\ |\ T32 = \text{t32}\ |\ ZZ = \text{zz}\ |\ T33 = \text{t33}\ ]$

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## 4.1.231 Intrinsic Permeability: Diagonal

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>variable</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

*Intrinsic Permeability: Diagonal {of species subindex} Species = Diagonal [ Variable = variable ]*

## 4.1.232 Intrinsic Permeability: Reordered Adagio

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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</tr>
<tr>
<td>variable</td>
<td>&quot;string&quot;</td>
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</tr>
<tr>
<td>restriction</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

*Intrinsic Permeability: Reordered Adagio {of species subindex} Species = Reordered_Adagio [ Variable = variable | Restriction = restriction ]*

## 4.1.233 Intrinsic Permeability: Log Reordered Adagio

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
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<tbody>
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</tr>
<tr>
<td>variable</td>
<td>&quot;string&quot;</td>
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</tr>
</tbody>
</table>

*Intrinsic Permeability: Log Reordered Adagio {of species subindex} Species = Log_Reordered_Adagio [ Variable = variable ]*

Summary: Constant tensor value for the intrinsic permeability
### 4.1.234 Intrinsic Permeability: Volume Average

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Intrinsic Permeability: Volume Average \[ \{ \text{of|species|subindex} \ Species \} = \text{Volume Average} \[ \] \]

**Summary**

Average intrinsic permeability: \( \text{sum} ( \text{volume_fractions} \times \text{permeabilities} ) / \text{sum} ( \text{volume_fractions} ) \)

### 4.1.235 Intrinsic Permeability: Young And Todd

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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</tr>
<tr>
<td>pore_diameter</td>
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</tr>
<tr>
<td>Liquid_Phase_Name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Intrinsic Permeability: Young And Todd \[ \{ \text{of|species|subindex} \ Species \} = \text{Young And Todd} \[ \text{Pore_Diameter} = \text{pore_diameter} | \text{Liquid_Phase_Name} = \text{Liquid_Phase_Name} \] \]

**Summary**

The Young and Todd permeability model described in SAND2010-0254.

### 4.1.236 Intrinsic Permeability Scaling

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
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<tr>
<td>s</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Intrinsic Permeability Scaling \[ \{ \text{of|species|subindex} \ Species \} = \text{Constant} \[ S = s \] \]

**Summary**

Constant value

### 4.1.237 Intrinsic Permeability Scaling: Kozeny

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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<tr>
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</tr>
<tr>
<td>Phi_Ref</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Intrinsic Permeability Scaling: Kozeny \[ \{ \text{of|species|subindex} \ Species \} = \text{Kozeny} \[ \text{Multiplier} = \text{multiplier} | \text{Phi_Ref} = \text{phi_ref} \] \]
### Summary
Intrinsic Permeability scaling function using Kozeny’s model.

#### 4.1.238 Intrinsic Permeability Scaling: T Exponent

**Scope:** Aria Material

\[
\text{Intrinsic Permeability Scaling: T Exponent [\{of|species|subindex\} Species ]= T_Exponent}
\]

\[
[ K_{\text{Ref}} = k_{\text{ref}} \ | \ T_{\text{Ref}} = t_{\text{ref}} \ | \ N = n ]
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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</tr>
<tr>
<td>(k_{\text{ref}})</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>(t_{\text{ref}})</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>(n)</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Intrinsic permeability scaling factor as a power of normalized temperature.

#### 4.1.239 Intrinsic Permeability Scaling: Forchheimer

**Scope:** Aria Material

\[
\text{Intrinsic Permeability Scaling: Forchheimer [\{of|species|subindex\} Species ]= Forchheimer}
\]

\[
[ \text{Inertia Coeff} = \text{inertia\_coeff} \ | \ G_x = g_x \ | \ G_y = g_y \ | \ G_z = g_z \ | X_x = x_x \ | Y_y = y_y \ | Z_z = z_z \ | X_y = x_y \ | X_z = x_z \ | Y_z = y_z \ | Y_x = y_x \ | Z_x = z_x \ | \text{Fluid Phase} = \text{fluid\_phase} ]
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
<td>inertia_coeff</td>
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<td>undefined</td>
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<tr>
<td>(g_x)</td>
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<td>undefined</td>
</tr>
<tr>
<td>(g_y)</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>(g_z)</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>(x_x)</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>(y_y)</td>
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<tr>
<td>(z_z)</td>
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<td>undefined</td>
</tr>
<tr>
<td>(x_y)</td>
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<td>undefined</td>
</tr>
<tr>
<td>(x_z)</td>
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<td>undefined</td>
</tr>
<tr>
<td>(y_z)</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>(y_x)</td>
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<td>undefined</td>
</tr>
<tr>
<td>(z_x)</td>
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<tr>
<td>(z_y)</td>
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<tr>
<td>fluid_phase</td>
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</tr>
</tbody>
</table>

**Summary**
Intrinsic Permeability scaling for Forchheimer flow.

#### 4.1.240 Inv Neutron Velocity

**Scope:** Aria Material
### 4.1.241 Irradiation

**Scope:** Aria Material

Irradiation \( \{ \text{of|species|subindex} \} \) \( \text{Species} \) = Constant \( [ I = i ] \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( i )</td>
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<td>undefined</td>
</tr>
<tr>
<td>( i )</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

### 4.1.242 Irradiation: Calore User Sub

**Scope:** Aria Material

Irradiation: Calore User Sub \( \{ \text{of|species|subindex} \} \) \( \text{Species} \) = Calore_User_Sub \( [ \text{Name} = \text{name} \text{|Type = type \text{|Multiplier = multiplier \text{|Material_Data_Block = material_data_block \text{|Data = data \text{|Scaling_Field = scaling_field}}}}] } \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \text{Name} )</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>( \text{type} )</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>( \text{Multiplier} )</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>( \text{material_data_block} )</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>( \text{data} )</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>( \text{scaling_field} )</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Values from a Calore user subroutine

### 4.1.243 Isobaric Compressibility

**Scope:** Aria Material

Isobaric Compressibility \( \{ \text{of|species|subindex} \} \) \( \text{Species} \) = Constant \( [ \text{Beta} = \text{beta} ] \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \text{Beta} )</td>
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</tr>
<tr>
<td>( \text{beta} )</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value
4.1.244  K Factor
Scope: Aria Material

K Factor [ {of|species|subindex} Species ]= Constant [ Value = value ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary      Constant value

4.1.245  Kinematic Viscosity: Correlation
Scope: Aria Material

Kinematic Viscosity: Correlation [ {of|species|subindex} Species ]= Correlation [ Nu = nu |Bulk_Node_Interface = bulk_node_interface ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
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<tr>
<td>nu</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>bulk_node_interface</td>
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</tr>
</tbody>
</table>

Summary          Correlation kinematic viscosity

4.1.246  Laser Power
Scope: Aria Material

Laser Power [ {of|species|subindex} Species ]= Constant [ P = p ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>p</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary      Constant value

4.1.247  Latent Heat
Scope: Aria Material

Latent Heat [ {of|species|subindex} Species ]= Constant [ Value = value ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary      Constant value
4.1.248  Level Set
Scope:  Aria Material

Level Set \[ \{ \text{of|species|subindex} \ S\text{pecies} \} = \text{Constant} \ [ \ \text{Value} = \text{value} \ ] \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
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</tbody>
</table>

Summary  Constant value

4.1.249  Level Set Curvature
Scope:  Aria Material

Level Set Curvature \[ \{ \text{of|species|subindex} \ S\text{pecies} \} = \text{Constant} \ [ \ K\text{appa} = \kappa \ ] \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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<tbody>
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</table>

Summary  Constant value

4.1.250  Level Set Diffusive Flux: Shock Capturing
Scope:  Aria Material

Level Set Diffusive Flux: Shock Capturing \[ \{ \text{of|species|subindex} \ S\text{pecies} \} = \text{Shock}\_\text{Capturing} \ [ \ \text{Multiplier} = \text{multiplier} \ ] \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
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<tbody>
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<tr>
<td>multiplier</td>
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</tbody>
</table>

Summary  Shock capturing operator for the level set equation

4.1.251  Level Set Diffusive Flux: Unit Gradient
Scope:  Aria Material

Level Set Diffusive Flux: Unit Gradient \[ \{ \text{of|species|subindex} \ S\text{pecies} \} = \text{Unit}\_\text{Gradient} \ [ \ ] \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tbody>
</table>

Summary  Unit gradient model of level set diffusive flux
4.1.252  Level Set Diffusive Flux

Scope:  Aria Material

Level Set Diffusive Flux $[\{\text{of|species|subindex}\} \text{Species}] = \text{Constant} [\text{Coeff} = \text{coeff}]$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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<td>coeff</td>
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</tbody>
</table>

Summary  Constant model of level set diffusive flux

4.1.253  Level Set Heaviside

Scope:  Aria Material

Level Set Heaviside $[\{\text{of|species|subindex}\} \text{Species}] = \text{Constant} [\text{Value} = \text{value}]$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>value</td>
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</tr>
</tbody>
</table>

Summary  Constant value

4.1.254  Level Set Heaviside: Smooth

Scope:  Aria Material

Level Set Heaviside: Smooth $[\{\text{of|species|subindex}\} \text{Species}] = \text{Smooth} [\text{Use\_Crossing\_Time} = \text{use\_crossing\_time}]$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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<tbody>
<tr>
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<tr>
<td>use_crossing_time</td>
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</table>

Summary  Smooth level set Heaviside

4.1.255  Level Set Heaviside: Interpolated

Scope:  Aria Material

Level Set Heaviside: Interpolated $[\{\text{of|species|subindex}\} \text{Species}] = \text{Interpolated} [\text{Epsilon} = \text{epsilon}]$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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</table>

Summary  Interpolated level set Heaviside
4.1.256  **Level Set Heaviside: Bf Interpolated**  
**Scope:**  Aria Material

<table>
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<tr>
<th>Parameter</th>
<th>Value</th>
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Summary  
BF interpolated level set Heaviside

4.1.257  **Level Set Heaviside: Sharp Analytic**  
**Scope:**  Aria Material

<table>
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<th>Value</th>
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</table>

Summary  
Sharp analytic level set Heaviside

4.1.258  **Level Set Width**  
**Scope:**  Aria Material

<table>
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<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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<tr>
<td>width</td>
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</table>

Summary  
Specification of a constant width diffuse level set

4.1.259  **Liquid Phase Retardation**  
**Scope:**  Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
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<tbody>
<tr>
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</tr>
<tr>
<td>r</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary  
Constant value

4.1.260  **Lubrication Height**  
**Scope:**  Aria Material

128
### Lubrication Height \( \text{Species} \) = Constant \( H = h \)

<table>
<thead>
<tr>
<th>Parameter ( \text{Species} )</th>
<th>Value ( h )</th>
<th>Default undefined</th>
</tr>
</thead>
</table>

**Summary**  
Constant value

#### 4.1.261 Lubrication Height: Combined

**Scope**: Aria Material

<table>
<thead>
<tr>
<th>Parameter ( \text{Species} )</th>
<th>Value ( h )</th>
<th>Default undefined</th>
</tr>
</thead>
</table>

**Summary**  
Calculate the lubrication height by combining the upper and lower heights

#### 4.1.262 Lubrication Height Lower

**Scope**: Aria Material

<table>
<thead>
<tr>
<th>Parameter ( \text{Species} )</th>
<th>Value ( h_{\text{lower}} )</th>
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</table>

**Summary**  
Constant value

#### 4.1.263 Lubrication Height Upper

**Scope**: Aria Material

<table>
<thead>
<tr>
<th>Parameter ( \text{Species} )</th>
<th>Value ( h_{\text{upper}} )</th>
<th>Default undefined</th>
</tr>
</thead>
</table>

**Summary**  
Constant value

#### 4.1.264 Lubrication K

**Scope**: Aria Material

<table>
<thead>
<tr>
<th>Parameter ( \text{Species} )</th>
<th>Value ( k )</th>
<th>Default undefined</th>
</tr>
</thead>
</table>
Summary  Constant value

4.1.265  Lubrication Velocity Lower
Scope:  Aria Material

Lubrication Velocity Lower [ {of|species|subindex} Species ] = Constant [ V_Lowerx = v_lowerx \ |
V_Lowery = v_lowery \ | V_Lowerz = v_lowerz ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>v_lowerx</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>v_lowery</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>v_lowerz</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Constant value

4.1.266  Lubrication Velocity Upper
Scope:  Aria Material

Lubrication Velocity Upper [ {of|species|subindex} Species ] = Constant [ V_Upperx = v_upperx \ |
V_Uppery = v_uppery \ | V_Upperz = v_upperz ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>v_upperx</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>v_uppery</td>
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<td>undefined</td>
</tr>
<tr>
<td>v_upperz</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Constant value

4.1.267  Mach Number
Scope:  Aria Material

Mach Number [ {of|species|subindex} Species ] = Constant [ Ma = Ma ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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</tr>
<tr>
<td>Ma</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Constant value
4.1.268  Masking
Scope:  Aria Material

Masking [ {of|species|subindex} Species ]= Constant [ Value = value ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Constant value

4.1.269  Mass Balance Advective Flux: Porous
Scope:  Aria Material

Mass Balance Advective Flux: Porous [ {of|species|subindex} Species ]= Porous [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Mass balance advection term from mass fraction and mass flux.

4.1.270  Mass Balance Advective Flux: Porous Upwind
Scope:  Aria Material

Mass Balance Advective Flux: Porous Upwind [ {of|species|subindex} Species ]= Porous_Upwind [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Mass balance advection term from mass fraction and mass flux.

4.1.271  Mass Balance Advective Flux: Single Phase
Scope:  Aria Material

Mass Balance Advective Flux: Single Phase [ {of|species|subindex} Species ]= Single_Phase [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Mass balance advection term from density and velocity

4.1.272  Mass Balance Diffusive Flux: Porous
Scope:  Aria Material
Mass Balance Diffusive Flux: Porous [ {of|species|subindex} Species ] = Porous [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Mass balance diffusive term from porosity, mass diffusivity, mass fraction and density.

4.1.273 Mass Balance Diffusive Flux: Nernst Planck
Scope: Aria Material

Mass Balance Diffusive Flux: Nernst Planck [ {of|species|subindex} Species ] = Nernst_Planck [ F = f \ R = r ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>f</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>r</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.274 Mass Balance Diffusive Flux: Density
Scope: Aria Material

Mass Balance Diffusive Flux: Density [ {of|species|subindex} Species ] = Density [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Mass balance diffusive term from porosity, mass diffusivity, mass fraction and density.

4.1.275 Mass Balance Diffusive Flux: Basic
Scope: Aria Material

Mass Balance Diffusive Flux: Basic [ {of|species|subindex} Species ] = Basic [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Mass balance diffusive term from mass diffusivity, mass fraction and density.

4.1.276 Mass Diffusivity
Scope: Aria Material

Mass Diffusivity [ {of|species|subindex} Species ] = Constant [ D = d ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>d</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>
### 4.1.277 Mass Diffusivity: Cantera

**Scope:** Aria Material

Mass Diffusivity: Cantera \( \{ \text{of|species|subindex} \} \) Species \( \Rightarrow \) Cantera \( \) 

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Cantera mass diffusivity

### 4.1.278 Mass Diffusivity: From Schmidt

**Scope:** Aria Material

Mass Diffusivity: From Schmidt \( \{ \text{of|species|subindex} \} \) Species \( \Rightarrow \) From_Schmidt \( \) 

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Mass diffusivity from a Schmidt number and viscosity

### 4.1.279 Mass Diffusivity: Air Water

**Scope:** Aria Material

Mass Diffusivity: Air Water \( \{ \text{of|species|subindex} \} \) Species \( \Rightarrow \) Air_Water \( \) Tortuosity \( = \) tortuosity \| P_Ref = p_ref \| D_Ref = d_ref \| Exponent = exponent \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>tortuosity</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>p_ref</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>d_ref</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>exponent</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Air/Water mass diffusivity

### 4.1.280 Mass Flux: Darcy

**Scope:** Aria Material

Mass Flux: Darcy \( \{ \text{of|species|subindex} \} \) Species \( \Rightarrow \) Darcy \( \) 

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Air/Water mass diffusivity
4.1.281  Mass Fraction
Scope:  Aria Material

Mass Fraction \( \{ \text{of|species|subindex} \} \SPECIES \) = Constant \( \{ \text{Value} = \text{value} \} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\SPECIES</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{value}</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Constant value

4.1.282  Mass Fraction: Fracbal
Scope:  Aria Material

Mass Fraction: Fracbal \( \{ \text{of|species|subindex} \} \SPECIES \) = Fracbal \[ \]  

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\SPECIES</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Mass fraction of one species to yield a unity sum over all species

4.1.283  Mass Fraction: From Chemeq
Scope:  Aria Material

Mass Fraction: From Chemeq \( \{ \text{of|species|subindex} \} \SPECIES \) = From_Chemeq \[ \]  

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\SPECIES</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Mass fraction of one species extracted from ChemEq

4.1.284  Mass Fraction Diffusive Flux: Basic
Scope:  Aria Material

Mass Fraction Diffusive Flux: Basic \( \{ \text{of|species|subindex} \} \SPECIES \) = Basic \[ \]  

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\SPECIES</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Basic model of mass fraction diffusive flux

Scope:  Aria Material

Mass Fraction Diffusive Flux: Energy Mass \( \{ \text{of|species|subindex} \} \SPECIES \) = Energy_Mass \[ \]
### Parameter | Value | Default
--- | --- | ---
Species | string | undefined

**Summary**
Energy_Mass model of mass fraction diffusive flux

#### 4.1.286 Material Data Block: Resource

**Scope:** Aria Material

Material Data Block: Resource [ {of|species|subindex} Species ]= Resource [ Data_Block_Name = data_block_name ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>data_block_name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Resource material data block

#### 4.1.287 Melt Temperature

**Scope:** Aria Material

Melt Temperature [ {of|species|subindex} Species ]= Constant [ Value = value ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Constant value

#### 4.1.288 Mesh Body Acceleration

**Scope:** Aria Material

Mesh Body Acceleration [ {of|species|subindex} Species ]= Constant [ Valuex = valuex | Valuey = valuey | Valuez = valuez ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>valuex</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>valuey</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>valuez</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Constant value

#### 4.1.289 Mesh Lambda

**Scope:** Aria Material

Mesh Lambda [ {of|species|subindex} Species ]= Constant [ Lambda = lambda ]
Summary: Constant value

4.1.290 Mesh Lambda: Converted
Scope: Aria Material

Parameter Value Default
Species string undefined
\( \text{lambda} \) real undefined

Summary: Lame coefficient computed from Poisson’s ratio and elastic modulus

4.1.291 Mesh Lambda: Converted Plane Stress
Scope: Aria Material

Parameter Value Default
Species string undefined

Summary: Lame coefficient applicable to plane stress conditions computed from Poisson’s ratio and elastic modulus

4.1.292 Mesh Lambda: Elemental Volume
Scope: Aria Material

Parameter Value Default
Species string undefined

Summary: Lambda computed from elemental volume

4.1.293 Mesh Lambda: Inverse Element Volume
Scope: Aria Material

Parameter Value Default
Species string undefined
lagged integer undefined

136
4.1.294 Mesh Poissons Ratio
Scope: Aria Material

Mesh Poissons Ratio \([ \{of|species|subindex\} \text{Species} ] = \text{Constant} \[ \nu = \nu \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>(\nu)</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Constant value

4.1.295 Mesh Stress: Isothermal
Scope: Aria Material

Mesh Stress: Isothermal \([ \{of|species|subindex\} \text{Species} ] = \text{Isothermal} \[ T = t \ | T_{\text{Ref}} = t_{\text{ref}} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>(t)</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>(t_{\text{ref}})</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.296 Mesh Stress: Linear Elastic
Scope: Aria Material

Mesh Stress: Linear Elastic \([ \{of|species|subindex\} \text{Species} ] = \text{Linear\_Elastic} \[ \text{Reference\_Frame} = \text{reference\_frame} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>\text{reference_frame}</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.297 Mesh Stress: Porous Effective
Scope: Aria Material

Mesh Stress: Porous Effective \([ \{of|species|subindex\} \text{Species} ] = \text{Porous\_Effective} \[ \text{Pressure} = \text{pressure} \ | \text{Phase} = \text{phase} \ | \text{Biot} = \text{biot} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>\text{pressure}</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{phase}</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{biot}</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Apply the pressure as an isotropic mesh stress. Primarily used as an effective stress in porous flow problems.
4.1.298  Mesh Stress: Saturation Weighted Porous Effective
Scope:  Aria Material

Mesh Stress: Saturation Weighted Porous Effective \([\{\text{of/species/subindex}\} \text{Species }]= \text{Saturation\_Weighted\_Porous\_Effective} \)

- **Parameter**
  - *Species*
  - *pressure*
  - *phase*
  - *biot*

- **Value**
  - *string*
  - "string"
  - "string"
  - real

- **Default**
  - undefined
  - undefined
  - undefined

Summary  Apply the saturation-weighted pressure as an isotropic mesh stress. Primarily used as an effective stress in porous flow problems.

4.1.299  Mesh Stress: Mooney Rivlin
Scope:  Aria Material

Mesh Stress: Mooney Rivlin \([\{\text{of/species/subindex}\} \text{Species }]= \text{Mooney\_Rivlin} \)

- **Parameter**
  - *Species*

- **Value**
  - *string*

- **Default**
  - undefined

4.1.300  Mesh Stress: Neoohookean Elastic
Scope:  Aria Material

Mesh Stress: Neoohookean Elastic \([\{\text{of/species/subindex}\} \text{Species }]= \text{Neoohookean\_Elastic} \)

- **Parameter**
  - *Species*

- **Value**
  - *string*

- **Default**
  - undefined

4.1.301  Mesh Stress: Nonlinear Elastic
Scope:  Aria Material

Mesh Stress: Nonlinear Elastic \([\{\text{of/species/subindex}\} \text{Species }]= \text{Nonlinear\_Elastic} \)

- **Parameter**
  - *Species*

- **Value**
  - *string*

- **Default**
  - undefined

4.1.302  Mesh Stress: Residual
Scope:  Aria Material

Mesh Stress: Residual \([\{\text{of/species/subindex}\} \text{Species }]= \text{Residual} \)

- **Parameter**
  - *Species*

- **Value**
  - *Sz = sz |
  - *Szz = szz |
  - *Sxz = szx |
  - *Syz = szy |
  - *Sy = sy |
  - *Syy = syy |
  - *Sxy = sxy |
  - *Sx = sx |
  - *Sxx = sxx |

138
<table>
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<td>szz</td>
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<td>sxz</td>
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<tr>
<td>syz</td>
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<td>sy</td>
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<td>syy</td>
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<tr>
<td>sxx</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

### 4.1.303  Mesh Stress: Thermal

Scope: Aria Material

Mesh Stress: Thermal [ of\{species\subindex\} Species ] = Thermal [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

### 4.1.304  Mesh Stress: Electrode

Scope: Aria Material

Mesh Stress: Electrode [ of\{species\subindex\} Species ] = Electrode [ Electrodefile = electrodeFile | F = f | Kmolconversion = kmolConversion | Jconversion = JConversion | Meterconversion = meterConversion | MinCapacity = minCapacity | DeactivationVoltage = deactivationVoltage | Voltagevariablename = voltageVariableName | Abortaction = abortAction | Inactivesolidfraction = inactiveSolidFraction ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>electrodeFile</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>f</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>kmolConversion</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>JConversion</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>meterConversion</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>minCapacity</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>deactivationVoltage</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>voltageVariableName</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>abortAction</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>inactiveSolidFraction</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

### 4.1.305  Mesh Stress: Electrode Old

Scope: Aria Material

Mesh Stress: Electrode Old [ of\{species\subindex\} Species ] = Electrode_Old [ F = f | Electrodefile = electrodeFile | Kmolconversion = kmolConversion | Jconversion = JConversion | Meterconversion = meterConversion | Max_Sub_Timesteps = max_sub_timesteps | Base_Delta = base_delta ]
### 4.1.306 Mesh Stress: Species

**Scope:** Aria Material

Mesh Stress: Species [ {of species subindex} Species ]= Species [ Sref = sref | Beta = beta | Phase = phase ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>sref</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>beta</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>phase</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
A linear mechanical strain due to a species concentration

### 4.1.307 Mesh Stress: Species Transversely Isotropic

**Scope:** Aria Material

Mesh Stress: Species Transversely Isotropic [ {of species subindex} Species ]= Species_Transversely_Isotropic [ Refval = refval | Beta_N = beta_n | Beta_T = beta_t | N_X = n_x | N_Y = n_y | N_Z = n_z | Phase = phase ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>refval</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>beta_n</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>beta_t</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>n_x</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>n_y</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>n_z</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>phase</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
A linear mechanical strain that is transversely isotropic due to a species concentration

### 4.1.308 Mesh Stress: J

**Scope:** Aria Material

Mesh Stress: J [ {of species subindex} Species ]= J [ H = h | Yield Stress = yield stress ]

140
### 4.1.309 Mesh Stress: Lame

**Scope:** Aria Material

Mesh Stress: Lame 

<table>
<thead>
<tr>
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<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
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</tbody>
</table>

### 4.1.310 Mesh Two Mu

**Scope:** Aria Material

Mesh Two Mu 

<table>
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<th>Value</th>
<th>Default</th>
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</table>

**Summary** Constant value

### 4.1.311 Mesh Two Mu: Converted

**Scope:** Aria Material

Mesh Two Mu: Converted 

<table>
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<tr>
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<th>Value</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
</tbody>
</table>

**Summary** Lame coefficient computed from Poisson’s ratio and elastic modulus

### 4.1.312 Mesh Two Mu: Elemental Volume

**Scope:** Aria Material

Mesh Two Mu: Elemental Volume 

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
<tr>
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</tr>
</tbody>
</table>

**Summary** 2 mu computed from elemental volume
4.1.313  Mesh Two Mu: Inverse Element Volume
Scope: Aria Material

Mesh Two Mu: Inverse Element Volume [ {of|species|subindex} Species ] = Inverse_Element_Volume [ Lagged = lagged ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
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<tr>
<td>lagged</td>
<td>integer</td>
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</tr>
</tbody>
</table>

Summary 2 mu computed from inverse of element volume

4.1.314  Mesh Youngs Modulus
Scope: Aria Material

Mesh Youngs Modulus [ {of|species|subindex} Species ] = Constant [ E = e ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>e</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Constant value

4.1.315  Mixture Fraction
Scope: Aria Material

Mixture Fraction [ {of|species|subindex} Species ] = Constant [ Value = value ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>value</td>
<td>real</td>
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</tr>
</tbody>
</table>

Summary Constant value

4.1.316  Mixture Fraction Diffusive Flux: Basic
Scope: Aria Material

Mixture Fraction Diffusive Flux: Basic [ {of|species|subindex} Species ] = Basic [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
</tbody>
</table>

Summary Basic model of mixture fraction diffusive flux

4.1.317  Mixture Fraction Diffusivity
Scope: Aria Material
Mixture Fraction Diffusivity [\{of|species|subindex\} Species ] = Constant [ \( D_z = D_z \) ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>( D_z )</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

### 4.1.318 Mixture Fraction Diffusivity: From Schmidt

Scope: Aria Material

Mixture Fraction Diffusivity: From Schmidt [\{of|species|subindex\} Species ] = From_Schmidt

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
</tbody>
</table>

Summary: Mixture fraction diffusivity from Schmidt number and viscosity

### 4.1.319 Mixture Mass Diffusivity

Scope: Aria Material

Mixture Mass Diffusivity [\{of|species|subindex\} Species ] = Constant [ \( D_{mix} = D_{mix} \) ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>( D_{mix} )</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

### 4.1.320 Mixture Mass Diffusivity: Mixture Average

Scope: Aria Material

Mixture Mass Diffusivity: Mixture Average [\{of|species|subindex\} Species ] = Mixture_Average

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Linear mixture-averaged mass diffusivity

### 4.1.321 Mixture Molecular Weight

Scope: Aria Material

Mixture Molecular Weight [\{of|species|subindex\} Species ] = Constant [ \( M_{mix} = M_{mix} \) ]

143
Parameter | Value | Default
---|---|---
Species | string | undefined
Mmix | real | undefined

Summary
Constant value

4.1.322  Mixture Molecular Weight: Cantera
Scope:  Aria Material

Mixture Molecular Weight: Cantera [{of|species|subindex} Species ] = Cantera [

Parameter | Value | Default
---|---|---
Species | string | undefined

Summary
Cantera mixture molecular weight

4.1.323  Mixture Molecular Weight: Mass Average
Scope:  Aria Material

Mixture Molecular Weight: Mass Average [{of|species|subindex} Species ] = Mass_Average [

Parameter | Value | Default
---|---|---
Species | string | undefined

Summary
Mass averaged mixture molecular weight

4.1.324  Mixture Molecular Weight: Mole Average
Scope:  Aria Material

Mixture Molecular Weight: Mole Average [{of|species|subindex} Species ] = Mole_Average [

Parameter | Value | Default
---|---|---
Species | string | undefined

Summary
Mole averaged mixture molecular weight

4.1.325  Molecular Weight
Scope:  Aria Material

Molecular Weight [{of|species|subindex} Species ] = Constant [ M = m ]

Parameter | Value | Default
---|---|---
Species | string | undefined
m | real | undefined

144
4.1.326 Molecular Weight: Cantera
Scope: Aria Material

Molecular Weight: Cantera [ {of|species|subindex} Species ] = Cantera [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary Cantera molecular weight

4.1.327 Molecular Weight: Dynamic Species
Scope: Aria Material

Molecular Weight: Dynamic Species [ {of|species|subindex} Species ] = Dynamic_Species [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Calculate the molecular weight dynamically from density/molar concentration.

4.1.328 Momentum Face Stabilization Scaling: Default
Scope: Aria Material

Summary Default face stabilization scaling for momentum equation

4.1.329 Momentum Stress: Bad Sensitivity
Scope: Aria Material

Momentum Stress: Bad Sensitivity [ {of|species|subindex} Species ] = Bad_Sensitivity [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.330 Momentum Stress: Balanced Force
Scope: Aria Material

Momentum Stress: Balanced Force [ {of|species|subindex} Species ] = Balanced_Force [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>
4.1.331 Momentum Stress: Grad Div

Scope: Aria Material

Momentum Stress: Grad Div [ {of|species|subindex} Species ] = Grad_Div [ Pressure = pressure | Multiplier = multiplier ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>pressure</td>
<td>&quot;string&quot;</td>
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<tr>
<td>multiplier</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.332 Momentum Stress: Formal Newtonian

Scope: Aria Material


<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
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<tbody>
<tr>
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<tr>
<td>pressure</td>
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<tr>
<td>phase</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
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<tr>
<td>biot</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>reference_frame</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.333 Momentum Stress: Incompressible Newtonian

Scope: Aria Material

Momentum Stress: Incompressible Newtonian [ {of|species|subindex} Species ] = Incompressible_Newtonian [ Pressure = pressure | Phase = phase | Biot = biot | Reference_Frame = reference_frame ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>pressure</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>phase</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
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<tr>
<td>biot</td>
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</tr>
<tr>
<td>reference_frame</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.334 Momentum Stress: Suspension

Scope: Aria Material

Momentum Stress: Suspension [ {of|species|subindex} Species ] = Suspension [ Pressure = pressure | Phase = phase | Biot = biot | Reference_Frame = reference_frame ]
### 4.1.335 Momentum Stress: Ls Capillary

Scope: Aria Material

Momentum Stress: Ls Capillary \( \{ \text{of|species|subindex} \, Species \} = \text{Ls\_Capillary} \) [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

### 4.1.336 Momentum Stress: Ls Implicit Capillary

Scope: Aria Material

Momentum Stress: Ls Implicit Capillary \( \{ \text{of|species|subindex} \, Species \} = \text{Ls\_Implicit\_Capillary} \) [ \( \mu_0 = \mu_0 \) ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>( \mu_0 )</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

### 4.1.337 Momentum Stress: Ls Implicit Normal Capillary

Scope: Aria Material

Momentum Stress: Ls Implicit Normal Capillary \( \{ \text{of|species|subindex} \, Species \} = \text{Ls\_Implicit\_Normal\_Capillary} \) [ \( \mu_0 = \mu_0 \) ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>( \mu_0 )</td>
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<td>undefined</td>
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</tbody>
</table>

### 4.1.338 Momentum Stress: Newtonian Dilational

Scope: Aria Material

Momentum Stress: Newtonian Dilational \( \{ \text{of|species|subindex} \, Species \} = \text{Newtonian\_Dilational} \) [ \( \text{Reference\_Frame = reference\_frame} \) ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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</tr>
</thead>
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<tr>
<td>Species</td>
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</table>

### 4.1.339 Momentum Stress: Newtonian Viscous

Scope: Aria Material
Momentum Stress: Newtonian Viscous \( \{\text{of|species|subindex}\} \) \( \text{Species} \) = Newtonian_Viscous
[Reference Frame = reference_frame]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
<tr>
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</tr>
<tr>
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</tbody>
</table>

4.1.340  Momentum Stress: Newtonian Pressure
Scope:  Aria Material

Momentum Stress: Newtonian Pressure \( \{\text{of|species|subindex}\} \) \( \text{Species} \) = Newtonian_Pressure
[Pressure = pressure | Phase = phase | Biot = biot]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
<tr>
<td>Species</td>
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</tbody>
</table>

4.1.341  Momentum Stress: Maxwell
Scope:  Aria Material

Momentum Stress: Maxwell \( \{\text{of|species|subindex}\} \) \( \text{Species} \) = Maxwell [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary
The gradient of the Maxwell stress tensor is the force on a fluid due to an electrical field and is non-zero when there is either a net charge density or a non-uniform electric permittivity.
\[
\nabla \cdot \mathbf{M} = \rho_{\text{charge}} \mathbf{E} - \frac{1}{2} (\mathbf{E} \cdot \mathbf{E}) \nabla \epsilon
\n\]

4.1.342  Neutron
Scope:  Aria Material

Neutron \( \{\text{of|species|subindex}\} \) \( \text{Species} \) = Constant [ Value = value ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
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</tbody>
</table>

Summary
Constant value

4.1.343  Neutron Diffusion: Ficks Law
Scope:  Aria Material

Neutron Diffusion: Ficks Law \( \{\text{of|species|subindex}\} \) \( \text{Species} \) = Ficks_Law [ ]
### 4.1.344 Neutron Diffusion: Basic

**Scope:** Aria Material

<table>
<thead>
<tr>
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<th>Value</th>
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<tbody>
<tr>
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</table>

Summary: Basic neutron diffusion

### 4.1.345 Neutron Diffusivity

**Scope:** Aria Material

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<tr>
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<th>Value</th>
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</tr>
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</table>

Summary: Constant value

### 4.1.346 Neutron Diffusivity: Five Ten

**Scope:** Aria Material

<table>
<thead>
<tr>
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<th>Value</th>
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</tr>
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<tbody>
<tr>
<td><em>Species</em></td>
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</tbody>
</table>

Summary: Five ten neutron diffusivity

### 4.1.347 Nonwetting Phase Retardation

**Scope:** Aria Material

<table>
<thead>
<tr>
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<th>Value</th>
<th>Default</th>
</tr>
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<tbody>
<tr>
<td><em>Species</em></td>
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<td>undefined</td>
</tr>
<tr>
<td><em>r</em></td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value
4.1.348 Partial Enclosure Area

Scope: Aria Material

Partial Enclosure Area [ {of|species|subindex} Species ] = Constant [ A = a ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>a</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.349 Partial Enclosure Area: Calore User Sub

Scope: Aria Material

Partial Enclosure Area: Calore User Sub [ {of|species|subindex} Species ] = Calore_User_Sub

[ Name = name | Type = type | Multiplier = multiplier | Material_Data_Block = material_data_block | Data = data | Scaling_Field = scaling_field ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
<td>name</td>
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<tr>
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</tr>
<tr>
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<td>undefined</td>
</tr>
<tr>
<td>data</td>
<td>&quot;string&quot;</td>
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</tr>
<tr>
<td>scaling_field</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Values from a Calore user subroutine

4.1.350 Partial Enclosure Emissivity

Scope: Aria Material

Partial Enclosure Emissivity [ {of|species|subindex} Species ] = Constant [ E = e ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>e</td>
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</tbody>
</table>

Summary: Constant value

4.1.351 Partial Enclosure Emissivity: Calore User Sub

Scope: Aria Material

Partial Enclosure Emissivity: Calore User Sub [ {of|species|subindex} Species ] = Calore_User_Sub

[ Name = name | Type = type | Multiplier = multiplier | Material_Data_Block = material_data_block | Data = data | Scaling_Field = scaling_field ]
### 4.1.352 Partial Enclosure Temperature

**Scope:** Aria Material

Partial Enclosure Temperature \( \{ \text{of|species|subindex} \) Species \} = \text{Constant [ T = t ]}

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<tbody>
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</tbody>
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**Summary**

Values from a Calore user subroutine

### 4.1.353 Partial Enclosure Temperature: Calore User Sub

**Scope:** Aria Material

Partial Enclosure Temperature: Calore User Sub \( \{ \text{of|species|subindex} \) Species \} = \text{Calore_User_Sub [ Name = name |Type = type |Multiplier = multiplier |Material_Data_Block = material_data_block |Data = data |Scaling_Field = scaling_field ]}

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**Summary**

Values from a Calore user subroutine

### 4.1.354 Partial Molar Volume

**Scope:** Aria Material

Partial Molar Volume \( \{ \text{of|species|subindex} \) Species \} = \text{Constant [ Value = value ]}

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<th>Value</th>
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<tbody>
<tr>
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<tr>
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</table>
4.1.355  Peltier Coefficient
Scope:  Aria Material

\[
\text{Peltier Coefficient \ [ \{of\,species\,subindex\} \ Specie s \] = Constant \ [ \ Value = \ value \ ]}
\]

<table>
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<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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<tr>
<td>value</td>
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<td>undefined</td>
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</tbody>
</table>

4.1.356  Phase Change Specific Heat
Scope:  Aria Material

\[
\text{Phase Change Specific Heat \ [ \{of\,species\,subindex\} \ Specie s \] = Constant \ [ \ Cp = \ cp \ ]}
\]

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</table>

4.1.357  Phase Change Specific Heat: Calore User Sub
Scope:  Aria Material

\[
\text{Phase Change Specific Heat: Calore User Sub \ [ \{of\,species\,subindex\} \ Specie s \] = Calore_User_Sub \ [ \ Name = \ name \ | Type = \ type \ | Multiplier = \ multiplier \ | Material_Data_Block = \ material\_data\_block \ | Data = \ data \ | Scaling_Field = \ scaling\_field \ ]}
\]

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</table>

Summary  Values from a Calore user subroutine

4.1.358  Phase Change Specific Heat: Curing Foam
Scope:  Aria Material

\[
\text{Phase Change Specific Heat: Curing Foam \ [ \{of\,species\,subindex\} \ Specie s \] = Curing_Foam \ [ \ ]}
\]
Summary  Phase change specific heat for a curing foam

4.1.359  Phi
Scope:  Aria Material

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</table>

Summary  Constant value

4.1.360  Polymerization Shift Factor
Scope:  Aria Material

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Summary  Constant value

4.1.361  Polymerization Shift Factor: Wlf
Scope:  Aria Material

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4.1.362  Porosity
Scope:  Aria Material

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</tbody>
</table>
4.1.363 Porosity: Deforming
Scope: Aria Material

Porosity: Deforming [of{species|subindex} Species] = Deforming [Phi_Init = phi_init]

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Summary Porosity accounting for deforming media

4.1.364 Porosity: Mesh Deforming
Scope: Aria Material


<table>
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</table>

Summary Rock compressibility linearized model of porosity for deforming media

4.1.365 Porosity: Solid Deforming
Scope: Aria Material

Porosity: Solid Deforming [of{species|subindex} Species] = Solid_Deforming [Biot = biot]

<table>
<thead>
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<th>Default</th>
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<tbody>
<tr>
<td>Species</td>
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<tr>
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</table>

Summary Rock compressibility linearized model of porosity for deforming media

4.1.366 Porosity: Rock Compressible
Scope: Aria Material

Porosity: Rock Compressible [of{species|subindex} Species] = Rock_Compressible [Cr = cr |Ref_Porosity = ref_porosity |Ref_Pressure = ref_pressure]
4.1.367 Porosity: Deformable Rock Compressible

Scope: Aria Material

Porosity: Deformable Rock Compressible [ {of|species|subindex} Species ]= Deformable_Rock_Compressible [ Cr = cr | Ref_Porosity = ref_porosity | Ref_Pressure = ref_pressure | Biot = biot ]

Summary Rock compressibility linearized model of porosity for deforming media

4.1.368 Porosity: Coussy

Scope: Aria Material

Porosity: Coussy [ {of|species|subindex} Species ]= Coussy [ M_Inv = m_inv | Biot = biot | Ref_Porosity = ref_porosity | Ref_Pressure = ref_pressure ]

Summary Coussy’s model of porosity for deforming media (based on pore pressure and div(u))

4.1.369 Porosity: From Density Ratio

Scope: Aria Material

Porosity: From Density Ratio [ {of|species|subindex} Species ]= From_Density_Ratio [ Solid_Density = solid_density ]

Summary
Porosity computed as one minus the density ratio between solved for solid-phase density and solid density of non-porous material.

4.1.370 Porosity: One Minus Volume Fractions
Scope: Aria Material

Porosity: One Minus Volume Fractions [ {of|species|subindex} Species ] = One_Minus_Volume_Fractions

<table>
<thead>
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<tbody>
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</tr>
</tbody>
</table>

Summary Porosity computed directly from volume fractions of species.

4.1.371 Porosity: Constant From Electrode Object Old
Scope: Aria Material

Porosity: Constant From Electrode Object Old [ {of|species|subindex} Species ] = Constant_From_Electrode_Object_Old [ Electrodefile = electrodeFile ]

<table>
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<th>Parameter</th>
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</table>

Summary Constant porosity extracted from Cantera thermal battery electrode object.

4.1.372 Porosity Def
Scope: Aria Material

Porosity Def [ {of|species|subindex} Species ] = Constant [ Phi = phi ]

<table>
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<tbody>
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</table>

Summary Constant value

4.1.373 Porosity Def: Deforming
Scope: Aria Material

Porosity Def: Deforming [ {of|species|subindex} Species ] = Deforming [ ]

<table>
<thead>
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<th>Value</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</table>

Summary Deforming model for porosity def
### 4.1.374 Porous Flow System: Single Phase

**Scope:** Aria Material


<table>
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<th>Value</th>
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<tbody>
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<tr>
<td>gz</td>
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### 4.1.375 Porous Flow System: Mixed Single Phase

**Scope:** Aria Material


<table>
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<tr>
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<td>undefined</td>
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<tr>
<td>gz</td>
<td>real</td>
<td>undefined</td>
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</table>

### 4.1.376 Porous Flow System: Two Phase Immiscible

**Scope:** Aria Material

Porous Flow System: Two Phase Immiscible [ of species subindex Species ] = Two_Phase_Immiscible [ Gx = gx | Gy = gy | Gz = gz ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
<td>gz</td>
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</table>

### 4.1.377 Porous Flow System: Air Water

**Scope:** Aria Material


<table>
<thead>
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<th>Value</th>
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<tr>
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</table>
4.1.378 Porous Flow System: Co2 Brine Salt
Scope: Aria Material

Porous Flow System: Co2 Brine Salt [ {of|species|subindex} Species ]= Co2_Brine_Salt [ X_Sat_Co2_File_Name = x_sat_co2_file_name | Rho_G_File_Name = rho_g_file_name | Rho_Co2_File_Name = rho_co2_file_name | Gx = gx | Gy = gy | Gz = gz ]

<table>
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</table>

4.1.379 Pp: Mesh Von Mises
Scope: Aria Material

Pp: Mesh Von Mises [ {of|species|subindex} Species ]= Mesh_Von_Mises [ ]

<table>
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<tr>
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Summary       Mesh Von Mises model

4.1.380 Pp: Solid Von Mises
Scope: Aria Material

Pp: Solid Von Mises [ {of|species|subindex} Species ]= Solid_Von_Mises [ ]

<table>
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Summary       Solid Von Mises model

4.1.381 Pp: Fluid Traction
Scope: Aria Material

Pp: Fluid Traction [ {of|species|subindex} Species ]= Fluid_Traction [ ]

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Summary       Fluid Traction model
4.1.382  Pp: Viscous Traction
Scope:  Aria Material

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Summary  Viscous Traction model - like Surface_Traction but omits the pressure contribution

4.1.383  Pp: Mesh Traction
Scope:  Aria Material

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Summary  Mesh Traction model

4.1.384  Pp: Solid Traction
Scope:  Aria Material

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Summary  Solid Traction model

4.1.385  Pp: Cvfem Yplus
Scope:  Aria Material

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Summary  CVFEM YPLUS model

4.1.386  Pp: Yplus
Scope:  Aria Material

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4.1.387 Pp: Turbulent Kinetic Energy Source
Scope: Aria Material

Pp: Turbulent Kinetic Energy Source $\text{[of|species|subindex} \text{Species}] = \text{Turbulent\_Kinetic\_Energy\_Source}$

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4.1.388 Pp: Surface Normal Shell
Scope: Aria Material

Pp: Surface Normal Shell $\text{[of|species|subindex} \text{Species}] = \text{Surface\_Normal\_Shell}$

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4.1.389 Prandtl Number
Scope: Aria Material

Prandtl Number $\text{[of|species|subindex} \text{Species}] = \text{Constant \[ Pr = Pr \]}$

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Summary  Constant value

4.1.390 Prandtl Number: General
Scope: Aria Material

Prandtl Number: General $\text{[of|species|subindex} \text{Species}] = \text{General}$

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</table>

Summary  General Prandtl number computed from material properties
4.1.391 Prandtl Number: Correlation
Scope: Aria Material

\[
\text{Prandtl Number: Correlation } [\text{ of species subindex } \text{ Species }] = \text{ Correlation} [\text{ Pr } = \text{ Pr } | \text{ Prw } = \text{ Prw } | \text{ Bulk Node Interface } = \text{ bulk node interface }]
\]

<table>
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Summary: Correlation Prandtl number computed from material properties

4.1.392 Prandtl Number: Correlation Wall
Scope: Aria Material

\[
\text{Prandtl Number: Correlation Wall } [\text{ of species subindex } \text{ Species }] = \text{ Correlation Wall} [\text{ Pr } = \text{ Pr } | \text{ Prw } = \text{ Prw } | \text{ Bulk Node Interface } = \text{ bulk node interface }]
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\text{Species}</td>
<td>\text{string}</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{Pr}</td>
<td>\text{real}</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{Prw}</td>
<td>\text{real}</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{bulk node interface}</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Correlation wall Prandtl number computed from material properties

4.1.393 Precursor Concentration
Scope: Aria Material

\[
\text{Precursor Concentration } [\text{ of species subindex } \text{ Species }] = \text{ Constant} [\text{ Value } = \text{ value }]
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\text{Species}</td>
<td>\text{string}</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{value}</td>
<td>\text{real}</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.394 Precursor Conservation
Scope: Aria Material

\[
\text{Precursor Conservation } [\text{ of species subindex } \text{ Species }] = \text{ Constant} [\text{ Value } = \text{ value }]
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\text{Species}</td>
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<td>undefined</td>
</tr>
<tr>
<td>\text{value}</td>
<td>\text{real}</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Summary
Constant value

### 4.1.395 Pressure

**Scope:** Aria Material

Pressure \[ \{ \text{of|species|subindex} \ Species \} = \text{Constant} \ [ \text{Value} = \text{value} ] \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Constant value

### 4.1.396 Pressure: Copy Phase All

**Scope:** Aria Material

Pressure: Copy Phase All \[ \{ \text{of|species|subindex} \ Species \} = \text{Copy_Phase_All} \ [ \] \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Copy the overall multi-phase pressure to the individual phase.

### 4.1.397 Pressure: From No Material Phase

**Scope:** Aria Material

Pressure: From No Material Phase \[ \{ \text{of|species|subindex} \ Species \} = \text{From_No_Material_Phase} \ [ \] \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Copy the overall pressure to the material phase (i.e. gas phase or solid phase).

### 4.1.398 Pressure: From Other Material Phase

**Scope:** Aria Material

Pressure: From Other Material Phase \[ \{ \text{of|species|subindex} \ Species \} = \text{From_Other_Material_Phase} \ [ \ Other_Phase = \text{other_phase} ] \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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</tr>
<tr>
<td>other_phase</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Copy the overall pressure from a different phase (i.e. gas phase or solid phase).
4.1.399 Pressure: Pressurization Model

Scope: Aria Material

- Pressure: Pressurization Model [ {of|species|subindex} Species ] = Pressurization_Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Pressure from a pressurization model

4.1.400 Pressure: Ideal Gas

Scope: Aria Material

- Pressure: Ideal Gas [ {of|species|subindex} Species ] = Ideal_Gas [ R = r ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>r</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Calculate the pressure based on the ideal gas law

4.1.401 Pressure: Equation Of State

Scope: Aria Material

- Pressure: Equation Of State [ {of|species|subindex} Species ] = Equation_Of_State [ Model_Name = model_name ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>model_name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Pressure calculated based on species, temperature, and equation of state for the specified pressurization model.

4.1.402 Radiation Form Factor

Scope: Aria Material

- Radiation Form Factor [ {of|species|subindex} Species ] = Constant [ F = f ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>f</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value
4.1.403  Radiation Form Factor: Calore User Sub
Scope: Aria Material

Radiation Form Factor: Calore User Sub [ {of|species|subindex} Species ]= Calore_User_Sub [ Name = name |Type = type |Multiplier = multiplier |Material_Data_Block = material_data_block |Data = data |Scaling_Field = scaling_field ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>type</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>multiplier</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>material_data_block</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>data</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>scaling_field</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Values from a Calore user subroutine

4.1.404  Radiative Conductivity
Scope: Aria Material

Radiative Conductivity [ {of|species|subindex} Species ]= Constant [ Krad = Krad ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
<td>Krad</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Constant value

4.1.405  Radiative Conductivity: Optically Thick
Scope: Aria Material

Radiative Conductivity: Optically Thick [ {of|species|subindex} Species ]= Optically_Thick [ Beta_R = beta_r |N = n ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>beta_r</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>n</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Radiation conductivity for optically thick materials

4.1.406  Radiative Conductivity: Volume Average
Scope: Aria Material

Radiative Conductivity: Volume Average [ {of|species|subindex} Species ]= Volume_Average [ ]
Parameter | Value | Default
---|---|---
Species | string | undefined

Summary: Radiation conductivity computed from normalized volume average of species radiation conductivities

4.1.407 Radiative Conductivity: Chemeq Foam
Scope: Aria Material

Radiative Conductivity: Chemeq Foam [ {of|species|subindex} Species ]= Chemeq_Foam [ Kc0 = Kc0 | Kc1 = Kc1 ]

Parameter | Value | Default
---|---|---
Species | string | undefined
Kc0 | real | undefined
Kc1 | real | undefined

Summary: Radiation conductivity for ChemEq foams, adjusting for the mass fraction of solids in the foam

4.1.408 Reaction Rate Multiplier
Scope: Aria Material

Reaction Rate Multiplier [ {of|species|subindex} Species ]= Constant [ Value = value ]

Parameter | Value | Default
---|---|---
Species | string | undefined
value | real | undefined

Summary: Constant value

4.1.409 Relative Permeability
Scope: Aria Material

Relative Permeability [ {of|species|subindex} Species ]= Constant [ K = k ]

Parameter | Value | Default
---|---|---
Species | string | undefined
k | real | undefined

Summary: Constant value

4.1.410 Relative Permeability: T Exponent
Scope: Aria Material

Relative Permeability: T Exponent [ {of|species|subindex} Species ]= T_Exponent [ K_Ref = k_ref | T_Ref = t_ref | N = n ]
Summary: Relative permeability as a power of normalized temperature

4.1.411 Relative Permeability: Volume Average
Scope: Aria Material

Relative Permeability: Volume Average [ \{of|species|subindex} Species ] = Volume_Average

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Relative permeability computed from volume average of species relative permeabilities

4.1.412 Relative Permeability: Mass Average
Scope: Aria Material

Relative Permeability: Mass Average [ \{of|species|subindex} Species ] = Mass_Average

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
</tbody>
</table>

Summary: Linear mixture-averaged mass diffusivity

4.1.413 Relative Permeability: Van Genuchten Gas
Scope: Aria Material

[ Liquid_Phase_Name = liquid_phase_name | Beta = beta | Residual_Liquid_Phase_Saturation = residual_liquid_phase_saturation ]
[ Residual_Gas_Phase_Saturation = residual_gas_phase_saturation ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
<td>liquid_phase_name</td>
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</tr>
<tr>
<td>beta</td>
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<td>undefined</td>
</tr>
<tr>
<td>residual_liquid_phase_saturation</td>
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<td>undefined</td>
</tr>
<tr>
<td>residual_gas_phase_saturation</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: A Van Genuchten gas model for relative permeability
4.1.414 Relative Permeability: Udell Cubic Gas

Scope: Aria Material

Relative Permeability: Udell Cubic Gas [ \{of\{species\{subindex\} Species \}= Udell_Cubic_Gas
[ Liquid_Phase_Name = liquid_phase_name |Residual_Liquid_Phase_Saturation = residual_liquid_phase_saturation
|Residual_Gas_Phase_Saturation = residual_gas_phase_saturation ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
<tr>
<td>Species</td>
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</tbody>
</table>

Summary  
A Udell cubic gas model for relative permeability

4.1.415 Relative Permeability: Van Genuchten Liquid

Scope: Aria Material

Relative Permeability: Van Genuchten Liquid [ \{of\{species\{subindex\} Species \}= Van_Genuchten_Liquid
[ Liquid_Phase_Name = liquid_phase_name |Beta = beta |Residual_Liquid_Phase_Saturation = residual_liquid_phase_saturation
|Residual_Gas_Phase_Saturation = residual_gas_phase_saturation ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
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</tr>
<tr>
<td>residual_gas_phase_saturation</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary  
A Van Genuchten liquid model for relative permeability

4.1.416 Relative Permeability: Udell Cubic Liquid

Scope: Aria Material

Relative Permeability: Udell Cubic Liquid [ \{of\{species\{subindex\} Species \}= Udell_Cubic_Liquid
[ Liquid_Phase_Name = liquid_phase_name |Residual_Liquid_Phase_Saturation = residual_liquid_phase_saturation
|Residual_Gas_Phase_Saturation = residual_gas_phase_saturation ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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<td>residual_liquid_phase_saturation</td>
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<tr>
<td>residual_gas_phase_saturation</td>
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</tbody>
</table>

Summary  
A Udell cubic liquid model for relative permeability

4.1.417 Relative Permeability: Von Mises Damage

Scope: Aria Material

Summary  
A Von Mises damage model for relative permeability
Relative Permeability: Von Mises Damage \[ \{ \text{of} \{ \text{species} \{ \text{subindex} \} \} \text{Species} \} = \text{Von\_Mises\_Damage} \]
\[ \text{Alpha} = \alpha \]
\[ \text{Residual\_Stress} = \text{residual\_stress} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>{ \text{of} { \text{species} { \text{subindex} } } \text{Species} }</td>
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<tr>
<td>\alpha</td>
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<td>undefined</td>
</tr>
<tr>
<td>\text{residual_stress}</td>
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</tr>
</tbody>
</table>

4.1.418 Reservoir Depth
Scope: Aria Material

Reservoir Depth \[ \{ \text{of} \{ \text{species} \{ \text{subindex} \} \} \text{Species} \} = \text{Constant} \]
\[ \text{H} = h \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>{ \text{of} { \text{species} { \text{subindex} } } \text{Species} }</td>
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</tr>
<tr>
<td>h</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Constant value

4.1.419 Retardation: Saturated Distributed
Scope: Aria Material

Retardation: Saturated Distributed \[ \{ \text{of} \{ \text{species} \{ \text{subindex} \} \} \text{Species} \} = \text{Saturated\_Distributed} \]

<table>
<thead>
<tr>
<th>Parameter</th>
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</tbody>
</table>

Summary
Distributed retardation factor for saturated porous flow. The corresponding distribution coefficient should be defined with a DISTRIBUTION COEFFICIENT model.

4.1.420 Retardation: Gas Phase Distributed
Scope: Aria Material

Retardation: Gas Phase Distributed \[ \{ \text{of} \{ \text{species} \{ \text{subindex} \} \} \text{Species} \} = \text{Gas\_Phase\_Distributed} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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<tr>
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</tbody>
</table>

Summary
Distributed retardation factor for gas phase in porous flow. The corresponding distribution coefficient should be defined with a DISTRIBUTION COEFFICIENT model.

4.1.421 Retardation: Liquid Phase Distributed
Scope: Aria Material
### 4.1.422 Retardation: Wetting Phase Distributed

**Scope:** Aria Material

Summary: Distributed retardation factor for wetting phase porous flow. The corresponding distribution coefficient should be defined with a DISTRIBUTION COEFFICIENT model.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tbody>
</table>

### 4.1.423 Retardation: Nonwetting Phase Distributed

**Scope:** Aria Material

Summary: Distributed retardation factor for non-wetting phase in porous flow. The corresponding distribution coefficient should be defined with a DISTRIBUTION COEFFICIENT model.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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</thead>
<tbody>
<tr>
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</tr>
</tbody>
</table>

### 4.1.424 Retardation

**Scope:** Aria Material

Summary: Constant value

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>r</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

### 4.1.425 Reynolds Number

**Scope:** Aria Material
Reynolds Number [ \{ \text{of|species|subindex}\} \text{Species} ] = \text{Constant} [ \text{Re} = \text{Re} ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
<td>\text{Re}</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.426 Reynolds Number: General
Scope: Aria Material

Reynolds Number: General [ \{ \text{of|species|subindex}\} \text{Species} ] = General [ \text{L} = \text{l} ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
<td>\text{L}</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Reynolds number computed from a specified length scale

4.1.427 Reynolds Number: Correlation
Scope: Aria Material

Reynolds Number: Correlation [ \{ \text{of|species|subindex}\} \text{Species} ] = Correlation [ \text{Re} = \text{Re} | \text{Bulk\_Node\_Interface} = \text{bulk\_node\_interface} ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>\text{Re}</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{bulk_node_interface}</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Correlation Reynolds number computed from material properties

4.1.428 Saturated Retardation
Scope: Aria Material

Saturated Retardation [ \{ \text{of|species|subindex}\} \text{Species} ] = \text{Constant} [ \text{R} = \text{r} ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>\text{r}</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.429 Saturation
Scope: Aria Material
Saturation \[ \{ \text{of|species|subindex} \} \text{ Species } \] = Constant \[ S = s \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>s</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.430 Saturation: Van Genuchten
Scope: Aria Material

Saturation: Van Genuchten \[ \{ \text{of|species|subindex} \} \text{ Species } \] = Van Genuchten \[ \text{ Is\_Wetting\_Phase } = \text{is\_wetting\_phase} \text{ Nonwetting\_Phase } = \text{nonwetting\_phase} \text{ Reference\_Capillary\_Pressure } = \text{reference\_capillary\_pressure} \text{ Beta } = \text{beta} \text{ Residual\_Wetting\_Phase\_Saturation } = \text{residual\_wetting\_phase\_saturation} \text{ Residual\_Nonwetting\_Phase\_Saturation } = \text{residual\_nonwetting\_phase\_saturation} \text{ Wetting\_Phase } = \text{wetting\_phase} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>is_wetting_phase</td>
<td>integer</td>
<td>undefined</td>
</tr>
<tr>
<td>nonwetting_phase</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>reference_capillary_pressure</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>beta</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>residual_wetting_phase_saturation</td>
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<td>undefined</td>
</tr>
<tr>
<td>residual_nonwetting_phase_saturation</td>
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<td>undefined</td>
</tr>
<tr>
<td>wetting_phase</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: A Van Genuchten model for saturation using capillary pressure

4.1.431 Saturation: From Other Phase
Scope: Aria Material

Saturation: From Other Phase \[ \{ \text{of|species|subindex} \} \text{ Species } \] = From Other Phase [

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Get the saturation from one minus the other phase saturation

4.1.432 Scaling
Scope: Aria Material

Scaling \[ \{ \text{of|species|subindex} \} \text{ Species } \] = Constant \[ \text{ Value } = \text{value} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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<tr>
<td>value</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value
4.1.433 Scattering Coefficient
Scope: Aria Material

Scattering Coefficient \[ \text{Species} \]= Constant \( \text{Value} = \text{value} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.434 Scattering Cross Section
Scope: Aria Material

Scattering Cross Section \[ \text{Species} \]= Constant \( \text{Scat} = \text{scat} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
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<tr>
<td>scat</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.435 Scattering Cross Section: Calore User Sub
Scope: Aria Material

Scattering Cross Section: Calore User Sub \[ \text{Species} \]= Calore_User_Sub \[ \text{Name} = \text{name} | \text{Type} = \text{type} | \text{Multiplier} = \text{multiplier} | \text{Material_Data_Block} = \text{material_data_block} | \text{Data} = \text{data} | \text{Scaling_Field} = \text{scaling_field} \]

Summary: Calore user subroutine scattering cross section

4.1.436 Schmidt Number
Scope: Aria Material

Schmidt Number \[ \text{Species} \]= Constant \( \text{Sc} = \text{Sc} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Sc</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>
4.1.437 Seebeck Coefficient
Scope: Aria Material

\[ \text{Seebeck Coefficient} = \text{Constant} \]

<table>
<thead>
<tr>
<th>Parameter ( \text{Species} )</th>
<th>Value ( a )</th>
<th>Default undefined</th>
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</thead>
</table>

Summary Constant value

4.1.438 Shell Lofting Factor
Scope: Aria Material

\[ \text{Shell Lofting Factor} = \text{Constant} \]

<table>
<thead>
<tr>
<th>Parameter ( \text{Species} )</th>
<th>Value ( f )</th>
<th>Default undefined</th>
</tr>
</thead>
</table>

Summary Constant value

4.1.439 Shell Lofting Factor: Element Attribute
Scope: Aria Material

\[ \text{Shell Lofting Factor: Element Attribute} = \text{Element Attribute} \]

<table>
<thead>
<tr>
<th>Parameter ( \text{Species} )</th>
<th>Value ( \text{Multiplier} )</th>
<th>Default undefined</th>
</tr>
</thead>
</table>

Summary Element Distribution Factor for the lofting of shell surface

4.1.440 Shell Thickness
Scope: Aria Material

\[ \text{Shell Thickness} = \text{Constant} \]

<table>
<thead>
<tr>
<th>Parameter ( \text{Species} )</th>
<th>Value ( t )</th>
<th>Default undefined</th>
</tr>
</thead>
</table>

Summary Constant value
4.1.441  Shell Thickness: Element Attribute
Scope: Aria Material

Shell Thickness:  Element Attribute [ {of|species|subindex} Species ]= Element_Attribute [ Multiplier = multiplier |Attribute_Name = attribute_name |Contact = contact ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>attribute_name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>contact</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Distribution Factor for the shell thickness

4.1.442  Skeleton Density
Scope: Aria Material

Skeleton Density [ {of|species|subindex} Species ]= Constant [ Rho = rho ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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<tbody>
<tr>
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<td>undefined</td>
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</table>

Summary  Constant value

4.1.443  Skeleton Enthalpy: Cpt
Scope: Aria Material

Skeleton Enthalpy:  Cpt [ {of|species|subindex} Species ]= Cpt [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tbody>
</table>

Summary  CPT model for the the solid skeleton enthalpy

4.1.444  Skeleton Internal Energy
Scope: Aria Material

Skeleton Internal Energy [ {of|species|subindex} Species ]= Constant [ E = e ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>e</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Constant value
4.1.445  Skeleton Internal Energy: Cpt
Scope:  Aria Material

Skeleton Internal Energy: Cpt [ {of|species|subindex} Species ]= Cpt [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
</tbody>
</table>

Summary  CPT model for the the solid skeleton internal energy

4.1.446  Skeleton Internal Energy: Linear
Scope:  Aria Material

Skeleton Internal Energy: Linear [ {of|species|subindex} Species ]= Linear [ Cp = cp | T_Ref = T_ref ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>cp</td>
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<td>undefined</td>
</tr>
<tr>
<td>T_ref</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  CPT model for the the solid skeleton internal energy

4.1.447  Skeleton Specific Heat
Scope:  Aria Material

Skeleton Specific Heat [ {of|species|subindex} Species ]= Constant [ Cp = cp ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>cp</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Constant value

4.1.448  Solid Body Acceleration
Scope:  Aria Material

Solid Body Acceleration [ {of|species|subindex} Species ]= Constant [ Valuex = valuex | Valuey = valuey | Valuez = valuez ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>valuex</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>valuey</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>valuez</td>
<td>real</td>
<td>undefined</td>
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</tbody>
</table>

Summary  Constant value
4.1.449  Solid Density
Scope:  Aria Material

Solid Density [ {of|species|subindex} Species ] = Constant [ Rho_Solid = rho_solid ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>rho_solid</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Constant value

4.1.450  Solid Lambda
Scope:  Aria Material

Solid Lambda [ {of|species|subindex} Species ] = Constant [ Lambda = lambda ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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<tr>
<td>lambda</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Constant value

4.1.451  Solid Lambda: Converted
Scope:  Aria Material

Solid Lambda: Converted [ {of|species|subindex} Species ] = Converted [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
</tbody>
</table>

Summary  Lame coefficient computed from Poisson’s ratio and elastic modulus

4.1.452  Solid Lambda: Converted Plane Stress
Scope:  Aria Material

Solid Lambda: Converted Plane Stress [ {of|species|subindex} Species ] = Converted_Plane_Stress [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Lame coefficient applicable to plane stress conditions computed from Poisson’s ratio and elastic modulus

4.1.453  Solid Poissons Ratio
Scope:  Aria Material
Solid Poissons Ratio \( \{ \text{of|species|subindex} \} \text{Species} \) = Constant \( \text{Nu} = \nu \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>( \nu )</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Constant value

4.1.454 Solid Pressure Gradient: Gidaspow
Scope: Aria Material

Solid Pressure Gradient: Gidaspow \( \{ \text{of|species|subindex} \} \text{Species} \) = Gidaspow \( \text{GO} = \text{g}0 \) \( \text{C} = \text{c} \) \( \text{F}_{\text{max}} = \text{f}_{\text{max}} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>( \text{g}0 )</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>( \text{c} )</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>( \text{f}_{\text{max}} )</td>
<td>real</td>
<td>undefined</td>
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</tbody>
</table>

4.1.455 Solid Stress: Isothermal
Scope: Aria Material

Solid Stress: Isothermal \( \{ \text{of|species|subindex} \} \text{Species} \) = Isothermal \( \text{T} = \t \) \( \text{T}_{\text{Ref}} = \t_{\text{ref}} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>( \t )</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>( \t_{\text{ref}} )</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.456 Solid Stress: Linear Elastic
Scope: Aria Material

Solid Stress: Linear Elastic \( \{ \text{of|species|subindex} \} \text{Species} \) = Linear_Elastic \( \text{Reference\_Frame} = \text{reference\_frame} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
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<tbody>
<tr>
<td>Species</td>
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<tr>
<td>reference_frame</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.457 Solid Stress: Porous Effective
Scope: Aria Material

Solid Stress: Porous Effective \( \{ \text{of|species|subindex} \} \text{Species} \) = Porous_Effective \( \text{Pressure} = \text{pressure} \) \( \text{Phase} = \text{phase} \) \( \text{Biot} = \text{biot} \)
### 4.1.458 Solid Stress: Saturation Weighted Porous Effective

#### Scope: Aria Material

Solid Stress: Saturation Weighted Porous Effective \( \{\text{of species subindex} \} \text{Species} \) = Saturation_Weighted_Porous_Effective \( \{ \text{Pressure} = \text{pressure} \mid \text{Phase} = \text{phase} \mid \text{Biot} = \text{biot} \} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
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<td>pressure</td>
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<td>undefined</td>
</tr>
<tr>
<td>phase</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
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<tr>
<td>biot</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Apply the pressure as an isotropic mesh stress. Primarily used as an effective stress in porous flow problems.

### 4.1.459 Solid Stress: Mooney Rivlin

#### Scope: Aria Material

Solid Stress: Mooney Rivlin \( \{\text{of species subindex} \} \text{Species} \) = Mooney_Rivlin \( \)  

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
</tbody>
</table>

### 4.1.460 Solid Stress: Neochookean Elastic

#### Scope: Aria Material

Solid Stress: Neochookean Elastic \( \{\text{of species subindex} \} \text{Species} \) = Neochookean_Elastic \( \) 

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
</tbody>
</table>

### 4.1.461 Solid Stress: J

#### Scope: Aria Material

Solid Stress: J \( \{\text{of species subindex} \} \text{Species} \) = J \( \{ \text{H = h} \mid \text{Yield Stress} = \text{yield stress} \) 

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4.1.462  Solid Stress: Hyperelastic Damage

Scope:  Aria Material

\[
\text{Solid Stress: Hyperelastic Damage [ } \{ \text{of|species|subindex}\} \text{ Species } = \text{Hyperelastic_Damage} \\
\text{ [ Max\_Damage = max\_damage ] }
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
<td>h</td>
<td>&quot;string&quot;</td>
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<tr>
<td>yield_stress</td>
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<td>undefined</td>
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</table>

4.1.463  Solid Stress: Incompressible Newtonian

Scope:  Aria Material

\[
\text{Solid Stress: Incompressible Newtonian [ } \{ \text{of|species|subindex}\} \text{ Species } = \text{Incompressible_Newtonian} \\
\text{ [ Pressure = pressure | Phase = phase | Biot = biot | Reference\_Frame = reference\_frame ] }
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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</tr>
<tr>
<td>pressure</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>phase</td>
<td>&quot;string&quot;</td>
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<td>biot</td>
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<td>undefined</td>
</tr>
<tr>
<td>reference_frame</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
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</tbody>
</table>

4.1.464  Solid Stress: Nonlinear Elastic

Scope:  Aria Material

\[
\text{Solid Stress: Nonlinear Elastic [ } \{ \text{of|species|subindex}\} \text{ Species } = \text{Nonlinear\_Elastic} [ ]
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
</tbody>
</table>

4.1.465  Solid Stress: Residual

Scope:  Aria Material

\[
\text{Solid Stress: Residual [ } \{ \text{of|species|subindex}\} \text{ Species } = \text{Residual } [ \text{ Sz = sz | Szz = szz | Sxz = szz | Syz = syz | Sy = sy | Syy = syy | Sxy = sxy | Sx = sx | Sxx = szz } ]
\]

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### 4.1.466 Solid Stress: Thermal

**Scope:** Aria Material

<table>
<thead>
<tr>
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<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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<td>undefined</td>
</tr>
</tbody>
</table>

Solid Stress: Thermal \{of|species|subindex\} Species = Thermal []

### 4.1.467 Solid Stress: Species

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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<td>undefined</td>
</tr>
<tr>
<td>sref</td>
<td>real undefined</td>
<td>undefined</td>
</tr>
<tr>
<td>beta</td>
<td>real undefined</td>
<td>undefined</td>
</tr>
<tr>
<td>phase</td>
<td>string undefined</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Solid Stress: Species \{of|species|subindex\} Species = Species [ Sref = sref | Beta = beta | Phase = phase ]

**Summary**
A linear mechanical strain due to a species concentration

### 4.1.468 Solid Stress: Species Transversely Isotropic

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string undefined</td>
<td>undefined</td>
</tr>
<tr>
<td>refval</td>
<td>real undefined</td>
<td>undefined</td>
</tr>
<tr>
<td>beta_n</td>
<td>real undefined</td>
<td>undefined</td>
</tr>
<tr>
<td>beta_t</td>
<td>real undefined</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Solid Stress: Species Transversely Isotropic \{of|species|subindex\} Species = Species_Transversely_Isotropic [ Refval = refval | Beta_N = beta_n | Beta_T = beta_t | N_X = n_x | N_Y = n_y | N_Z = n_z | Phase = phase ]
Summary  A linear mechanical strain that is transversely isotropic due to a species concentration

4.1.469  Solid Stress: Lame
Scope:  Aria Material

Solid Stress: Lame [ {of|species|subindex} Species ]= Lame [ Model = model ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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</tr>
<tr>
<td>model</td>
<td>&quot;string&quot;</td>
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</tbody>
</table>

4.1.470  Solid Two Mu
Scope:  Aria Material

Solid Two Mu [ {of|species|subindex} Species ]= Constant [ Two_Mu = two.mu ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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<tr>
<td>two.mu</td>
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<td>undefined</td>
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</table>

Summary  Constant value

4.1.471  Solid Two Mu: Converted
Scope:  Aria Material

Solid Two Mu: Converted [ {of|species|subindex} Species ]= Converted [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Lame coefficient computed from Poisson’s ratio and elastic modulus

4.1.472  Solid Youngs Modulus
Scope:  Aria Material

Solid Youngs Modulus [ {of|species|subindex} Species ]= Constant [ E = e ]
Parameter | Value | Default
--- | --- | ---
Species | string | undefined
\( e \) | real | undefined

Summary | Constant value

4.1.473 Soret Coefficient
Scope: Aria Material

Soret Coefficient \([ \text{of} \{\text{species}\} \text{Species} \] = Constant \([ S = s ]\)

Parameter | Value | Default
--- | --- | ---
Species | string | undefined
\( s \) | real | undefined

Summary | Constant value

4.1.474 Sound Speed
Scope: Aria Material

Sound Speed \([ \text{of} \{\text{species}\} \text{Species} \] = Constant \([ C = c ]\)

Parameter | Value | Default
--- | --- | ---
Species | string | undefined
\( c \) | real | undefined

4.1.475 Species
Scope: Aria Material

Species \([ \text{of} \{\text{species}\} \text{Species} \] = Constant \([ \text{Value} = \text{value} ]\)

Parameter | Value | Default
--- | --- | ---
Species | string | undefined
\( \text{value} \) | real | undefined

Summary | Constant value

4.1.476 Species: Partial Molar Volume
Scope: Aria Material

Species: Partial Molar Volume \([ \text{of} \{\text{species}\} \text{Species} \] = Partial_Molar_Volume \([ ]\)

Parameter | Value | Default
--- | --- | ---
Species | string | undefined
### 4.1.477 Species: Sum All Species

**Scope:** Aria Material

Species: Sum All Species

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

### 4.1.478 Species: Solvent Concentration From Solute Concentrations And Partial Molar Volumes

**Scope:** Aria Material

Species: Solvent Concentration From Solute Concentrations And Partial Molar Volumes

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
</tbody>
</table>

### 4.1.479 Species: Chemeq Gas

**Scope:** Aria Material

Species: Chemeq Gas

<table>
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<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

### 4.1.480 Species: From Chemeq

**Scope:** Aria Material

Species: From Chemeq

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
</tbody>
</table>

**Summary**
Species concentration of one species extracted from ChemEq

### 4.1.481 Species: From Mass Fraction

**Scope:** Aria Material

Species: From Mass Fraction

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Molar concentration of a single species calculated from its mass fraction, molecular weight, and overall density
### 4.1.482 Species Diffusion: Ficks Law

**Scope:** Aria Material

Species Diffusion: Ficks Law [ \{of\mid species\mid subindex\} Species ]= Ficks_Law [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

### 4.1.483 Species Diffusion: Nernst Planck

**Scope:** Aria Material

Species Diffusion: Nernst Planck [ \{of\mid species\mid subindex\} Species ]= Nernst_Planck [ F = f | R = r ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>f</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>r</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

### 4.1.484 Species Diffusion: All Subindices Use Stefan Maxwell

**Scope:** Aria Material

Species Diffusion: All Subindices Use Stefan Maxwell [ \{of\mid species\mid subindex\} Species ]= All_Subindices_Use_Stefan_Maxwell [ VelocityConversion = velocityConversion | Canteraxmlfile = canteraXMLFile | Velocitybasis = velocityBasis ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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<td>velocityBasis</td>
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</table>

### 4.1.485 Species Diffusion: Basic

**Scope:** Aria Material

Species Diffusion: Basic [ \{of\mid species\mid subindex\} Species ]= Basic [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

### 4.1.486 Species Diffusion: Constant Tensorial

**Scope:** Aria Material

Species Diffusion: Constant Tensorial [ \{of\mid species\mid subindex\} Species ]= Constant_Tensorial [ D_{Xx} = d_{xx} | D_{Xy} = d_{xy} | D_{Xz} = d_{xz} | D_{Yx} = d_{yx} | D_{Yy} = d_{yy} | D_{Yz} = d_{yz} | D_{Zx} = d_{zx} | D_{Zy} = d_{zy} | D_{Zz} = d_{zz} ]
<table>
<thead>
<tr>
<th>Parameter</th>
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<td>d_xz</td>
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</tr>
<tr>
<td>d_zz</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.487  Species Diffusion: Tensorial
Scope:  Aria Material

Species Diffusion: Tensorial [ {of|species|subindex} Species ]= Tensorial [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
</tbody>
</table>

4.1.488  Species Diffusion: Tensorial Dispersive
Scope:  Aria Material

Species Diffusion: Tensorial Dispersive [ {of|species|subindex} Species ]= Tensorial_Dispersive [ Alpha_T = alpha_t |Alpha_L = alpha_l |Tau = tau ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>alpha_t</td>
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<td>alpha_l</td>
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<td>undefined</td>
</tr>
<tr>
<td>tau</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.489  Species Diffusion: Thermophoresis
Scope:  Aria Material

Species Diffusion: Thermophoresis [ {of|species|subindex} Species ]= Thermophoresis [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
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<th>Default</th>
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</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.490  Species Diffusion: Tensor Thermophoresis
Scope:  Aria Material

Species Diffusion: Tensor Thermophoresis [ {of|species|subindex} Species ]= Tensor_Thermophoresis [ ]
4.1.491  Species Diffusion: Electromigration
Scope: Aria Material

\[ \text{Species Diffusion: Electromigration } \{\text{of|species|subindex}} \text{ Species } \} = \text{Electromigration} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
<tr>
<td>Species</td>
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</tbody>
</table>

4.1.492  Species Diffusion: Tensor Electromigration
Scope: Aria Material

\[ \text{Species Diffusion: Tensor Electromigration } \{\text{of|species|subindex}} \text{ Species } \} = \text{Tensor_Electromigration} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.493  Species Diffusion: Darcy
Scope: Aria Material

\[ \text{Species Diffusion: Darcy } \{\text{of|species|subindex}} \text{ Species } \} = \text{Darcy} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.494  Species Diffusion: Chemical Potential
Scope: Aria Material

\[ \text{Species Diffusion: Chemical Potential } \{\text{of|species|subindex}} \text{ Species } \} = \text{Chemical_Potential} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  
Model species diffusion using a flux determined from the gradient of the chemical potential. \[ \vec{J} = -M c \nabla \mu \] where M is the mobility, c the species concentration, and \( \mu \) the chemical potential.

4.1.495  Species Diffusion: Tensor Chemical Potential
Scope: Aria Material

\[ \text{Species Diffusion: Tensor Chemical Potential } \{\text{of|species|subindex}} \text{ Species } \} = \text{Tensor_Chemical_Potential} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

186
Model species diffusion using a flux determined from the gradient of the chemical potential. \( \vec{J} = -M\dot{c}\nabla \mu \) where M is the tensor mobility, c the species concentration, and \( \mu \) the chemical potential.

**4.1.496 Species Diffusion: Mass Balance Fracbal**

**Scope:** Aria Material

Summary: Species diffusion for the fracbal species in a mass balance equation system. \( \vec{J}_i = -1 \sum_{j \neq i} (\dot{m}_j)/MW_i \).

**4.1.497 Species Diffusivity**

**Scope:** Aria Material

Summary: Constant value

**4.1.498 Species Diffusivity: Arrhenius**

**Scope:** Aria Material

Summary: Arrhenius species diffusivity
### 4.1.499 Species Diffusivity: Dissolution

**Scope:** Aria Material

Species Diffusivity: Dissolution  

\[ \text{Species} \text{ Diffusivity: Dissolution} \left[ \{\text{of/species/subindex}\} \text{ Species } \right] = \text{Dissolution} \left[ D = d \right] \]

<table>
<thead>
<tr>
<th>Parameter ( \text{Species} )</th>
<th>Value ( d )</th>
<th>Default ( )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \text{string} )</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  
Dissolution species diffusivity for a species disolving in a liquid from a gas.

### 4.1.500 Species Enthalpy

**Scope:** Aria Material

Species Enthalpy  

\[ \text{Species} \text{ Enthalpy} \left[ \{\text{of/species/subindex}\} \text{ Species } \right] = \text{Constant} \left[ H = h \right] \]

<table>
<thead>
<tr>
<th>Parameter ( \text{Species} )</th>
<th>Value ( h )</th>
<th>Default ( )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \text{string} )</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  
Constant value

### 4.1.501 Species Enthalpy: Cantera

**Scope:** Aria Material

Species Enthalpy: Cantera  

\[ \text{Species} \text{ Enthalpy: Cantera} \left[ \{\text{of/species/subindex}\} \text{ Species } \right] = \text{Cantera} \left[ \right] \]

<table>
<thead>
<tr>
<th>Parameter ( \text{Species} )</th>
<th>Value ( )</th>
<th>Default ( )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \text{string} )</td>
<td></td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  
Cantera species enthalpy

### 4.1.502 Species Face Stabilization Scaling: Default

**Scope:** Aria Material

**Summary**  
Default face stabilization scaling for energy equation

### 4.1.503 Species Fraction

**Scope:** Aria Material

Species Fraction  

\[ \text{Species} \text{ Fraction} \left[ \{\text{of/species/subindex}\} \text{ Species } \right] = \text{Constant} \left[ \text{Value} = \text{value} \right] \]
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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<td>undefined</td>
</tr>
<tr>
<td></td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

**4.1.504 Species Fraction: From Chemeq**

**Scope:** Aria Material

Species Fraction: From Chemeq [ {of|species|subindex} Species ] = From_Chemeq [ ]

Summary: Species fraction of one species extracted from ChemEq

**4.1.505 Species Mobility**

**Scope:** Aria Material

Species Mobility [ {of|species|subindex} Species ] = Constant [ Value = value ]

Summary: Constant value

**4.1.506 Species Mobility: Nernst Einstein**

**Scope:** Aria Material

Species Mobility: Nernst Einstein [ {of|species|subindex} Species ] = Nernst_Einstein [ R = r [Temperature_Material_Phase = temperature_material_phase ]

Summary: Mobility from diffusivity, $M = \frac{D}{RT}$.

**4.1.507 Species Surface**

**Scope:** Aria Material

Species Surface [ {of|species|subindex} Species ] = Constant [ Value = value ]
Summary    Constant value

4.1.508  Species Surface: Equivalent To Bulk
Scope:  Aria Material

Species Surface: Equivalent To Bulk [ {of|species|subindex} Species ] = Equivalent_To_Bulk

Summary    Approximate a surface species concentration as identical to the bulk concentration

4.1.509  Species Surface: Diffusion To Surf Correction
Scope:  Aria Material

Species Surface: Diffusion To Surf Correction [ {of|species|subindex} Species ] = Diffusion_To_Surf_Correction[
Species_A = species_a |Species_B = species_b |D = d |R = r |F = f |K = k |Eqpot = eqPot |Dir = dir ]

Summary    Correction for surface concentration based on balancing reaction rate and diffusion from pore center to wall.

4.1.510  Species Valence
Scope:  Aria Material

Species Valence [ {of|species|subindex} Species ] = Constant [ Z = z ]

Summary    Constant value

190
4.1.511 Specific Dissipation Rate
Scope: Aria Material

Specific Dissipation Rate [ \{of\|species\|subindex\} Species ] = Constant [ Value = value ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Constant value

4.1.512 Specific Dissipation Rate Density
Scope: Aria Material

Specific Dissipation Rate Density [ \{of\|species\|subindex\} Species ] = Constant [ Value = value ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Constant value

4.1.513 Specific Dissipation Rate Diffusive Flux: Basic
Scope: Aria Material

Specific Dissipation Rate Diffusive Flux: Basic [ \{of\|species\|subindex\} Species ] = Basic [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Basic model of specific dissipation rate diffusive flux

4.1.514 Specific Heat: Calore User Sub
Scope: Aria Material

Specific Heat: Calore User Sub [ \{of\|species\|subindex\} Species ] = Calore_User_Sub [ Name = name | Type = type | Multiplier = multiplier | Material_Data_Block = material_data_block | Data = data | Scaling_Field = scaling_field ]
### 4.1.515 Specific Heat: Curing Foam

**Scope:** Aria Material

**Summary:** Values from a Calore user subroutine

#### Specific Heat: Curing Foam

Specific Heat: Curing Foam [ {of|species|subindex} Species ] = Curing_Foam [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** Specific heat for a curing foam

### 4.1.516 Specific Heat: Phase Average

**Scope:** Aria Material

**Summary:** Phase-averaged values

#### Specific Heat: Phase Average

Specific Heat: Phase Average [ {of|species|subindex} Species ] = Phase_Average [ Subindex_A = subindex_a | Subindex_B = subindex_b ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>subindex_a</td>
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<td>undefined</td>
</tr>
<tr>
<td>subindex_b</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

### 4.1.517 Specific Heat: Use Phase Change

**Scope:** Aria Material

**Summary:**

#### Specific Heat: Use Phase Change

Specific Heat: Use Phase Change [ {of|species|subindex} Species ] = Use_Phase_Change [ Flh = flh | Ts = ts | Tl = tl | Type = type | Field = field ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>flh</td>
<td>real</td>
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</tr>
<tr>
<td>field</td>
<td>&quot;string&quot;</td>
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</tr>
</tbody>
</table>
Summary: Specific heat with a step function accounting for phase change.

**Known Issue:** Using this model can cause poor convergence. Use the 'Melting' source for energy for much better numerical performance.

### 4.1.518 Specific Heat: Interpolated Phase Average

**Scope:** Aria Material

Specific Heat: Interpolated Phase Average \( \{ \text{species|subindex} \} \) \( \text{Species} \) = Interpolated_Phase_Average \( \{ \text{subindex}_A = \text{subindex}_a \ | \text{subindex}_B = \text{subindex}_b \} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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<tbody>
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<tr>
<td>subindex_b</td>
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</table>

Summary: Phase-averaged values

### 4.1.519 Specific Heat: Porous Phase Specific Average

**Scope:** Aria Material

Specific Heat: Porous Phase Specific Average \( \{ \text{species|subindex} \} \) \( \text{Species} \) = Porous_Phase_Specific_Average \( \{ \text{Liquid_Phase_Name} = \text{Liquid_Phase_Name} \ | \text{Gas_Phase_Name} = \text{Gas_Phase_Name} \} \)

<table>
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<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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<td>Gas_Phase_Name</td>
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</tr>
</tbody>
</table>

Summary: Specific heat for a multiphase material of two or three phases

### 4.1.520 Specific Heat: Mass Average

**Scope:** Aria Material

Specific Heat: Mass Average \( \{ \text{species|subindex} \} \) \( \text{Species} \) = Mass_Average \[

<table>
<thead>
<tr>
<th>Parameter</th>
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<tbody>
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<td>Species</td>
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</tr>
</tbody>
</table>

Summary: Specific heat computed from mass average of species specific heats

### 4.1.521 Specific Heat: Cantera

**Scope:** Aria Material

Specific Heat: Cantera \( \{ \text{species|subindex} \} \) \( \text{Species} \) = Cantera \[

193
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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</tr>
<tr>
<td>$h_0$ real</td>
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<td></td>
</tr>
</tbody>
</table>

**Summary**  
Cantera specific heat for low Mach number applications

### 4.1.522 Specific Heat

**Scope:** Aria Material

Specific Heat \[ \{ \text{of \ species \ subindex} \ \text{Species} \} = \text{Constant} [ \ C_p = C_p \ | \ H_0 = h_0 ] \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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<tbody>
<tr>
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<tr>
<td>$C_p$ real</td>
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<tr>
<td>$h_0$ real</td>
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</tr>
</tbody>
</table>

**Summary**  
Constant specific heat from an evaluator

### 4.1.523 Specific Heat: T Exponent

**Scope:** Aria Material

Specific Heat: T Exponent \[ \{ \text{of \ species \ subindex} \ \text{Species} \} = \text{T_Exponent} [ \ C_p_{\text{ref}} = C_p_{\text{ref}} \ | \ T_{\text{ref}} = T_{\text{ref}} \ | \ N = n \ | \ H_0 = h_0 ] \]

<table>
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<tr>
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<th>Value</th>
<th>Default</th>
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<tbody>
<tr>
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<td>$T_{\text{ref}}$ real</td>
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<td>$h_0$ real</td>
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**Summary**  
Temperature exponent specific heat from an evaluator

### 4.1.524 Specific Heat Cp

**Scope:** Aria Material

Specific Heat Cp \[ \{ \text{of \ species \ subindex} \ \text{Species} \} = \text{Constant} [ \ \text{Value} = \text{value} ] \]

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**Summary**  
Constant value

### 4.1.525 Specific Heat Cp: Cantera

**Scope:** Aria Material
Specific Heat $C_p$: Cantera [ of \{species \} subindex $Species$ ] = Cantera [ ]

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<td>Species</td>
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Summary: Cantera specific heat for low Mach number applications

### 4.1.526 Specific Surface Area

**Scope:** Aria Material

Specific Surface Area [ of \{species \} $Species$ ] = Constant [ Value = value ]

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<th>Value</th>
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<tbody>
<tr>
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</table>

Summary: Constant value

### 4.1.527 Supg Tau: Shakib Energy

**Scope:** Aria Material

Supg Tau: Shakib Energy = Shakib_Energy [ $H = h$ | Use_Advection = use_advection | Use_Diffusion = use_diffusion | Use_Time = use_time | Multiplier = multiplier ]

<table>
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</table>

Summary: A Shakib SUPG Tau model

### 4.1.528 Supg Tau: Shakib Enthalpy

**Scope:** Aria Material

Supg Tau: Shakib Enthalpy = Shakib_Enthalpy [ Multiplier = multiplier | Use_Diffusion = use_diffusion | Use_Time = use_time ]

<table>
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Summary: A Shakib SUPG Tau model
4.1.529  **Supg Tau: Shakib Momentum**  
Scope:  Aria Material

\[
\text{Supg Tau: Shakib Momentum} = \text{Shakib\_Momentum} [ \ H = \ h \mid \text{Use\_Advection} = \ use\_advection \mid \text{Use\_Diffusion} = \ use\_diffusion \mid \text{Use\_Time} = \ use\_time \mid \text{Multiplier} = \ multiplier ]
\]

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Summary  A Shakib SUPG Tau model

4.1.530  **Supg Tau: Shakib Level Set**  
Scope:  Aria Material

\[
\text{Supg Tau: Shakib Level Set} = \text{Shakib\_Level\_Set} [ \ H = \ h \mid \text{Use\_Advection} = \ use\_advection \mid \text{Use\_Diffusion} = \ use\_diffusion \mid \text{Use\_Time} = \ use\_time \mid \text{Multiplier} = \ multiplier ]
\]

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Summary  A Shakib SUPG Tau model

4.1.531  **Supg Tau: Shakib Suspension**  
Scope:  Aria Material

\[
\text{Supg Tau: Shakib Suspension} = \text{Shakib\_Suspension} [ \ H = \ h \mid \text{Use\_Advection} = \ use\_advection \mid \text{Use\_Diffusion} = \ use\_diffusion \mid \text{Use\_Time} = \ use\_time \mid \text{Multiplier} = \ multiplier ]
\]

<table>
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Summary  A Shakib SUPG Tau model for the suspension equation

4.1.532  **Supg Tau: Classic Energy**  
Scope:  Aria Material
Supg Tau: Classic Energy = Classic_Energy [ Epsilon = epsilon |Multiplier = multiplier ]

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<tr>
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</table>

Summary A Shakib SUPG Tau model

4.1.533 Supg Tau: Classic Momentum
Scope: Aria Material

Supg Tau: Classic Momentum = Classic_Momentum [ Epsilon = epsilon |Multiplier = multiplier ]

<table>
<thead>
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<th>Parameter</th>
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<th>Default</th>
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<tbody>
<tr>
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</tr>
<tr>
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Summary A Shakib SUPG Tau model

4.1.534 Supg Tau: Shakib Cvfem Momentum
Scope: Aria Material

Supg Tau: Shakib Cvfem Momentum = Shakib_Cvfem_Momentum [ Multiplier = multiplier |Use_Diffusion = use_diffusion |Use_Time = use_time ]

<table>
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<tr>
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</table>

Summary A Shakib SUPG Tau model

4.1.535 Supg Tau: Shakib Cvfem Mixture Fraction
Scope: Aria Material

Supg Tau: Shakib Cvfem Mixture Fraction = Shakib_Cvfem_Mixture_Fraction [ Multiplier = multiplier |Use_Diffusion = use_diffusion |Use_Time = use_time ]

<table>
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<th>Default</th>
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</table>

Summary A Shakib SUPG Tau model
4.1.536  Supg Tau: Shakib Cvfem Level Set
Scope: Aria Material

Supg Tau: Shakib Cvfem Level Set = Shakib_Cvfem_Level_Set [ Multiplier = multiplier | Use_Diffusion = use_diffusion | Use_Time = use_time ]

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Summary  A Shakib SUPG Tau model

4.1.537  Supg Tau: Shakib Cvfem Dispersed Phase Momentum
Scope: Aria Material

Supg Tau: Shakib Cvfem Dispersed Phase Momentum = Shakib_Cvfem_Dispersed_Phase_Momentum [ Multiplier = multiplier | Use_Diffusion = use_diffusion | Use_Time = use_time ]

<table>
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</table>

Summary  A Shakib SUPG Tau model

4.1.538  Supg Tau: Classic Mixture Fraction
Scope: Aria Material

Supg Tau: Classic Mixture Fraction = Classic_Mixture_Fraction [ Epsilon = epsilon | Multiplier = multiplier ]

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<th>Default</th>
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Summary  A Shakib SUPG Tau model

4.1.539  Supg Tau: Shakib Mixture Fraction
Scope: Aria Material

Supg Tau: Shakib Mixture Fraction = Shakib_Mixture_Fraction [ Multiplier = multiplier | Use_Diffusion = use_diffusion | Use_Time = use_time ]

<table>
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</table>
4.1.540 Supg Tau: Classic Species
Scope: Aria Material

Supg Tau: Classic Species = Classic_Species [ Epsilon = \( \epsilon \) |Multiplier = multiplier ]

<table>
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<th>Parameter</th>
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<th>Default</th>
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</thead>
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<tr>
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</table>

4.1.541 Supg Tau: Shakib Species
Scope: Aria Material

Supg Tau: Shakib Species = Shakib_Species [ Multiplier = multiplier |Use_Diffusion = use_diffusion |Use_Time = use_time ]

<table>
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<th>Parameter</th>
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</table>

4.1.542 Supg Tau: Shakib Mass Balance
Scope: Aria Material

Supg Tau: Shakib Mass Balance = Shakib_Mass_Balance [ Multiplier = multiplier |Use_Diffusion = use_diffusion |Use_Time = use_time ]

<table>
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<tr>
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<tr>
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</table>

4.1.543 Supg Tau: Shakib Charge Density
Scope: Aria Material

Supg Tau: Shakib Charge Density = Shakib_Charge_Density [ H = h |Use_Advection = use_advection |Use_Diffusion = use_diffusion |Use_Time = use_time |Multiplier = multiplier ]
### 4.1.544 Surface Tension

**Scope:** Aria Material

Surface Tension [ {of|species|subindex} Species ]= Constant [ Sigma = sigma ]

<table>
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</table>

**Summary** Constant value

### 4.1.545 Surface Tension: Linear T

**Scope:** Aria Material

Surface Tension: Linear T [ {of|species|subindex} Species ]= Linear_T [ Sigma0 = sigma0 | Dsigmadt = dsigmadT | T_Ref = T_ref ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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<tbody>
<tr>
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</table>

**Summary** Linear T surface tension

### 4.1.546 Suspension Flux: Phillips

**Scope:** Aria Material

Suspension Flux: Phillips [ {of|species|subindex} Species ]= Phillips [ K_Mu = k_mu | K_C = k_c | Phi_Max = phi_max | Beta = beta | Particle_Radius = particle_radius | K_0 = k_0 | Phi_Tol = phi_tol | Use_Fd = use_fd ]

---

Parameter | Value | Default |
<table>
<thead>
<tr>
<th></th>
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<tbody>
<tr>
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</table>

Summary A Shakib SUPG Tau model
### 4.1.547 Suspension Flux: Fad Phillips

**Scope:** Aria Material

Suspension Flux: Fad Phillips \[ \{ \text{of|species|subindex} \} \text{Species} \] = Fad_Phillips \[ K_Mu = k_mu \mid K_C = k_c \mid Phi_Max = phi_max \mid Beta = beta \mid Particle_Radius = particle_radius \mid K_0 = k_0 \mid Phi_Tol = phi_tol \]

<table>
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<tr>
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</tr>
</tbody>
</table>

### 4.1.548 Suspension Flux: Balance

**Scope:** Aria Material

Suspension Flux: Balance \[ \{ \text{of|species|subindex} \} \text{Species} \] = Balance \[ A = a \mid Mu_S = mu_s \mid Lambda_1 = lambda_1 \mid Lambda_2 = lambda_2 \mid Lambda_3 = lambda_3 \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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</tr>
<tr>
<td>a</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>mu_s</td>
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</tr>
<tr>
<td>lambda_1</td>
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</tr>
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<td>lambda_2</td>
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<td>undefined</td>
</tr>
<tr>
<td>lambda_3</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

### 4.1.549 Suspension Flux: Hydrostatic

**Scope:** Aria Material

Suspension Flux: Hydrostatic \[ \{ \text{of|species|subindex} \} \text{Species} \] = Hydrostatic \[ Gx = gx \mid Gy = gy \mid Gz = gz \mid Lambda = lambda \mid Fluid_Density = fluid_density \mid B = b \mid Phi_Max = phi_max \]

<table>
<thead>
<tr>
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<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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<td>Lambda</td>
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<tr>
<td>Fluid_Density</td>
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<td>undefined</td>
</tr>
<tr>
<td>B</td>
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</tr>
<tr>
<td>Phi_Max</td>
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</tr>
</tbody>
</table>
### Summary

Hydrostatic source for suspension equation

### 4.1.550 Suspension Hindrance Function: Morris

**Scope:** Aria Material

Suspension Hindrance Function: Morris \( \{ \text{of|species|subindex} \text{Species} \} = \text{Morris} \ [ \text{Alpha} = \text{alpha} \ | \text{Phi}\_\text{Max} = \text{phi\_max} \ | \text{Phi}\_\text{Tol} = \text{phi\_tol} ] \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
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</tr>
<tr>
<td>phi_tol</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

### 4.1.551 Suspension Normal Viscosity: Morris

**Scope:** Aria Material

Suspension Normal Viscosity: Morris \( \{ \text{of|species|subindex} \text{Species} \} = \text{Morris} \ [ \text{K}\_\text{N} = \text{K}\_\text{n} \ | \text{Phi}\_\text{Max} = \text{phi\_max} \ | \text{Phi}\_\text{Tol} = \text{phi\_tol} \ | \text{Mu}\_\text{S} = \text{mu}\_\text{s} ] \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
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</tr>
<tr>
<td>mu_s</td>
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</tr>
</tbody>
</table>

### 4.1.552 Suspension Q Tensor: Isotropic

**Scope:** Aria Material

Suspension Q Tensor: Isotropic \( \{ \text{of|species|subindex} \text{Species} \} = \text{Isotropic} \ [ ] \)

<table>
<thead>
<tr>
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<th>Value</th>
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</thead>
<tbody>
<tr>
<td>Species</td>
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</tbody>
</table>

### 4.1.553 Suspension Q Tensor: Ct Aligned

**Scope:** Aria Material
Suspension Q Tensor: Ct Aligned [ {of|species|subindex} Species ]= Ct_Aligned [ Lambda_1 = lambda_1 | Lambda_2 = lambda_2 | Lambda_3 = lambda_3 ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
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</tr>
<tr>
<td>lambda_2</td>
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<td>undefined</td>
</tr>
<tr>
<td>lambda_3</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.554 Suspension Q Tensor: Flow Aligned
Scope: Aria Material

Suspension Q Tensor: Flow Aligned [ {of|species|subindex} Species ]= Flow_Aligned_ [ Lambda_A = lambda_a | Lambda_B = lambda_b ]

<table>
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<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>lambda_a</td>
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<tr>
<td>lambda_b</td>
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</tbody>
</table>

4.1.555 Species Names
Scope: Aria Material

Species Names {=|are|is} Names...

<table>
<thead>
<tr>
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<th>Default</th>
</tr>
</thead>
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<td>Names</td>
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</tbody>
</table>

Description: This defines the available chemical species, if they are not already specified by an external property TPL.

4.1.556 Temperature
Scope: Aria Material

Temperature [ {of|species|subindex} Species ]= Constant [ Power_Output = power_output | Flux_Output = flux_output | Toggle = toggle | Value = value ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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</tr>
<tr>
<td>power_output</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>flux_output</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>toggle</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
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<tr>
<td>value</td>
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</tbody>
</table>

Summary: Constant value

4.1.557 Temperature: Calore User Sub
Scope: Aria Material
4.1.558 Temperature: From No Material Phase

Scope: Aria Material

Summary: Copy the temperature from NO_MATERIAL_PHASE to the desired material phase.

4.1.559 Temperature: Cantera

Scope: Aria Material

Summary: Cantera enthalpy computed from temperature
4.1.560 Temperature: Fortran User Sub
Scope: Aria Material

Temperature: Fortran_User_Sub [ {of|species|subindex} Species ] = Fortran_User_Sub [ Power_Output = power_output | Flux_Output = flux_output | Toggle = toggle | Multiplier = multiplier | Name = name | Type = type | Real_Data = real_data | Int_Data = int_data | Resource_Name = resource_name | Data = data | Material_Data_Block = material_data_block ]

<table>
<thead>
<tr>
<th>Parameter</th>
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<th>Default</th>
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</thead>
<tbody>
<tr>
<td>Species</td>
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<td>flux_output</td>
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<tr>
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<tr>
<td>int_data</td>
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</tr>
<tr>
<td>material_data_block</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary Dirichlet value from a Fortran user subroutine

4.1.561 Temperature: Cht Robin
Scope: Aria Material

Temperature: Cht_Robin [ {of|species|subindex} Species ] = Cht_Robin [ Power_Output = power_output | Flux_Output = flux_output | Toggle = toggle | Temperature_Field = temperature_field ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
<tr>
<td>Species</td>
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<td>power_output</td>
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<td>flux_output</td>
<td>&quot;string&quot;</td>
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<tr>
<td>toggle</td>
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</tr>
<tr>
<td>temperature_field</td>
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</table>

Summary Dirichlet boundary condition for interface coupled with Robin style BC CHT_ROBIN. This BC computes the heat flux for the other side.

4.1.562 Temperature: Correlation Wall
Scope: Aria Material

Temperature: Correlation_Wall [ {of|species|subindex} Species ] = Correlation_Wall [ Tw = Tw | Bulk_Node_Interface = bulk_node_interface ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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</tr>
</thead>
<tbody>
<tr>
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<tr>
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<td>&quot;string&quot;</td>
<td>undefined</td>
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</tbody>
</table>
4.1.563 Temperature: Correlation Fluid

**Scope:** Aria Material

Temperature: Correlation Fluid [ {of|species|subindex} Species ] = Correlation_Fluid [ T = t [Bulk_Node_Interface = bulk_node_interface ] ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tbody>
</table>

Summary: Correlation fluid temperature

4.1.564 Tensor Bulk Conductivity

**Scope:** Aria Material

Tensor Bulk Conductivity [ {of|species|subindex} Species ] = Constant [ Xx = xx | T11 = t11 | Xy = xy | T12 = t12 | Xz = xz | T13 = t13 | Yx = yx | T21 = t21 | Yy = yy | T22 = t22 | Yz = yz | T23 = t23 | Zx = xz | T31 = t31 | Zy = yz | T32 = t32 | Zz = zz | T33 = t33 ]

<table>
<thead>
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<th>Parameter</th>
<th>Value</th>
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<tbody>
<tr>
<td>Species</td>
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<td>t12</td>
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<td>undefined</td>
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<tr>
<td>xz</td>
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<td>undefined</td>
</tr>
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<td>t13</td>
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</tr>
<tr>
<td>yx</td>
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<td>undefined</td>
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<tr>
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</tbody>
</table>

Summary: Constant tensor value for the bulk conductivity.

4.1.565 Tensor Bulk Conductivity: Diagonal

**Scope:** Aria Material

Summary: Constant tensor value for the bulk conductivity.
Tensor Bulk Conductivity: Diagonal \[
\text{Species} = \text{Diagonal [Variable = variable]}
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</tbody>
</table>

4.1.566 Tensor Bulk Conductivity: Volume Average
Scope: Aria Material

Tensor Bulk Conductivity: Volume Average \[
\text{Species} = \text{Volume\_Average [ ]}
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
</tbody>
</table>

Summary Bulk conductivity tensor computed from volume average of species bulk conductivities

4.1.567 Tensor Bulk Conductivity Scaling
Scope: Aria Material

Tensor Bulk Conductivity Scaling \[
\text{Species} = \text{Constant [ S = s ]}
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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<td>s</td>
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</table>

Summary Constant value

4.1.568 Tensor Bulk Conductivity Scaling: T Exponent
Scope: Aria Material

Tensor Bulk Conductivity Scaling: T Exponent \[
\text{Species} = \text{T\_Exponent [ \text{Kbulk\_Ref = kbulk\_ref} \mid \text{T\_Ref = t\_ref} \mid \text{N = n} ]}
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<tr>
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</tbody>
</table>

Summary Tensor thermal conductivity scaling for bulk material

4.1.569 Tensor Electrical Conductivity: Calore User Sub
Scope: Aria Material
Tensor Electrical Conductivity: Calore User Sub [ {of|species|subindex} Species ] = Calore_User_Sub [ Name = name | Type = type | Multiplier = multiplier | Material_Data_Block = material_data_block | Data = data ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tbody>
</table>

Summary: Tensor values for temperature dependent electrical conductivity from a calore user subroutine

4.1.570 Tensor Electrical Conductivity

Scope: Aria Material

Tensor Electrical Conductivity [ {of|species|subindex} Species ] = Constant [ Xx = xx | T11 = t11 | Xy = xy | T12 = t12 | Xz = xz | T13 = t13 | Yx = yx | T21 = t21 | Yy = yy | T22 = t22 | Yz = yz | T23 = t23 | Zx = xz | T31 = t31 | Zy = yz | T32 = t32 | Zz = zz | T33 = t33 ]

<table>
<thead>
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<th>Value</th>
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<tbody>
<tr>
<td>Species</td>
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Summary: Constant tensor value for the electrical conductivity

4.1.571 Tensor Electromigration Coefficient

Scope: Aria Material

Tensor Electromigration Coefficient [ {of|species|subindex} Species ] = Constant [ Xx = xx | T11 = t11 | Xy = xy | T12 = t12 | Xz = xz | T13 = t13 | Yx = yx | T21 = t21 | Yy = yy | T22 = t22 | Yz = yz | T23 = t23 | Zx = xz | T31 = t31 | Zy = yz | T32 = t32 | Zz = zz | T33 = t33 ]

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Parameter | Value | Default
--- | --- | ---
Species | string | undefined
xx | real | undefined
t11 | real | undefined
xy | real | undefined
t12 | real | undefined
xz | real | undefined
t13 | real | undefined
yx | real | undefined
t21 | real | undefined
yy | real | undefined
t22 | real | undefined
yz | real | undefined
t23 | real | undefined
zx | real | undefined
t31 | real | undefined
zy | real | undefined
t32 | real | undefined
zz | real | undefined
t33 | real | undefined

Summary | Constant tensor value for the Soret coefficient

### 4.1.572 Tensor Soret Coefficient

**Scope:** Aria Material
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**Summary**  
Constant tensor value for the Soret coefficient

### 4.1.573 Tensor Species Diffusivity

**Scope:** Aria Material

Tensor Species Diffusivity [ {of|species|subindex} Species ] = Constant [ Xx = xx | T11 = t11 | Xy = xy | T12 = t12 | Xz = xz | T13 = t13 | Yx = yx | T21 = t21 | Yy = yy | T22 = t22 | Yz = yz | T23 = t23 | Zx = xx | T31 = t31 | Zy = yz | T32 = t32 | Zz = zz | T33 = t33 ]

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**Summary**  
Constant tensor value for the species diffusivity
4.1.574 Tensor Species Mobility: Nernst Einstein
Scope: Aria Material

Tensor Species Mobility: Nernst Einstein \( \{ \text{of|species|subindex} \} \text{Species} \) = Nernst_Einstein
\( \{ R = r \ | \text{Temperature\_Material\_Phase} = \text{temperature\_material\_phase} \} \)

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Summary: Tensor mobility from diffusivity, \( M = \frac{D}{RT} \).

4.1.575 Tensor Species Mobility
Scope: Aria Material

Tensor Species Mobility \( \{ \text{of|species|subindex} \} \text{Species} \) = Constant \( \{ \text{Xx} = xx \ | \text{T11} = t11 \ | \text{xy} = xy \ | \text{T12} = t12 \ | \text{Xz} = xz \ | \text{T13} = t13 \ | \text{Yx} = yx \ | \text{T21} = t21 \ | \text{Yy} = yy \ | \text{T22} = t22 \ | \text{Yz} = yz \ | \text{T23} = t23 \ | \text{Zx} = zx \ | \text{T31} = t31 \ | \text{Zy} = zy \ | \text{T32} = t32 \ | \text{Zz} = zz \ | \text{T33} = t33 \} \)

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Summary: Constant tensor value for the electrical conductivity

4.1.576 Tensor Thermal Conductivity: Calore User Sub
Scope: Aria Material

Tensor Thermal Conductivity: Calore User Sub \( \{ \text{of|species|subindex} \} \text{Species} \) = Calore_User_Sub
\( \{ \text{Name} = name \ | \text{Type} = type \ | \text{Multiplier} = multiplier \ | \text{Material\_Data\_Block} = \text{material\_data\_block} \ | \text{Data} = data \} \)
### Parameter Value Default

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**Summary**

Tensor values for the thermal conductivity from a calore user subroutine

#### 4.1.577 Tensor Thermal Conductivity

**Scope:** Aria Material

\[
\text{Tensor Thermal Conductivity} \{ \text{species} \} \text{ Species } = \text{ Constant} \[ Xx = xx \mid T11 = t11 \mid Xy = xy \mid T12 = t12 \mid Xz = xz \mid T13 = t13 \mid Yx = yx \mid T21 = t21 \mid Yy = yy \mid T22 = t22 \mid Yz = yz \mid T23 = t23 \mid Zx = zx \mid T31 = t31 \mid Zy = zy \mid T32 = t32 \mid Zz = zz \mid T33 = t33 \]

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<tr>
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</table>

**Summary**

Constant tensor value for the thermal conductivity

#### 4.1.578 Tensor Thermal Conductivity: Mesh Input

**Scope:** Aria Material

\[
\text{Tensor Thermal Conductivity: Mesh Input} \{ \text{species} \} \text{ Species } = \text{ Mesh Input} \[ \text{Multiplier} = \text{multiplier} \mid \text{Name} = \text{name} \mid \text{Type} = \text{type} \mid \text{Axis1} = \text{axis1} \mid \text{Axis2} = \text{axis2} \mid \text{Tolerance} = \text{tolerance} \]

212
Parameter | Value | Default
--- | --- | ---
Species | string | undefined
multiplier | real | undefined
name | "string" | undefined
type | "string" | undefined
axis1 | "string" | undefined
axis2 | "string" | undefined
tolerance | real | undefined

Summary: Tensor thermal conductivity from the input mesh database

### 4.1.579 Tensor Thermal Conductivity: Mass Average

**Scope:** Aria Material

Tensor Thermal Conductivity: Mass Average

\[ \text{Species} \] = Mass_Average

Parameter | Value | Default
--- | --- | ---
Species | string | undefined

Summary: Thermal conductivity computed from mass average of species conductivities

### 4.1.580 Tensor Thermal Conductivity: Summed

**Scope:** Aria Material

Tensor Thermal Conductivity: Summed

\[ \text{Species} \] = Summed [ Contributions = contributions ]

Parameter | Value | Default
--- | --- | ---
Species | string | undefined
contributions | "string" | undefined

Summary: Tensor thermal conductivity as a sum of user-specified conductivities

### 4.1.581 Thermal Conductivity

**Scope:** Aria Material

Thermal Conductivity

\[ \text{Species} \] = Constant [ \( K = k \) ]

Parameter | Value | Default
--- | --- | ---
Species | string | undefined
k | real | undefined

Summary: Constant value

### 4.1.582 Thermal Conductivity: Calore User Sub

**Scope:** Aria Material
4.1.583 Thermal Conductivity: Cantera
Scope: Aria Material

Summary: Cantera thermal conductivity

4.1.584 Thermal Conductivity: Curing Foam
Scope: Aria Material

Summary: ?

4.1.585 Thermal Conductivity: From Prandtl
Scope: Aria Material

Summary: ?

4.1.586 Thermal Conductivity: Optically Thick
Scope: Aria Material
Thermal Conductivity: Optically Thick \[ {\text{Species}} = \text{Optically\_Thick} \]
\[ K = k \ | \beta_r = \beta_r \ | \eta = \eta \]

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<td>( \eta )</td>
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Summary: Thermal conductivity model for optically thick materials

4.1.587 Thermal Conductivity: Phase Average

Scope: Aria Material

Thermal Conductivity: Phase Average \[ {\text{Species}} = \text{Phase\_Average} \]
\[ \text{Subindex}_A = \text{subindex}_a \ | \text{Subindex}_B = \text{subindex}_b \]

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Summary: Phase-averaged values

4.1.588 Thermal Conductivity: Interpolated Phase Average

Scope: Aria Material

Thermal Conductivity: Interpolated Phase Average \[ {\text{Species}} = \text{Interpolated\_Phase\_Average} \]
\[ \text{Subindex}_A = \text{subindex}_a \ | \text{Subindex}_B = \text{subindex}_b \]

<table>
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Summary: Phase-averaged values

4.1.589 Thermal Conductivity: Power Law

Scope: Aria Material

Thermal Conductivity: Power Law \[ {\text{Species}} = \text{Power\_Law} \]
\[ A = a \ | \Gamma = \gamma \]

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Summary: Thermal conductivity as a power law in temperature

215
4.1.590 Thermal Conductivity: Saturation Power Law

Scope: Aria Material

Thermal Conductivity: Saturation Power Law \[ \text{Species} = \text{Saturation\_Power\_Law} \]

\[ K_{\text{Dry}} = k_{\text{dry}} \quad K_{\text{Wet}} = k_{\text{wet}} \quad \text{Power} = \text{power} \]

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Summary: Thermal conductivity as a power law in liquid saturation for two phase porous flows

4.1.591 Thermal Conductivity: Thermal

Scope: Aria Material

Thermal Conductivity: Thermal \[ \text{Species} = \text{Thermal} \]

\[ A = a \quad B = b \quad C = c \quad D = d \]

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Summary: Thermal conductivity as a cubit polynomial in temperature

4.1.592 Thermal Conductivity: T Exponent

Scope: Aria Material

Thermal Conductivity: T Exponent \[ \text{Species} = \text{T\_Exponent} \]

\[ K_{\text{Ref}} = \text{k\_ref} \quad T_{\text{Ref}} = \text{t\_ref} \quad N = n \]

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Summary: Thermal conductivity as a power of normalized temperature

4.1.593 Thermal Conductivity: Volume Average

Scope: Aria Material

Thermal Conductivity: Volume Average \[ \text{Species} = \text{Volume\_Average} \]
### 4.1.594 Thermal Conductivity: Summed

**Scope:** Aria Material

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**Summary**
Thermal conductivity computed from volume average of species thermal conductivities

### 4.1.595 Thermal Conductivity: Activation User Function

**Scope:** Aria Material

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**Summary**
Thermal conductivity as a sum of user-specified conductivities

### 4.1.596 Thermal Conductivity: Linear Temperature And Density

**Scope:** Aria Material

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<td>undefined</td>
</tr>
<tr>
<td>c_0</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>C_rho</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>c_t</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Thermal conductivity that is a linear function of both temperature and density. \[ \kappa = C_0 + C_{\rho} \rho + C_T T \]

### 4.1.597 Thermal Conductivity: Mass Average

**Scope:** Aria Material
Thermal Conductivity: Mass Average \[ \{\text{of species subindex}\} \text{Species} \] = Mass_Average \[

<table>
<thead>
<tr>
<th>Parameter \text{Species}</th>
<th>Value \text{string}</th>
<th>Default \text{undefined}</th>
</tr>
</thead>
</table>

Summary Thermal conductivity computed from mass average of species conductivities

4.1.598 Thermal Diffusivity
Scope: Aria Material

Thermal Diffusivity \[ \{\text{of species subindex}\} \text{Species} \] = Constant \[ D = d \]

<table>
<thead>
<tr>
<th>Parameter \Species</th>
<th>Value \text{string}</th>
<th>Default \text{undefined}</th>
</tr>
</thead>
<tbody>
<tr>
<td>\text{d}</td>
<td>\text{real}</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Constant value

4.1.599 Thermal Diffusivity: Calore User Sub
Scope: Aria Material

Thermal Diffusivity: Calore User Sub \[ \{\text{of species subindex}\} \text{Species} \] = Calore_User_Sub [ Name = \text{name} \ Type = \text{type} \ Multiplier = \text{multiplier} \ Material_Data_Block = \text{material_data_block} \ Data = \text{data} \ Scaling_Field = \text{scaling_field} ]

<table>
<thead>
<tr>
<th>Parameter \Species</th>
<th>Value \text{string}</th>
<th>Default \text{undefined}</th>
</tr>
</thead>
<tbody>
<tr>
<td>\text{name}</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{type}</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{multiplier}</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{material_data_block}</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{data}</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>\text{scaling_field}</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Calore user subroutine for the thermal diffusivity

4.1.600 Thermodynamic Pressure
Scope: Aria Material

Thermodynamic Pressure \[ \{\text{of species subindex}\} \text{Species} \] = Constant \[ \text{Value} = \text{value} \]

<table>
<thead>
<tr>
<th>Parameter \Species</th>
<th>Value \text{string}</th>
<th>Default \text{undefined}</th>
</tr>
</thead>
<tbody>
<tr>
<td>\text{value}</td>
<td>\text{real}</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Constant value
### 4.1.601 Tic Drag: Basic

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** Model for drag created in the Theory of Interacting Continua

### 4.1.602 Tic Drag Coeff

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>alpha_TIC</td>
<td>real</td>
<td>undefined</td>
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</tbody>
</table>

**Summary:** Constant value

### 4.1.603 Tortuosity Factor

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** Constant value

### 4.1.604 Tortuosity Factor: Bruggeman

**Scope:** Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>exponent</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** Bruggeman model for tortuosity factor.

### 4.1.605 Tortuosity Factor: Lanfrey

**Scope:** Aria Material
### Tortuosity Factor: Lanfrey

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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<td>undefined</td>
</tr>
<tr>
<td>sphericity</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>


#### 4.1.606 Tortuosity Factor: Comiti

Scope: Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>


#### 4.1.607 Total Internal Energy

Scope: Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

#### 4.1.608 Total Internal Energy: Porous Flow

Scope: Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Total internal energy for porous flow

#### 4.1.609 Transport Cross Section

Scope: Aria Material

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>Trans</td>
<td>trans</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Transport cross section
Parameter | Value | Default
Species | string | undefined
trans | real | undefined

Summary | Constant value

4.1.610 Transport Cross Section: Linearized
Scope: Aria Material

Transport Cross Section: Linearized [ \{of\|species\|subindex\} Species ]= Linearized [ Sigma_0 = sigma_0 \| D_Sigma_D_Rho = d\_sigma\_d\_rho \| D_Sigma_D_T = d\_sigma\_d\_t \| T_Sigma_0 = t\_sigma\_0 \| Rho_Sigma_0 = rho\_sigma\_0 ]

Parameter | Value | Default
Species | string | undefined
sigma_0 | real | undefined
d\_sigma\_d\_rho | real | undefined
d\_sigma\_d\_t | real | undefined
t\_sigma\_0 | real | undefined
rho\_sigma\_0 | real | undefined

Summary | Linearized transport cross section

4.1.611 Transported Enthalpy: Standard
Scope: Aria Material

Transported Enthalpy: Standard [ \{of\|species\|subindex\} Species ]= Standard [ ]

Parameter | Value | Default
Species | string | undefined

Summary | Transported quantity for porous enthalpy equation

4.1.612 Transported Enthalpy: Porous
Scope: Aria Material

Transported Enthalpy: Porous [ \{of\|species\|subindex\} Species ]= Porous [ ]

Parameter | Value | Default
Species | string | undefined

Summary | Transported quantity for porous enthalpy equation

4.1.613 Transported Enthalpy: Standard Fd Sens
Scope: Aria Material
Transported Enthalpy: Standard Fd Sens \[ \{ \text{of|species|subindex} \} \text{Species } ]= \text{Standard\_Fd\_Sens} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary DEPRECATED, same model as TRANSPORTED\_ENTHALPY = STANDARD

4.1.614 Transported Enthalpy: Porous Fd Sens

Scope: Aria Material

Transported Enthalpy: Porous Fd Sens \[ \{ \text{of|species|subindex} \} \text{Species } ]= \text{Porous\_Fd\_Sens} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary DEPRECATED, same model as TRANSPORTED\_ENTHALPY = POROUS

4.1.615 Turbulence Dissipation Rate

Scope: Aria Material

Turbulence Dissipation Rate \[ \{ \text{of|species|subindex} \} \text{Species } ]= \text{Constant} [ \text{Value } = \text{value} ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Constant value

4.1.616 Turbulence Dissipation Rate Diffusive Flux: Basic

Scope: Aria Material

Turbulence Dissipation Rate Diffusive Flux: Basic \[ \{ \text{of|species|subindex} \} \text{Species } ]= \text{Basic} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Basic model of turbulence dissipation rate diffusive flux

4.1.617 Turbulent Bulk Viscosity

Scope: Aria Material

Turbulent Bulk Viscosity \[ \{ \text{of|species|subindex} \} \text{Species } ]= \text{Constant} [ \text{Kappa } = \text{kappa} ]
4.1.618  Turbulent Energy Diffusive Flux: Gradient Transport
Scope:  Aria Material

\[
\text{Turbulent Energy Diffusive Flux: Gradient Transport } = \text{Gradient Transport}
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>(\kappa)</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Gradient transport model of turbulent energy diffusive flux

4.1.619  Turbulent Kinetic Energy
Scope:  Aria Material

\[
\text{Turbulent Kinetic Energy } = \text{Constant}
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>(value)</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Constant value

4.1.620  Turbulent Kinetic Energy Density
Scope:  Aria Material

\[
\text{Turbulent Kinetic Energy Density } = \text{Constant}
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
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<tr>
<td>(value)</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Constant value

4.1.621  Turbulent Kinetic Energy Diffusive Flux: Basic
Scope:  Aria Material

\[
\text{Turbulent Kinetic Energy Diffusive Flux: Basic }
\]

Summary
Parameter | Value | Default
---|---|---
Species | string | undefined

Summary  
Basic model of turbulent kinetic energy diffusive flux

4.1.622 Turbulent Mass Diffusivity

Scope: Aria Material

Turbulent Mass Diffusivity $\{of|species|subindex\} Species =$ Constant $\{Dt = Dt\}$

Parameter | Value | Default
---|---|---
Species | string | undefined
Dt | real | undefined

Summary  
Constant value

4.1.623 Turbulent Mass Diffusivity: From Schmidt

Scope: Aria Material

Turbulent Mass Diffusivity: From Schmidt $\{of|species|subindex\} Species =$ From_Schmidt

Parameter | Value | Default
---|---|---
Species | string | undefined

Summary  
Turbulent mass diffusivity from Schmidt number and viscosity

4.1.624 Turbulent Mass Fraction Diffusive Flux: Gradient Transport

Scope: Aria Material

Turbulent Mass Fraction Diffusive Flux: Gradient Transport $\{of|species|subindex\} Species =$ Gradient_Transport

Parameter | Value | Default
---|---|---
Species | string | undefined

Summary  
Gradient transport model of turbulent mass fraction diffusive flux

4.1.625 Turbulent Mixture Fraction Diffusive Flux: Gradient Transport

Scope: Aria Material

Turbulent Mixture Fraction Diffusive Flux: Gradient Transport $\{of|species|subindex\} Species =$ Gradient_Transport

Parameter | Value | Default
---|---|---
Species | string | undefined
4.1.626 Turbulent Mixture Fraction Diffusivity

Scope: Aria Material

Turbulent Mixture Fraction Diffusivity [ {of|species|subindex} Species ] = Constant [ Dt = Dt ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Dt</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.627 Turbulent Mixture Fraction Diffusivity: From Schmidt

Scope: Aria Material

Turbulent Mixture Fraction Diffusivity: From Schmidt [ {of|species|subindex} Species ] = From_Schmidt []

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Turbulent mixture fraction diffusivity from trb Schmidt number and trb viscosity

4.1.628 Turbulent Momentum Stress: Formal Newtonian Isotropic

Scope: Aria Material


<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>reference_frame</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.629 Turbulent Momentum Stress: Incompressible Newtonian Isotropic

Scope: Aria Material

Turbulent Momentum Stress: Incompressible Newtonian Isotropic [ {of|species|subindex} Species ] = Incompressible_Newtonian_Isotropic [ Reference_Frame = reference_frame ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>reference_frame</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>
4.1.630 Turbulent Momentum Stress: Newtonian Dilational Isotropic
Scope: Aria Material

\[
\text{Turbulent Momentum Stress: Newtonian Dilational Isotropic } \{ \text{Species } \}_{\text{species}_{\text{subindex}}} = \text{Newtonian}_{\text{Dilational Isotropic}} \{ \text{Reference Frame } \}_{\text{reference}_{\text{frame}}} 
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>reference_frame</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.631 Turbulent Momentum Stress: Newtonian Isotropic
Scope: Aria Material

\[
\text{Turbulent Momentum Stress: Newtonian Isotropic } \{ \text{Species } \}_{\text{species}_{\text{subindex}}} = \text{Newtonian}_{\text{Isotropic}} \{ \text{Reference Frame } \}_{\text{reference}_{\text{frame}}} 
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>reference_frame</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.632 Turbulent Prandtl Number
Scope: Aria Material

\[
\text{Turbulent Prandtl Number } \{ \text{Species } \}_{\text{species}_{\text{subindex}}} = \text{Constant } \{ \text{Pr}_T = \text{Pr}_t \}
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
</tr>
<tr>
<td>Pr_t</td>
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</tr>
</tbody>
</table>

Summary Constant value

4.1.633 Turbulent Schmidt Number
Scope: Aria Material

\[
\text{Turbulent Schmidt Number } \{ \text{Species } \}_{\text{species}_{\text{subindex}}} = \text{Constant } \{ \text{Sc}_T = \text{Sc}_t \}
\]

<table>
<thead>
<tr>
<th>Parameter</th>
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<th>Default</th>
</tr>
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<tr>
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<tr>
<td>Sc_t</td>
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</table>

Summary Constant value

4.1.634 Turbulent Thermal Conductivity: From Prandtl
Scope: Aria Material

\[
\text{Turbulent Thermal Conductivity: From Prandtl } \{ \text{Species } \}_{\text{species}_{\text{subindex}}} = \text{From Prandtl} 
\]

226
Summary  Turbulent thermal conductivity when closing energy term by \( \text{grad} T \)

### 4.1.635 Turbulent Thermal Diffusivity: From Prandtl

**Scope:** Aria Material

Turbulent Thermal Diffusivity: From Prandtl

<table>
<thead>
<tr>
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</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
</tbody>
</table>

Summary  ?

### 4.1.636 Use Data Block

**Scope:** Aria Material

Use Data Block *Name*

<table>
<thead>
<tr>
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<th>Value</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Reference to predefined data to be used by the user subroutine. These values may be changed by the user subroutine.

### 4.1.637 Valence

**Scope:** Aria Material

Valence

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
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</tr>
<tr>
<td>( z )</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Constant value

### 4.1.638 Velocity: Melting Capillary Darcy

**Scope:** Aria Material

Velocity: Melting Capillary Darcy

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td></td>
</tr>
<tr>
<td>( \text{Melt}<em>{\text{Dt}} = \text{melt}</em>{\text{DT}} )</td>
<td>real</td>
<td></td>
</tr>
<tr>
<td>( \text{Gx} = \text{gx} )</td>
<td>real</td>
<td></td>
</tr>
<tr>
<td>( \text{Gy} = \text{gy} )</td>
<td>real</td>
<td></td>
</tr>
<tr>
<td>( \text{Gz} = \text{gz} )</td>
<td>real</td>
<td></td>
</tr>
<tr>
<td>Parameter</td>
<td>Value</td>
<td>Default</td>
</tr>
<tr>
<td>-------------</td>
<td>---------</td>
<td>-----------</td>
</tr>
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<td>undefined</td>
</tr>
<tr>
<td>gx</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>gy</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>gz</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

### 4.1.639 Velocity: Darcy

**Scope:** Aria Material

Velocity: Darcy \[ \text{[of|species|subindex] Species} \] = Darcy \[ G_x = g_x \mid G_y = g_y \mid G_z = g_z \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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<td>gy</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>gz</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

### 4.1.640 Velocity: Darcy Solvent

**Scope:** Aria Material

Velocity: Darcy Solvent \[ \text{[of|species|subindex] Species} \] = Darcy_Solvent \[ G_x = g_x \mid G_y = g_y \mid G_z = g_z \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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<tbody>
<tr>
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</tr>
<tr>
<td>gz</td>
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</tbody>
</table>

**Summary**

This model is intended for problems where the porous species equation is being solved for pressure with a solvent species concentration as the mass term in order to solve for the velocity of that solvent species.

### 4.1.641 Velocity: Schloegl

**Scope:** Aria Material

Velocity: Schloegl \[ \text{[of|species|subindex] Species} \] = Schloegl \[ K_{\text{Phi}} = k_{\phi} \mid F = f \mid G_x = g_x \mid G_y = g_y \mid G_z = g_z \mid \text{Proton\_Name = proton\_name} \]

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<td>gy</td>
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4.1.642  Velocity: From Mesh Displacement
Scope:  Aria Material

Velocity: From Mesh Displacement [ {of|species|subindex} Species ]= From_Mesh_Displacement [ ]

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</table>

4.1.643  Velocity
Scope:  Aria Material

Velocity [ {of|species|subindex} Species ]= Constant [ X = x | Y = y | Z = z ]

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<tr>
<td>z</td>
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</table>

Summary  Constant value

4.1.644  Velocity: Correlation Fluid
Scope:  Aria Material

Velocity: Correlation Fluid [ {of|species|subindex} Species ]= Correlation_Fluid [ V = v | Prereq_Model = prereq_model | Bulk_Node_Interface = bulk_node_interface ]

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Summary  Correlation fluid velocity

4.1.645  Venting Volumetric Flow Rate
Scope:  Aria Material

Venting Volumetric Flow Rate [ {of|species|subindex} Species ]= Constant [ Value = value ]

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<th>Value</th>
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<tbody>
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<td>value</td>
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</table>

Summary  Constant value
### 4.1.646 Venting Volumetric Flow Rate: K Factor

**Scope:** Aria Material

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</thead>
<tbody>
<tr>
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</tbody>
</table>

**Summary**

Volumetric flow rate calculated as \( \text{rate} = K \times \sqrt{(P - P_{\text{ambient}}) / \text{density}} \).

### 4.1.647 Venting Volumetric Flow Rate: K Factor With Choking

**Scope:** Aria Material

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**Summary**

Volumetric flow rate calculated as \( \text{rate} = K \times \sqrt{(P - P_{\text{ambient}}) / \text{density}} \) for \( P < P_{\text{crit}} \).

Above \( P_{\text{crit}} \) the flow rate is calculated as \( \text{rate} = K \times \sqrt{(1 - 1/critical\_pressure\_ratio) \times \frac{P}{\text{density}}} \).

### 4.1.648 Viscosity

**Scope:** Aria Material

<table>
<thead>
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**Summary**

Constant value

### 4.1.649 Viscosity: Arrhenius

**Scope:** Aria Material

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<td>mu_max</td>
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</table>
4.1.650  Viscosity: Arrhenius Carreau

Scope:  Aria Material

Viscosity: Arrhenius Carreau [{of|species|subindex} Species ]= Arrhenius_Carreau [ Mu_Zero = mu_zero | Mu_Inf = mu_inf | N = n | A = a | Lambda = lambda | K = k | T_Ref = t_ref | Skip_Sensitivities = skip_sensitivities | Sensitivity_Scaling = sensitivity_scaling | Ramp_Sensitivities_Over_Time = ramp_sensitivities_over_time ]

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</tbody>
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4.1.651  Viscosity: Bingham Wlf

Scope:  Aria Material

Viscosity: Bingham Wlf [{of|species|subindex} Species ]= Bingham_Wlf [ Mu_Zero = mu_zero | Mu_Inf = mu_inf | F = f | N = n | A = a | Lambda = lambda | Tau_Y = tau_y | Skip_Sensitivities = skip_sensitivities ]

<table>
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<td>skip_sensitivities</td>
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4.1.652  Viscosity: Bingham Wlft

Scope:  Aria Material

Viscosity: Bingham Wlft [{of|species|subindex} Species ]= Bingham_Wlft [ Mu_Zero = mu_zero | Mu_Inf = mu_inf | F = f | N = n | A = a | Lambda = lambda | Tau_Y = tau_y | T_Ref = t_ref | C1 = c1 | C2 = c2 | Skip_Sensitivities = skip_sensitivities ]
Parameter | Value | Default
---|---|---
Species | string | undefined
mu_zero | real | undefined
mu_inf | real | undefined
f | real | undefined
n | real | undefined
a | real | undefined
lambda | real | undefined
tau_y | real | undefined
t_ref | real | undefined
c1 | real | undefined
c2 | real | undefined
skip_sensitivities | integer | undefined

4.1.653 Viscosity: Carreau
Scope: Aria Material

Viscosity: Carreau [ {of|species|subindex} Species ]= Carreau [ Mu_Zero = mu_zero | Mu_Inf = mu_inf | N = n | A = a | Lambda = lambda | Skip_Sensitivities = skip_sensitivities ]

Parameter | Value | Default
---|---|---
Species | string | undefined
mu_zero | real | undefined
mu_inf | real | undefined
n | real | undefined
a | real | undefined
lambda | real | undefined
tau_y | real | undefined
t_ref | real | undefined
c1 | real | undefined
c2 | real | undefined
skip_sensitivities | integer | undefined

4.1.654 Viscosity: Carreau T
Scope: Aria Material

Viscosity: Carreau T [ {of|species|subindex} Species ]= Carreau_T [ Mu_Zero = mu_zero | Mu_Inf = mu_inf | N = n | A = a | K = k | Skip_Sensitivities = skip_sensitivities ]

Parameter | Value | Default
---|---|---
Species | string | undefined
mu_zero | real | undefined
mu_inf | real | undefined
n | real | undefined
a | real | undefined
k | real | undefined
skip_sensitivities | integer | undefined

4.1.655 Viscosity: Cantera
Scope: Aria Material

Viscosity: Cantera [ {of|species|subindex} Species ]= Cantera [ ]

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4.1.656  Viscosity: Clsm

Scope:  Aria Material

Summary  CLSM viscosity

4.1.657  Viscosity: Curing Epoxy

Scope:  Aria Material

4.1.658  Viscosity: Curing Foam

Scope:  Aria Material
### 4.1.659 Viscosity: Keyes
**Scope:** Aria Material

Viscosity: Keyes \{of|species|subindex\} Species = Keyes \[ A_0 = a_0 \mid A = a \mid A_1 = a_1 \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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<tbody>
<tr>
<td>Species</td>
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<td>undefined</td>
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<tr>
<td>a1</td>
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</table>

### 4.1.660 Viscosity: Krieger
**Scope:** Aria Material

Viscosity: Krieger \{of|species|subindex\} Species = Krieger \[ Beta = beta \mid Phi_{Max} = \phi_{max} \mid Mu_S = \mu_s \mid Use_Fd = use_fd \]

<table>
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<th>Value</th>
<th>Default</th>
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<tbody>
<tr>
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</tbody>
</table>

### 4.1.661 Viscosity: Mixture Fraction
**Scope:** Aria Material

Viscosity: Mixture Fraction \{of|species|subindex\} Species = Mixture_Fraction \[ Primaryproperty = primaryProperty \mid Secondaryproperty = secondaryProperty \]

<table>
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<th>Default</th>
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</thead>
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### 4.1.662 Viscosity: Mixture Fraction Turbulent
**Scope:** Aria Material

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Viscosity: Mixture Fraction Turbulent  
\{ of | \text{species} | \text{subindex} \} \ \text{Species} = \text{Mixture\_Fraction\_Turbulent}  
\{ \text{Primaryproperty} = \text{primaryProperty} | \text{Secondaryproperty} = \text{secondaryProperty} \}

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4.1.663  Viscosity: Morris Boulay
Scope:  Aria Material

Viscosity: Morris Boulay  
\{ of | \text{species} | \text{subindex} \} \ \text{Species} = \text{Morris\_Boulay}  
\{ K_S = K_s | \text{Phi\_Max} = \text{phi\_max} | \text{Phi\_Tol} = \text{phi\_tol} | \text{Mu\_S} = \text{mu\_s} \}

<table>
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</table>

4.1.664  Viscosity: Phase Average
Scope:  Aria Material

Viscosity: Phase Average  
\{ of | \text{species} | \text{subindex} \} \ \text{Species} = \text{Phase\_Average}  
\{ \text{Subindex\_A} = \text{subindex\_a} | \text{Subindex\_B} = \text{subindex\_b} \}

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Summary  Phase-averaged values

4.1.665  Viscosity: Interpolated Phase Average
Scope:  Aria Material

Viscosity: Interpolated Phase Average  
\{ of | \text{species} | \text{subindex} \} \ \text{Species} = \text{Interpolated\_Phase\_Average}  
\{ \text{Subindex\_A} = \text{subindex\_a} | \text{Subindex\_B} = \text{subindex\_b} \}

<table>
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Summary  Phase-averaged values
### 4.1.666 Viscosity: Power Law

**Scope:** Aria Material

Viscosity: Power Law \( \text{Species} = \text{Power Law} \ [ K = k \ | N = n ] \)

<table>
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<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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<tbody>
<tr>
<td>Species</td>
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<tr>
<td>n</td>
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### 4.1.667 Viscosity: Sutherland

**Scope:** Aria Material

Viscosity: Sutherland \( \text{Species} = \text{Sutherland} \ [ T_{\text{ref}} = t_{\text{ref}} \ | C = c ] \)

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<th>Default</th>
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<tr>
<td>c</td>
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### 4.1.668 Viscosity: Weld

**Scope:** Aria Material

Viscosity: Weld \( \text{Species} = \text{Weld} \ [ \text{Interpolated} = \text{interpolated} \ | \text{Beta} = \beta \ | \text{C} = c_0 \ | \text{C} = c_1 \ | \text{C} = c_2 \ | \text{C} = c_3 \ | \text{T}_{\text{Liq}} = T_{\text{liq}} \ | \text{T}_{\text{90}} = t_{\text{90}} \ | \text{T}_{\text{Max}} = T_{\text{max}} ] \)

<table>
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### 4.1.669 Viscosity: Thermal

**Scope:** Aria Material

Viscosity: Thermal \( \text{Species} = \text{Thermal} \ [ A = a \ | B = b \ | C = c \ | D = d ] \)

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4.1.670  Viscosity: Mass Average  
Scope: Aria Material

Viscosity: Mass Average \[ \{of\{species\{subindex}\} Species \} = \text{Mass Average} \]

<table>
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<tr>
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Summary  
Viscosity computed from mass average of species viscosities

4.1.671  Viscosity: Species Average  
Scope: Aria Material

Viscosity: Species Average \[ \{of\{species\{subindex\} Species \} = \text{Species Average} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  
Viscosity computed from mass average of species viscosities

4.1.672  Voltage  
Scope: Aria Material

Voltage \[ \{of\{species\{subindex\} Species \} = \text{Constant} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>toggle</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  
Constant value

4.1.673  Volume Fraction  
Scope: Aria Material

Volume Fraction \[ \{of\{species\{subindex\} Species \} = \text{Constant} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>
4.1.674 Volume Fraction: From Mass Fraction
Scope: Aria Material

Volume Fraction: From Mass Fraction [ {of|species|subindex} Species ] = From_Mass_Fraction [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Compute the volume fraction from the ratio of species porous density (\(\rho_{\text{bulk}} \times Y_i\)) to species solid density.

4.1.675 Volume Fraction: From Porosity
Scope: Aria Material

Volume Fraction: From Porosity [ {of|species|subindex} Species ] = From_Porosity [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Compute the volume fraction from a species porosity.

4.1.676 Volume Fraction: From Molar Volume
Scope: Aria Material

Volume Fraction: From Molar Volume [ {of|species|subindex} Species ] = From_Molar_Volume [ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Compute the volume fraction of a species as the product of it’s concentration and partial molar volume.

4.1.677 Volume Fraction: From Electrode Object
Scope: Aria Material

Summary: Get the solid volume fraction for all species tracked by the Electrode object being used to calculate electrochemical reaction source terms.

4.1.678 Volume Fraction: From Electrode Object Old

Scope: Aria Material

\[
\text{Volume Fraction: From Electrode Object Old } \{ \text{Species } \}_{\text{of}\{\text{species}\}_{\text{subindex}}} = \text{From\_Electrode\_Object\_Old} \\
n = \text{electrodeFile} \mid F = f \mid K = \text{kmolConversion} \mid J = \text{JConversion} \\
\mid M = \text{meterConversion} \mid M = \text{minCapacity} \mid D = \text{deactivationVoltage} \\
\mid V = \text{voltageVariableName} \mid A = \text{abortAction} \mid I = \text{inactiveSolidFraction} \\
\]

Parameter | Value | Default
--- | --- | ---
Species | string | undefined
electrodeFile | "string" | undefined
f | real | undefined
kmolConversion | real | undefined
JConversion | real | undefined
meterConversion | real | undefined
minCapacity | real | undefined
deactivationVoltage | real | undefined
voltageVariableName | "string" | undefined
abortAction | "string" | undefined
inactiveSolidFraction | real | undefined

Summary: Get the solid volume fraction for all species tracked by the Electrode object being used to calculate electrochemical reaction source terms.

4.1.679 Volume Fraction Gas

Scope: Aria Material

\[
\text{Volume Fraction Gas } \{ \text{Species } \}_{\text{of}\{\text{species}\}_{\text{subindex}}} = \text{Constant} \mid \text{Value } = \text{value} \\
\]

Parameter | Value | Default
--- | --- | ---
Species | string | undefined
value | real | undefined
4.1.680  Volume Fraction Gas: From Density

Scope: Aria Material

Summary  Compute the gas volume fraction from the gas density

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>rho_liquid</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>rho_gas</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>rho_initial</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.681  Volume Fraction Gas: Fromfoamtimetemp

Scope: Aria Material

Summary  Compute the gas volume fraction from the foam time/temperature profile

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>rho_liquidfluorinert</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>rho_vaporfluorinert</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>rho_air</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>massfraction_air</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>rho_initial</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

4.1.682  Volume Fraction Gas: Phase Average

Scope: Aria Material

Summary  Phase-averaged values

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>subindex_a</td>
<td>integer</td>
<td>undefined</td>
</tr>
<tr>
<td>subindex_b</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>
4.1.683 Volume Fraction Gas: Interpolated Phase Average

Scope: Aria Material

Volume Fraction Gas: Interpolated Phase Average \( \{ \text{of|species|subindex}\} \text{Species} \) = Interpolated_Phase_Average
\( \text{Subindex}_A = \text{subindex}_a \ | \text{Subindex}_B = \text{subindex}_b \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>subindex_a</td>
<td>integer</td>
<td>undefined</td>
</tr>
<tr>
<td>subindex_b</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Phase-averaged values

4.1.684 Volume Fraction Gas: From Reaction Extent

Scope: Aria Material

Volume Fraction Gas: From Reaction Extent \( \{ \text{of|species|subindex}\} \text{Species} \) = From_Reaction_Extent
\( \text{Al}_1 = \text{al}_1 \ | \text{Al}_2 = \text{al}_2 \ | \text{V}_{\text{Liq}} = \text{v}_{\text{liq}} \ | \text{N}_I = \text{n}_i \ | \text{N}_{\text{Max}} = \text{n}_{\text{max}} \ | \text{M}_w = \text{mw} \ | \text{T}_{\text{Ref}} = \text{t}_{\text{ref}} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>al_1</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>al_2</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>v_liq</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>n_i</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>n_max</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>mw</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>t_ref</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Gas volume fraction from reaction extent for foaming applications

4.1.685 Volume Fraction Gas: Species

Scope: Aria Material

Volume Fraction Gas: Species \( \{ \text{of|species|subindex}\} \text{Species} \) = Species \( \text{Gas}_{\text{Species}}_{\text{Subindex}} = \text{gas}_{\text{species}}_{\text{subindex}} \ | \text{Gas}_{\text{Species}}_{\text{Name}} = \text{gas}_{\text{species}}_{\text{name}} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>gas_species_subindex</td>
<td>integer</td>
<td>undefined</td>
</tr>
<tr>
<td>gas_species_name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Compute the gas volume fraction as \( C_g \times MW_g / \rho_g \) where \( C_g \) is the molar concentration of the gas, \( MW_g \) is the molecular weight, and \( \rho_g \) is the density of the pure gas at the current \( T, P \). The species to use for \( C_g \) and \( MW_g \) can be specified either by name or by subindex.
4.1.686 Volume Fraction Gas: From Thermal Expansion
Scope: Aria Material

Volume Fraction Gas: From Thermal Expansion [ {of|species|subindex} Species ] = From_Thermal_Expansion 
[ \Phi_0 = \phi_0 | \rho_{b0} = \rho_{b0} | \rho_{c0} = \rho_{c0} | \beta_0 = \beta_0 | \beta_1 = \beta_1 | T_0 = t_0 ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>\phi_0</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>\rho_{b0}</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>\rho_{c0}</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>\beta_0</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>\beta_1</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>t_0</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Gas volume fraction from thermal expansion of the condensed phase

4.1.687 Volume Fraction Gas: From Mass Fractions
Scope: Aria Material


<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Compute the gas volume fraction from the mass fractions and bulk densities

4.1.688 Volumetric Heat Transfer Coefficient
Scope: Aria Material

Volumetric Heat Transfer Coefficient [ {of|species|subindex} Species ] = Constant [ H = h ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>h</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.689 Wall Temperature
Scope: Aria Material

Wall Temperature [ {of|species|subindex} Species ] = Constant [ T_w = T_w ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>T_w</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>
4.1.690 Wetted Perimeter
Scope: Aria Material

Wetted Perimeter \[ \text{of} \{ \text{species} \| \text{subindex} \} \text{Species} \] = Constant \[ P = p \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>( p )</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value

4.1.691 Wetted Perimeter: Spatial User Function
Scope: Aria Material

Wetted Perimeter: Spatial User Function \[ \text{of} \{ \text{species} \| \text{subindex} \} \text{Species} \] = Spatial_User_Function \[ \text{Threshold} = \text{threshold} \| \text{Multiplier} = \text{multiplier} \| \text{Name} = \text{name} \| \text{Bar} = \text{bar} \| X = x \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>( \text{threshold} )</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>( \text{multiplier} )</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>( \text{name} )</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>( \text{bar} )</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>( x )</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Spatially varying wetted perimeter based on a coordinate dependent user function

4.1.692 Wetted Perimeter: Correlation Spatial User Function
Scope: Aria Material

Wetted Perimeter: Correlation Spatial User Function \[ \text{of} \{ \text{species} \| \text{subindex} \} \text{Species} \] = Correlation_Spatial_User_Function \[ \text{Threshold} = \text{threshold} \| \text{Multiplier} = \text{multiplier} \| \text{Name} = \text{name} \| \text{Bar} = \text{bar} \| X = x \| \text{Bulk_Node_Interface} = \text{bulk_node_interface} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>( \text{threshold} )</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>( \text{multiplier} )</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>( \text{name} )</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>( \text{bar} )</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>( x )</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>( \text{bulk_node_interface} )</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Correlation spatially varying wetted perimeter based on a coordinate dependent user function
4.1.693  Wetting Phase Retardation

Scope:  Aria Material

Wetting Phase Retardation [ {of|species|subindex} Species ]=< Constant [ R = r ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>r</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Constant value

4.1.694  Wisdom: Magic Eight Ball

Scope:  Aria Material

Wisdom: Magic Eight Ball [ {of|species|subindex} Species ]= Magic_Eight_Ball [ Question = question ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>question</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  The Eight ball knows all

4.2  BAR AREA

BAR AREA = MODEL[ param1 = val1, param2 = val2 . . . ]

Description  Specifies the material model for the bar cross-sectional area.

Summary  Specifies the material model and parameters for the bar cross-sectional area.

Parent Block(s)  ARIA MATERIAL

4.2.1  BAR AREA = CONSTANT

Parameters  A = REAL

Example  Bar Area = Constant A = 0.8

Description  A is the value of the bar cross-sectional area.
4.2.2 BAR AREA = ELEMENT_ATTRIBUTE

Parameters

MULTIPLIER = REAL

Example

Bar Area = Element_Attribute multiplier = 1.0

Description

For bar elements the ExodusII mesh file can contain area attribute values for each element within an element block. This value can then be retrieved and used by the application. This model is used to scale the value of area attribute by Multiplier to obtain the bar cross-sectional area for a particular element.

4.3 BETA

BETA = MODEL [param1 = val1, param2 = val2 ...]

Description

Specifies the coefficient for thermal stress.

Summary

The solid stress $T$ is given by

$$T = \lambda E_{kk} I + 2\mu E - \beta (T - T_{\text{ref}}) I$$

(4.1)

where $\lambda$ and $\mu$ are the Lamé coefficients, $E = \frac{1}{2} (\nabla d + \nabla d^T)$ is the deformation tensor, $\beta$ is the coefficient of thermal stress, $T$ is temperature and $T_{\text{ref}}$ is the thermal strain reference temperature.

These Lamé coefficients are related to the more standard Young’s modulus, Poisson’s ratio and CTE ($\alpha$) as follows:

$$2\mu = \frac{E}{(1 + \nu)}$$

(4.2)

$$\lambda = \frac{\nu E}{(1 + \nu) (1 - 2\nu)} = 2\mu \frac{\nu}{(1 - 2\nu)}$$

(4.3)

$$\beta = \frac{\alpha E}{(1 - 2\nu)} = \alpha (3\lambda + 2\mu)$$

(4.4)

When a user supplies the Young’s modulus, Poisson’s ratio and CTE properties Aria internally converts them into the Lamé coefficients.

Parent Block(s)

ARIA MATERIAL

4.3.1 BETA = CONSTANT

Parameters

BETA = REAL

Example

BETA = CONSTANT BETA = 1.0

Description

BETA is the value of $\beta$. 

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4.3.2 BETA = CONVERTED

Parameters (None)

Example BETA = Converted

Description Aria will use Young's modulus, Poisson ratio and CTE to compute the Lamé β coefficient. Supplying the Lamé coefficients is more computationally efficient but perhaps less convenient, especially if the material properties are varying (e.g., temperature dependent in a non-isothermal problem).

4.3.3 BETA = ENCORE_FUNCTION

Parameters
NAME = STRING
[EVAL_TYPE = STRING]

Example Beta = Encore_Function Name=My_Function

Description Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```
Begin String Function My_Function
  Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

4.3.4 BETA = LINEAR

Parameters
A = REAL
B = REAL

Example BETA = LINEAR A = 1.0 B = -.005

Description β is a linear function of temperature T,

\[ \beta = A + BT \]  (4.5)
4.4 BULK VISCOSITY

BULK VISCOSITY = \textit{MODEL}[\text{param}_1 = \text{val}_1, \text{param}_2 = \text{val}_2 \ldots ]

**Description**
Specifies the material model for the fluid bulk viscosity.

**Summary**
Specifies the material model for the fluid bulk viscosity.

**Parent Block(s)**
ARIA MATERIAL

### 4.4.1 BULK VISCOSITY = CONSTANT

**Parameters**
KAPPA = \textit{REAL}

**Example**
BULK VISCOSITY = CONSTANT KAPPA = 1.0e-5

**Description**
KAPPA is the value of the constant fluid bulk viscosity.

### 4.4.2 BULK VISCOSITY = CURING_FOAM

**Parameters**
VFRAC_SUBINDEX = \textit{INT}
EXTENT_SUBINDEX = \textit{INT}
PHI_ZERO = \textit{REAL}
[A = \textit{REAL}]
[B = \textit{REAL}]
[C = \textit{REAL}]
[\kappa = \textit{REAL}]

**Example**
Bulk Viscosity = Curing_Foam Vfrac_Subindex=1 Extent_Subindex=2 Phi_Zero=0.45

**Description**
For a curing epoxy with volume fraction \( \phi \) and extent of reaction \( \xi \) the viscosity is given by

\[
\kappa = \frac{4}{3} \mu_o \frac{\phi - \phi - 1}{\phi - \phi} \tag{4.6}
\]

where \( \mu_o \) is given by

\[
\mu_o = (a - bT) \left( \frac{\xi^2 - \xi^2_c}{\xi^2_c} \right)^c \tag{4.7}
\]

where \( T \) is the temperature. The remaining parameters \( a, b, c \) and \( \xi_c \) have default values of \( a = 20, b = 0.22, c = -4/3 \) and \( \xi_c = 0.45 \) though they can be overridden with the optional model parameters.

**NOTE:** The volume fraction is assumed to be a SPECIES field with the subindex provided by the VFRAC_SUBINDEX parameter. Likewise, the extent of reaction field is assumed to be a SPECIES field with the subindex provided by the EXTENT_SUBINDEX parameter.
4.4.3 BULK VISCOSITY = ENCORE_FUNCTION

Parameters

NAME = STRING
[EVAL_TYPE = STRING]

Example

Bulk Viscosity = Encore_Function Name=My Function

Description

Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

Begin String Function My Function
Value is "200 + 1.0*t"
End

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

4.5 CTE

CTE = MODEL [param1 = val1, param2 = val2 ...]

Description

Specifies the coefficient of thermal expansion.

Summary

The solid stress $\mathbf{T}$ is given by

$$
\mathbf{T} = \lambda E \mathbf{I} + 2\mu \mathbf{E} - \beta (T - T_{\text{ref}}) \mathbf{I}
$$

(4.8)

where $\lambda$ and $\mu$ are the Lamé coefficients, $\mathbf{E} = \frac{1}{2} \left( \nabla \mathbf{d} + \nabla \mathbf{d}^T \right)$ is the deformation tensor, $\beta$ is the coefficient of thermal stress, $T$ is temperature and $T_{\text{ref}}$ is the thermal strain reference temperature.

These Lamé coefficients are related to the more standard Young’s modulus, Poisson’s ratio and CTE ($\alpha$) as follows:

$$
2\mu = \frac{E}{(1+\nu)} \quad \text{(4.9)}
$$

$$
\lambda = \frac{\nu E}{(1+\nu)(1-2\nu)} = 2\mu \frac{\nu}{(1-2\nu)} \quad \text{(4.10)}
$$

$$
\beta = \alpha \frac{E}{(1-2\nu)} = \alpha \left( 3\lambda + 2\mu \right) \quad \text{(4.11)}
$$

When a user supplies the Young’s modulus, Poisson’s ratio and CTE properties Aria internally converts them into the Lamé coefficients.

Supplying the Lamé coefficients is more computationally efficient but perhaps less convenient, especially if the material properties are varying (e.g., temperature dependent in a non-isothermal problem).
4.5.1 CTE = CONSTANT

Parameters

CTE = REAL

Example

CTE = CONSTANT cte = 1.0

Description

CTE is the value of the coefficient of thermal expansion.

4.5.2 CTE = ENCORE_FUNCTION

Parameters

NAME = STRING
[EVAL_TYPE = STRING]

Example

CTE = Encore_Function Name=My_Function

Description

Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```
Begin String Function My_Function
  Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

4.6 CURRENT DENSITY

CURRENT DENSITY = MODEL[param1 = val1, param2 = val2 ...]

Description

Specifies the material (constitutive) model for the current density in the bulk.

Summary

Specifies the material (constitutive) model for the current density in the bulk.
4.6.1 CURRENT DENSITY = BASIC

This is an alias for OHMS_LAW.

Example  Current Density = Basic

4.6.2 CURRENT DENSITY = DIELECTRIC_DISPLACEMENT

Parameters  (none)

Example  Current Density = Dielectric_Displacement

Description  The current density $J$ is taken to be

$$J = -\varepsilon \nabla \frac{\partial V}{\partial t} - \frac{\partial \varepsilon}{\partial t} \nabla V$$  \hspace{1cm} (4.12)

where $\varepsilon$ is the electrical permittivity and $V$ is the voltage (electric potential). This model can be used to simulate dielectric materials adjoined to conducting materials in transient problems.

This arises from time-differentiating Gauss' Law

$$- \nabla \cdot (\varepsilon \nabla V) = \rho_f$$  \hspace{1cm} (4.13)

which gives

$$- \nabla \cdot \left( \varepsilon \nabla \frac{\partial V}{\partial t} + \frac{\partial \varepsilon}{\partial t} \nabla V \right) = \frac{\partial \rho_f}{\partial t}.$$  \hspace{1cm} (4.14)

The left hand side of this equation can be used to eliminate the $\partial \rho_f / \partial t$ from the current density equation.

4.6.3 CURRENT DENSITY = OHMS_LAW

Parameters  (none)

Example  Current Density = Ohms_Law

Description  The current density $J$ is given by Ohm’s Law,

$$J = -\sigma_e \nabla V$$  \hspace{1cm} (4.15)

where $\sigma_e$ is the electrical conductivity and $V$ is the voltage (electric potential).

4.6.4 CURRENT DENSITY = THERMOELECTRIC

Parameters  (none)

250
Current Density = Thermoelectric

The current density \( J \) is given by a combination of Ohm’s Law and a contribution related to the temperature gradient,

\[
J = -\sigma_e (\nabla V + \alpha \nabla T)
\]

(4.16)

where \( \sigma_e \) is the electrical conductivity, \( V \) is the voltage (electric potential), \( \alpha \) is the Seebeck coefficient and \( T \) is the temperature. This current density is used in conjunction with a THERMOELECTRIC heat conduction 4.15.5.

\section{4.7 DENSITY}

\[\text{DENSITY} = \text{MODEL}[\text{param}_1 = \text{val}_1, \text{param}_2 = \text{val}_2 \ldots]\]

\textbf{Description} Specifies the material model for the density.

\textbf{Summary} Specifies the material model for the density.

\textbf{Parent Block(s)} ARIA MATERIAL

\subsection{4.7.1 DENSITY = CALORE_USER_SUB}

\textbf{Parameters}

\begin{itemize}
  \item \texttt{NAME = STRING}
  \item \texttt{TYPE = STRING}
  \item \texttt{[MULTIPLIER = REAL]}
  \item \texttt{[NR = INT]}
  \item \texttt{[R0 = INT]}
  \item \texttt{[R1 = INT(etc.)]}
  \item \texttt{[NI = INT]}
  \item \texttt{[I0 = INT]}
  \item \texttt{[I1 = INT(etc.)]}
\end{itemize}

\textbf{Example}

\begin{verbatim}
Density = Calore_User_Sub name=w80afftuser type=element NR=2 R0=1000 R1=0.7
\end{verbatim}

\textbf{Description} \texttt{NAME} is the name of the user subroutine was registered with. \texttt{NR} and \texttt{NI} are the numbers of real and integer parameters, respectively. Each real (int) parameter is supplied by the \texttt{Rn} (\texttt{In}) parameter. \textbf{Note}: the parameters use zero based counting. The \texttt{TYPE} parameter denotes the API the user subroutine uses; currently Aria only supports types \texttt{ELEMENT} and \texttt{NODE}.

The optional \texttt{MULTIPLIER} parameter (defaults to 1.0) can be supplied to scale the output of the user subroutine, i.e., the flux \( f \) will be \( f = mf_{\text{user}} \) where \( m \) is the multiplier and \( f_{\text{user}} \) is the output of the user subroutine.

See section 9.2.2 for a more complete example of a Calore user subroutine.
4.7.2 DENSITY = COMPRESSIBLE_BOUSSINESQ

Parameters

REF_DENSITY
ALPHA = REAL
REF_PRESSURE
[BETA = REAL]
[REF_TEMPERATURE]

Example

Density = Compressible_Boussinesq Ref_Density=1.0 alpha=1e-8
Ref_Pressure=101235 beta=1e-3 Ref_Temperature=298

Description

\[ \rho = \rho_{ref} (1 + \alpha (P - P_{ref}) + \beta (T - T_{ref})) \] (4.17)

Here \( \rho_{ref} \) is specified by the \( \text{REF\_DENSITY} \) parameter and is the density at pressure \( P = P_{ref} \) (given by the \( \text{REF\_PRESSURE} \) parameter) and temperature \( T = T_{ref} \) (given by the \( \text{REF\_TEMPERATURE} \) parameter). The parameter \( \text{ALPHA} \) specifies the isothermal compressibility \( \alpha \). The parameter \( \text{BETA} \) specifies the isobaric compressibility \( \beta \).

However, if the temperature is not available, then the temperature effects are not included and the \( \text{BETA} \) and \( \text{REF\_TEMPERATURE} \) parameters do not need to be supplied.

4.7.3 DENSITY = CONSTANT

Parameters

RHO = REAL

Example

DENSITY = CONSTANT RHO = 1.0

Description

RHO is the value of the constant density.

4.7.4 DENSITY = CURING_FOAM

Parameters

R = REAL
RHO_E = REAL
RHO_F = REAL
PHI_ZERO = REAL
VFRAC_SUBINDEX = INT

Example

Molecular Weight = Constant Subindex=1 M = 17.0
Density = Curing_Foam R=8.314472E3 RHO_E=1 RHO_F=1.5 PHI_ZERO=0.2 VFRAC_SUBINDEX=1

Description

The density curing epoxy foam with volume fraction \( \phi \), molecular weight \( M \), temperature \( T \), and pressure \( p \) is given by

\[ \rho = (\phi_o - \phi) \frac{pM}{RT} + (1 - \phi_o) \rho_e + \phi \rho_f \] (4.18)

where \( R \) is the gas constant, \( \phi_o \) is the reference volume fraction in the flourinert \( \rho_e \) is the pure epoxy density and \( \rho_f \) is the pure flourinert density.
NOTE: The volume fraction is assumed to be a SPECIES field with the subindex provided by the VFRAC_SUBINDEX parameter.

### 4.7.5 DENSITY = ENCORE_FUNCTION

**Parameters**

- NAME = STRING
  - [EVAL_TYPE = STRING]

**Example**

Density = Encore_Function Name=My_Function

**Description**

Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```plaintext
Begin String Function My_Function
  Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

### 4.7.6 DENSITY = EXP_DECAY

**Parameters**

- RHOInicial = REAL
- RHOFinal = REAL
- K = REAL

**Example**

Density = Exp_Decay K=1.2 RHOInicial=1.0 RHOFinal=0.2

**Description**

This model supplies a density that is an exponential decay,

\[ \rho = \rho_f + (\rho_i - \rho_f) e^{-kt} \]  

(4.19)

where \( \rho_i \) is the initial density (RHOInicial), \( \rho_f \) is the final density (RHOFinal) and \( k \) (K) is the decay constant.

### 4.7.7 DENSITY = IDEAL_GAS

**Parameters**

- R = REAL
  - [P_REF = REAL]
  - [T_REF = REAL]

**Example**

Molecular Weight = Constant Subindex=1 M = 17.0
Molecular Weight = Constant Subindex=2 M = 23.0
Density = Ideal_Gas R=8.314472E3 T_ref=273.15 P_ref=101325.0
The density of a multicomponent ideal gas in kg m\(^{-3}\) may be written as
\[
\rho = \frac{P_{\text{ref}} + P}{R(T_{\text{ref}} + T)} \sum_{i} N_i M_i y_i
\]

where \(N\) is the number of species, \(P\) is the pressure in Pascals, \(P_{\text{ref}}\) is a reference pressure, \(R\) is the gas constant, \(T\) is the temperature, \(T_{\text{ref}}\) is a reference temperature, \(M_i\) is the molecular weight of species \(i\) in kg kmol\(^{-1}\) and \(y_i\) is the mole fraction of species \(i\). Note, the units given here and on the density card are SI units; any units may be used as long as internal consistency is maintained.

The optional reference values for the temperature and pressure allow you to solve for the temperature and pressure using relative units (e.g., Celsius temperature and gauge pressure) and but still use absolute values as required by this material model.

### 4.7.8 DENSITY = INCOMPRESSIBLE\_IDEAL\_GAS

**Parameters**
- \(R = \text{REAL}\)
- \(P_{\text{REF}} = \text{REAL}\)
- \([T_{\text{REF}} = \text{REAL}]\)

**Example**
- Molecular Weight = Constant Subindex=1 \(M = 17.0\)
- Molecular Weight = Constant Subindex=2 \(M = 23.0\)
- Density = Incompressible_Ideal_Gas \(R=8.314472\text{E3} T_{\text{ref}}=273.15\)
- \(P_{\text{ref}}=101325.0\)

**Description**
The density of a multicomponent ideal gas in kg m\(^{-3}\) may be written as
\[
\rho = \frac{P_{\text{ref}} + P}{R(T_{\text{ref}} + T)} \sum_{i} N_i M_i y_i
\]

where \(N\) is the number of species, \(P_{\text{ref}}\) is a reference pressure in pascal, \(R\) is the gas constant in J kmol\(^{-1}\) K\(^{-1}\), \(T\) is the temperature, \(T_{\text{ref}}\) is a reference temperature, \(M_i\) is the molecular weight of species \(i\) in kg kmol\(^{-1}\), and \(y_i\) is the mole fraction of species \(i\). Note, the units given here and on the density card are SI units; any units may be used as long as internal consistency is maintained.

The optional reference value for the temperature allow you to solve for the temperature using relative units (e.g., Celsius temperature) and but still use absolute values as required by this material model.

### 4.7.9 DENSITY = POLYNOMIAL

**Parameters**
- \(\text{VARIABLE} = \text{STRING}\)
- \(\text{ORDER} = \text{INT}\)
- \([\text{VARIABLE\_OFFSET} = \text{REAL}]\)
- \([\text{C0} = \text{REAL}]\)
- \([\text{C1} = \text{REAL}]\)
- \(\ldots\)
- \([\text{CN} = \text{REAL}]\)

**Example**
- Density= Polynomial Variable=Temperature Order=1 \(C0=401.0\) \(C1=88.5\)
4.7.10 DENSITY = SINGLE_COMPONENT_IDEAL_GAS

Parameters

- \( R = \text{REAL} \)
- \( M = \text{REAL} \)
- \( P_{\text{REF}} = \text{REAL} \)
- \( T_{\text{REF}} = \text{REAL} \)

Example

Density = Single_Component_Ideal_Gas R=8.314 T_ref=273.15 P_ref=101325.0

Description

This supplies the density for a single component ideal gas,

\[
\rho = \frac{(P_{\text{ref}} + P) M}{R(T_{\text{ref}} + T)}
\]

\( P \) is the pressure, \( P_{\text{ref}} \) is a reference pressure (defaults to zero), \( R \) is the gas constant, \( T \) is the temperature, \( T_{\text{ref}} \) is a reference temperature (defaults to zero), and \( M \) is the molecular weight (defaults to one).

The optional reference values for the temperature and pressure allow you to solve for the temperature and pressure using relative units (e.g., Celsius temperature and gauge pressure) and but still use absolute values as required by this material model.

4.7.11 DENSITY = THERMAL

Parameters

- \( A = \text{REAL} \)
- \( B = \text{REAL} \)
- \( C = \text{REAL} \)
- \( D = \text{REAL} \)

Example

DENSITY = THERMAL A = 1.0 B = -.005

Description

Cubic polynomial function of temperature for the density.

\[
\rho = A + BT + CT^2 + DT^3
\]

4.7.12 DENSITY = USER_FUNCTION

Parameters

- NAME = \text{STRING} \n- X = \text{STRING}
Example

Begin Definition for Function Water_Density
# Source Appendix 2 from "Transport Processes and
# Unit Operations" by C. J. Geankoplis
Type is Piecewise Linear
Begin Values
  # K kg * m^-3
  273.15 999.87
  277.15 1000.00
  283.15 999.73
  293.15 998.23
  298.15 997.08
  303.15 995.68
  313.15 992.25
  323.15 988.07
  333.15 983.24
  343.15 977.81
  353.15 971.83
  363.15 965.34
  373.15 958.38
End
End
...

Begin Aria Material Foo
...
  Density = User_Function Name=Water_Density X=Temperature
...
End Aria Material Foo

Description
A look-up function is used to compute the values of the density as a function of some other variable, i.e. \( f(x) \). The function type ("piecewise linear" in the example above) must support the differentiate() method for Newton’s method.

Here NAME is the name of the user-defined function (Water_Density in the example) and X is the Aria name of the abscissa variable (TEMPERATURE in the example). Note that X is not necessarily the same name as the abscissa variable identified in the user-defined function (T in the example).

4.8 ELECTRICAL CONDUCTIVITY

ELECTRICAL CONDUCTIVITY = MODEL [param_1 = val_1, param_2 = val_2 ...]

Description
Specifies the material model for the electrical conductivity.

Summary
Specifies the material model for the electrical conductivity.

Parent Block(s) ARIA MATERIAL
4.8.1 ELECTRICAL CONDUCTIVITY = CONSTANT

Parameters

SIGMA = REAL

Example

ELECTRICAL CONDUCTIVITY = CONSTANT SIGMA = 1.0

Description

SIGMA is the value of the constant electrical conductivity.

4.8.2 ELECTRICAL CONDUCTIVITY = ENCORE_FUNCTION

Parameters

NAME = STRING

[EVAL_TYPE = STRING]

Example

ELECTRICAL CONDUCTIVITY = Encore_Function Name=My_Function

Description

Using the Encore library, you can define your own functions in a number of ways, such as
analytically in the input file, based on the values of a Sierra field or your own compiled
functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

Begin String Function My_Function
    Value is "200 + 1.0*t"
End

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore
function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option
is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE
for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

4.8.3 ELECTRICAL CONDUCTIVITY = EXPONENTIAL

Parameters

VARIABLE = STRING

[CONSTANT = REAL]

[MULTIPLIER = REAL]

EXPONENT = REAL

Example

Electrical Conductivity = Exponential Variable=Temperature Multiplier=1.0
Exponent=-0.3

Description

Exponential function of in specified scalar variable. The electrical conductivity is computed as

\[ \sigma_e = C + Me^{EX} \]  \hspace{1cm} (4.25)

Here, \( C \) is the constant term supplied by the CONSTANT parameter which defaults to zero, \( M \)

is the value supplied by the MULTIPLIER parameter which defaults to unity, \( X \) is the variable

supplied by the VARIABLE parameter and \( E \) is the exponential multiplier provided by the

EXPONENT parameter.
4.8.4 **ELECTRICAL CONDUCTIVITY = FROM_RESISTANCE**

**Parameters**

None.

**Example**

ELECTRICAL CONDUCTIVITY = FROM_RESISTANCE

**Description**

The conductivity is computed as the inverse of the electrical resistance which must be provided separately.

4.8.5 **ELECTRICAL CONDUCTIVITY = POLYNOMIAL**

**Parameters**

| VARIABLE = STRING 
| ORDER = INT 
| [VARIABLE_OFFSET = REAL] 
| [C0 = REAL] 
| [C1 = REAL] 
| ... 
| [CN = REAL] 

**Example**

Electrical Conductivity= Polynomial Variable=Temperature Order=1 C0=401.0 C1=88.5

**Description**

Arbitrary order polynomial function of a specified scalar variable.

\[ \sigma_e = \sum_{i=0}^{N} C_i \left( X + X_o \right)^i \]  

Here, \( N \) is the order of the polynomial provided by the ORDER parameter and \( X \) is the variable supplied by the VARIABLE parameter, \( X_o \) is an optional offset (VARIABLE_OFFSET, default is zero) and \( C_i \) are the supplied coefficients. Coefficients that are not supplied default to a value of zero. The VARIABLE argument can be TIME or any internal Expression that evaluates to a scalar. For the latter case, the format of the VARIABLE argument is described in section 2.6.1.

4.8.6 **ELECTRICAL CONDUCTIVITY = TBC**

**Parameters**

| Ki = REAL 
| Ti = REAL 
| E = REAL 
| R = REAL 

**Example**

ELECTRICAL CONDUCTIVITY = TBC Ki=1.0 Ti=273 R=8.314 E=1e-3

**Description**

Thermal battery electrical conductivity model (see Ken Chen).

\[ \kappa(T) = \kappa_i \frac{T_i}{T} e^{-\frac{E}{R} \left( \frac{T}{T_i} - 1 \right)} \]  

Here, \( T \) is temperature, \( T_i \) is the initial temperature provided by Ti, \( \kappa_i \) is the electrical conductivity at \( T_i \) provided by Ki, \( R \) is the universal gas constant provided by R and \( E \) is the energy provided by E.
4.8.7 ELECTRICAL CONDUCTIVITY = THERMAL

Parameters

\[
A = \text{REAL} \\
B = \text{REAL} \\
C = \text{REAL} \\
D = \text{REAL}
\]

Example

ELECTRICAL CONDUCTIVITY = THERMAL \ A = 1.0 \ B = -0.01

Description

Cubic polynomial function of temperature for the conductivity.

\[
\sigma_e = A + BT + CT^2 + DT^3
\] (4.28)

4.9 ELECTRIC DISPLACEMENT

\[
\text{ELECTRIC DISPLACEMENT} = \text{MODEL[param}_1 \ = \ \text{val}_1, \ \text{param}_2 \ = \ \text{val}_2 \ldots]\n\]

Description

Specifies the material (constitutive) model for the electric displacement

Summary

Specifies the material (constitutive) model for the electric displacement

Parent Block(s) ARIA MATERIAL

4.9.1 ELECTRIC DISPLACEMENT = BASIC

This is an alias for LINEAR.

Example

Electric Displacement = Basic

4.9.2 ELECTRIC DISPLACEMENT = LINEAR

Parameters

\( \text{none} \)

Example

Electric Displacement = Linear

Description

The electric displacement \( D \) is linearly proportional to the electric field \( E = - \nabla V \)

\[
D = -\epsilon \nabla V
\] (4.29)

where \( \epsilon \) is the electrical permittivity and \( V \) is the voltage (electric potential).
4.10 ELECTRICAL PERMITTIVITY

ELECTRICAL PERMITTIVITY = MODEL[\text{param}_1 = \text{val}_1, \text{param}_2 = \text{val}_2 \ldots]

Description Specifies the material model for the electrical permittivity.

Summary Specifies the material model for the electrical permittivity.

Parent Block(s) ARIA MATERIAL

4.10.1 ELECTRICAL PERMITTIVITY = CONSTANT

Parameters $E = \text{REAL}$

Example ELECTRICAL PERMITTIVITY = CONSTANT E = 1.0

Description $E$ is the value of the constant electrical permittivity.

4.10.2 ELECTRICAL PERMITTIVITY = ENCORE_FUNCTION

Parameters NAME = STRING

Example ELECTRICAL PERMITTIVITY = Encore_Function Name=My_Function

Description Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```plaintext
Begin String Function My_Function
  Value is "$200 + 1.0\times t$"
End
```

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

4.11 ELECTRICAL RESISTANCE

ELECTRICAL RESISTANCE = MODEL[\text{param}_1 = \text{val}_1, \text{param}_2 = \text{val}_2 \ldots]
Description Specifies the material model for the electrical resistance.

Summary Specifies the material model for the electrical resistance.

Parent Block(s) ARIA MATERIAL

4.11.1 ELECTRICAL RESISTANCE = CONSTANT

Parameters None.

Example ELECTRICAL RESISTANCE = CONSTANT R = 1.0

Description R is the value of the constant electrical resistance.

4.11.2 ELECTRICAL RESISTANCE = ENCORE_FUNCTION

Parameters NAME = STRING
[EVAL_TYPE = STRING]

Example ELECTRICAL RESISTANCE = Encore Function Name=My_Function

Description Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```
Begin String Function My_Function
  Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

4.11.3 ELECTRICAL RESISTANCE = EXPONENTIAL

Parameters VARIABLE = STRING
[CONSTANT = REAL]
[MULTIPLIER = REAL]
EXPONENT = REAL

Example Electrical Resistance = Exponential Variable=Temperature Multiplier=1.0 Exponent=-0.3
Description  Exponential function of in specified scalar variable. The electrical resistance is computed as

\[ R = C + Me^{EX} \]  \hspace{1cm} (4.30) 

Here, \( C \) is the constant term supplied by the CONSTANT parameter which defaults to zero, \( M \) is the value supplied by the MULTIPLIER parameter which defaults to unity, \( X \) is the variable supplied by the VARIABLE parameter and \( E \) is the exponential multiplier provided by the EXPONENT parameter.

4.11.4 ELECTRICAL RESISTANCE = FROM_CONDUCTIVITY

Parameters  \( R = \text{REAL} \)

Example  ELECTRICAL RESISTANCE = FROM_CONDUCTIVITY

Description  The resistance is computed as the inverse of the electrical conductivity which must be provided separately.

4.11.5 ELECTRICAL RESISTANCE = POLYNOMIAL

Parameters  VARIABLE = \text{STRING} \hspace{1cm} \text{ORDER} = \text{INT} \hspace{1cm} [\text{VARIABLE_OFFSET} = \text{REAL}] \hspace{1cm} [\text{C0} = \text{REAL}] \hspace{1cm} [\text{C1} = \text{REAL}] \hspace{1cm} \ldots \hspace{1cm} [\text{CN} = \text{REAL}] 

Example  Electrical Resistance= Polynomial Variable=Temperature Order=1 C0=401.0 C1=88.5

Description  Arbitrary order polynomial function of a specified scalar variable.

\[ R = \sum_{i=0}^{N} C_i (X + X_o)^i \]  \hspace{1cm} (4.31) 

Here, \( N \) is the order of the polynomial provided by the ORDER parameter and \( X \) is the variable supplied by the VARIABLE parameter, \( X_o \) is an optional offset (VARIABLE_OFFSET, default is zero) and \( C_i \) are the supplied coefficients. Coefficients that are not supplied default to a value of zero. The VARIABLE argument can be TIME or any internal Expression that evaluates to a scalar. For the latter case, the format of the VARIABLE argument is described in section 2.6.1.

4.11.6 ELECTRICAL RESISTANCE = USER_FUNCTION

Parameters  NAME = \text{STRING} \hspace{1cm} X = \text{STRING} 

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Example

Begin definition for function RESISTANCE_DATA
Type is piecewise linear
Abscissa is T
Ordinate is Electrical_Resistance
Begin Values
  # [K]  [Ohm-um]
  273   1.00E-9
  323   7.99E-10
  ...
  873   1.09E-11
End Values
End definition for function RESISTANCE_DATA

... 

Begin Aria Material Foo

... 
  Electrical Resistance = User_Function Name=RESISTANCE_DATA X=Temperature
... 
End Aria Material Foo

Description
A look-up function is used to compute the values of the resistance as a function of some other variable, i.e. \( f(x) \). The function type ("piecewise linear" in the example above) must support the `differentiate()` method for Newton's method.

Here `NAME` is the name of the user-defined function (RESISTANCE_DATA in the example) and `X` is the Aria name of the abscissa variable (TEMPERATURE in the example). Note that `X` is not necessarily the same name as the abscissa variable identified in the user-defined function (T in the example).

4.12 EMISSIVITY

EMISSIVITY = MODEL[param1 = val1, param2 = val2 ...]

Description
Specifies the material model for the emissivity.

Summary
Specifies the material model for the emissivity.

Parent Block(s) ARIA MATERIAL

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4.12.1 EMISSIVITY = CALORE_USER_SUB

Parameters

- **NAME** = STRING
- **TYPE** = STRING
  - [**MULTIPLIER** = REAL]
  - [**NR** = INT]
  - [**R0** = INT]
  - [**R1** = INT(etc.)]
  - [**NI** = INT]
  - [**I0** = INT]
  - [**I1** = INT(etc.)]

Example

Emissivity = Calore_User_Sub name=w80afftuser type=element NR=2 R0=1000 R1=0.7

Description

**NAME** is the name of the user subroutine was registered with. **NR** and **NI** are the numbers of real and integer parameters, respectively. Each real (int) parameter is supplied by the **R** (**I**) parameter. **Note:** the parameters use zero based counting. The **TYPE** parameter denotes the API the user subroutine uses; currently Aria only supports types **ELEMENT** and **NODE**.

The optional **MULTIPLIER** parameter (defaults to 1.0) can be supplied to scale the output of the user subroutine, i.e., the flux \( f \) will be \( f = mf_{user} \) where \( m \) is the multiplier and \( f_{user} \) is the output of the user subroutine.

See section 9.2.2 for a more complete example of a Calore user subroutine.

4.12.2 EMISSIVITY = CONSTANT

Parameters

- **E** = REAL

Example

Emissivity = Constant E = 0.8

Description

**E** is the value of the constant emissivity.

4.12.3 EMISSIVITY = POLYNOMIAL

Parameters

- **VARIABLE** = STRING
- **ORDER** = INT
  - [**VARIABLE_OFFSET** = REAL]
  - [**C0** = REAL]
  - [**C1** = REAL]
  - ...
  - [**CN** = REAL]

Example

Emissivity= Polynomial Variable=Temperature Order=1 C0=401.0 C1=88.5

Description

Arbitrary order polynomial function of a specified scalar variable.

\[
\epsilon = \sum_{i=0}^{N} C_i (X + X_0)^i
\]        \( (4.32) \)
Here, $N$ is the order of the polynomial provided by the ORDER parameter and $X$ is the variable supplied by the VARIABLE parameter, $X_0$ is an optional offset (VARIABLE_OFFSET, default is zero) and $C_i$ are the supplied coefficients. Coefficients that are not supplied default to a value of zero. The VARIABLE argument can be TIME or any internal Expression that evaluates to a scalar. For the latter case, the format of the VARIABLE argument is described in section 2.6.1.

4.12.4 EMISSIVITY = ENCORE_FUNCTION

Parameters

- NAME = STRING
- [EVAL_TYPE = STRING]

Example

Emissivity = Encore_Function Name=My_Function

Description

Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```plaintext
Begin String Function My_Function
  Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

4.12.5 EMISSIVITY = USER_FUNCTION

Parameters

- NAME = STRING
- X = STRING

Example

Begin Definition for Function My_Emissivity
  Type is Piecewise Linear
  Begin Values
    100 0.7
    200 0.6
    300 0.5
  End
End

Begin Aria Material Foo
  ...
  Emissivity = User_Function Name=My_Emissivity X=Temperature
  ...
End Aria Material Foo

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A look-up function is used to compute the values of the emissivity as a function of some other variable, i.e. $f(x)$. The function type ("piecewise linear" in the example above) must support the `differentiate()` method for Newton’s method.

Here `NAME` is the name of the user-defined function (My_Emissivity in the example) and `X` is the Aria name of the abscissa variable (TEMPERATURE in the example). Note that `X` is not necessarily the same name as the abscissa variable identified in the user-defined function (T in the example).

### 4.13 ENTHALPY

**ENTHALPY = MODEL** [param$_1$ = val$_1$, param$_2$ = val$_2$ ...]

**Description**  
Specifies a model for the enthalpy of a material.

**Summary**  
Specifies a model for the enthalpy of a material.

**Parent Block(s)**  
ARIA MATERIAL

#### 4.13.1 ENTHALPY = CONSTANT

**Parameters**  
$H = REAL$

**Example**  
Enthalpy = Constant $H=1e-4$

**Description**  
The value is constant in space and time.

#### 4.13.2 ENTHALPY = ENCORE_FUNCTION

**Parameters**  
`NAME = STRING`  
[EVAL_TYPE = STRING]

**Example**  
Enthalpy = Encore_Function Name=My_Function

**Description**  
Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```plaintext
Begin String Function My_Function
  Value is "200 + 1.0*t"
End
```
Then, this function can be used as shown in the example above.
The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for \texttt{NO\_OP} expressions, DOT for \texttt{DT\_OP} expressions and GRADIENT for \texttt{GRAD\_OP} expressions.

4.14 EQUATION OF STATE

\texttt{EQUATION OF STATE = MODEL[\texttt{param}_1 = \texttt{val}_1, \texttt{param}_2 = \texttt{val}_2 \ldots]}

\textbf{Description} \hspace{1cm} Specifies the equation of state for gas dynamics problems.

\textbf{Summary} \hspace{1cm} Specifies the equation of state for gas dynamics problems.

\textbf{Parent Block(s)} ARIA MATERIAL

4.14.1 EQUATION OF STATE = IDEAL_GAS

\textbf{Parameters} \hspace{1cm} \texttt{R = REAL}

\hspace{1cm} \texttt{GAMMA = REAL}

\textbf{Example} \hspace{1cm} Equation of State = Ideal_Gas R=8.314 Gamma=1.4

\textbf{Description} \hspace{1cm} \texttt{R} is the gas constant and \texttt{GAMMA} is the ratio of heat capacities. This model is used for gas dynamics problems where the density is an unknown. In this case, the pressure is given by the ideal gas law,

\[ p = RT\rho \quad (4.33) \]

where \( T \) is the temperature and \( \rho \) is the density. Activating this model supplies several quantities that are related to this equation of state such as the pressure, temperature, and other gas dynamics related quantities.

4.15 HEAT CONDUCTION

\texttt{HEAT CONDUCTION = MODEL[\texttt{param}_1 = \texttt{val}_1, \texttt{param}_2 = \texttt{val}_2 \ldots]}

\textbf{Description} \hspace{1cm} Specifies the material (constitutive) model for the heat conduction (diffusive flux) in the bulk.

\textbf{Summary} \hspace{1cm} Specifies the material (constitutive) model for the heat conduction (diffusive flux) in the bulk.

\textbf{Parent Block(s)} ARIA MATERIAL
4.15.1 HEAT CONDUCTION = BASIC

This is an alias for FOURIERS_LAW.

Example Heat Conduction = Basic

4.15.2 HEAT CONDUCTION = CONVECTED_ENTHALPY

Parameters (none)

Example Heat Conduction = Convected_Enthalpy

Description The heat conduction (flux) $q$ is given by,

$$ q = -h \rho v $$

where $h$ is the enthalpy, $\rho$ is the density and $v$ is the velocity.

4.15.3 HEAT CONDUCTION = FOURIERS_LAW

Parameters (none)

Example Heat Conduction = Fouriers_Law

Description The heat conduction (flux) $q$ is given by Fourier’s Law,

$$ q = -\kappa \nabla T $$

where $\kappa$ is the thermal conductivity and $T$ is the temperature.

4.15.4 HEAT CONDUCTION = GENERALIZED

Parameters (none)

Example Heat Conduction = Generalized

Description The heat conduction (flux) $q$ is given by Fourier’s Law,

$$ q = -\kappa \nabla T $$

where $\kappa$ is the thermal conductivity tensor and $T$ is the temperature.
4.15.5 HEAT CONDUCTION = THERMOELECTRIC

Parameters  (none)

Example  Heat Conduction = Thermoelectric

Description  The heat conduction (flux) \( q \) is given by Fourier’s Law combined with an electric energy flux,

\[
q = -\kappa \nabla T + \alpha T J
\]  \hspace{1cm} (4.37)

where \( \kappa \) is the thermal conductivity, \( \alpha \) is the Seebeck coefficient, \( T \) is the temperature and \( J \) is the thermoelectric current density 4.6.4.

4.16 HEAT OF VAPORIZATION

HEAT OF VAPORIZATION = \textit{MODEL}[\text{param}_1 = \text{val}_1, \text{param}_2 = \text{val}_2 \ldots]

Description  Specifies the heat of vaporization for a material or for a particular species.

Summary  Quantifies the amount of energy consumed during evaporation per unit mass.

Parent Block(s)  ARIA MATERIAL

4.16.1 HEAT OF VAPORIZATION = CONSTANT

Parameters  \( Hv = \textit{REAL} \)  
[\text{SUBINDEX} = \textit{INT}]

Example  HEAT OF VAPORIZATION = CONSTANT SUBINDEX=0 Hv = 1.0  
HEAT OF VAPORIZATION = CONSTANT SUBINDEX=1 Hv = 2.0  
HEAT OF VAPORIZATION = CONSTANT SUBINDEX=3 Hv = 3.14

Description  \( Hv \) is the value of the constant heat of vaporization and \text{SUBINDEX} is the optional species index (used in multicomponent systems).

4.17 HEAT TRANSFER COEFFICIENT

HEAT TRANSFER COEFFICIENT = \textit{MODEL}[\text{param}_1 = \text{val}_1, \text{param}_2 = \text{val}_2 \ldots]

Description  Specifies the material model for the heat transfer coefficient.

Summary  Specifies the material model for the heat transfer coefficient.

Parent Block(s)  ARIA MATERIAL
4.17.1 HEAT TRANSFER COEFFICIENT = CALORE_USER_SUB

Parameters

- NAME = STRING
- TYPE = STRING
- [MULTIPLIER = REAL]
- [NR = INT]
- [R0 = INT]
- [R1 = INT(etc.)]
- [NI = INT]
- [I0 = INT]
- [I1 = INT(etc.)]

Example

Heat Transfer Coefficient = Calore_User_Sub name=w80afftuser type=element NR=2 R0=1000 R1=0.7

Description

NAME is the name of the user subroutine was registered with. NR and NI are the numbers of real and integer parameters, respectively. Each real (int) parameter is supplied by the Rn (In) parameter. Note: the parameters use zero based counting. The TYPE parameter denotes the API the user subroutine uses; currently Aria only supports types ELEMENT and NODE.

The optional MULTIPLIER parameter (defaults to 1.0) can be supplied to scale the output of the user subroutine, i.e., the flux \( f \) will be \( f = m f_{user} \) where \( m \) is the multiplier and \( f_{user} \) is the output of the user subroutine.

See section 9.2.2 for a more complete example of a Calore user subroutine.

4.17.2 HEAT TRANSFER COEFFICIENT = CONSTANT

Parameters

- H = REAL

Example

Heat Transfer Coefficient = Constant h = 0.8

Description

H is the value of the constant heat transfer coefficient.

4.17.3 HEAT TRANSFER COEFFICIENT = ENCORE_FUNCTION

Parameters

- NAME = STRING
- [EVAL_TYPE = STRING]

Example

Heat Transfer Coefficient = Encore_Function Name=My_Function

Description

Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```plaintext
Begin String Function My_Function
    Value is "200 + 1.0*t"
End
```
Then, this function can be used as shown in the example above. The optional argument `EVAL_TYPE` can be used to select the particular function on the Encore function object. Valid options include `VALUE`, `DOT`, `GRADIENT`, `FLUX` and `STRESS`. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – `VALUE` for `NO_OP` expressions, `DOT` for `DT_OP` expressions and `GRADIENT` for `GRAD_OP` expressions.

### 4.17.4 HEAT TRANSFER COEFFICIENT = POLYNOMIAL

**Parameters**

- `VARIABLE` = `STRING`
- `ORDER` = `INT`
  - `[VARIABLE_OFFSET = `REAL`]`
  - `[C0 = `REAL`]`
  - `[C1 = `REAL`]`
  - ...
  - `[CN = `REAL`]`

**Example**

Heat Transfer Coefficient = Polynomial Variable=Temperature Order=1 C0=401.0 C1=88.5

**Description**

Arbitrary order polynomial function of a specified scalar variable.

\[ h = \sum_{i=0}^{N} C_i (X + X_o)^i \]  

Here, \( N \) is the order of the polynomial provided by the `ORDER` parameter and \( X \) is the variable supplied by the `VARIABLE` parameter, \( X_o \) is an optional offset (`VARIABLE_OFFSET`, default is zero) and \( C_i \) are the supplied coefficients. Coefficients that are not supplied default to a value of zero. The `VARIABLE` argument can be `TIME` or any internal Expression that evaluates to a scalar. For the latter case, the format of the `VARIABLE` argument is described in section 2.6.1.

### 4.17.5 HEAT TRANSFER COEFFICIENT = USER_FUNCTION

**Parameters**

- `NAME` = `STRING`
- `X` = `STRING`

**Example**

Begin Definition for Function My_Heat_Transfer_Coefficient

  Type is Piecewise Linear
  Begin Values
    100  0.7
    200  0.6
    300  0.5
  End

End

Begin Aria Material Foo

  ... Heat Transfer Coefficient = User_Function Name=My_Heat_Transfer_Coefficient X=Temperature ...

End Aria Material Foo
A look-up function is used to compute the values of the heat transfer coefficient as a function of some other variable, i.e. \( f(x) \). The function type (“piecewise linear” in the example above) must support the `differentiate()` method for Newton’s method.

Here `NAME` is the name of the user-defined function (`My_Heat_Transfer_Coefficient` in the example) and \( X \) is the Aria name of the abscissa variable (`TEMPERATURE` in the example). Note that \( X \) is not necessarily the same name as the abscissa variable identified in the user-defined function (\( T \) in the example).

### 4.18 INTRINSIC PERMEABILITY

**INTRINSIC PERMEABILITY = MODEL[param\_1 = val\_1, param\_2 = val\_2 ...]**

**Description**

Specifies the material model for the intrinsic permeability tensor for porous flow in Darcy’s Law.

**Summary**

Specifies the material model for the intrinsic permeability tensor for porous flow in Darcy’s Law. In general, the permeability may be nonisotropic in porous media. In that case, Darcy’s law may be written as,

\[
\rho v_d = f = -\frac{k_r}{\mu} K \cdot (\nabla P - \rho g)
\]  

(4.39)

where \( K \) is the intrinsic permeability tensor, \( \rho \) is the density, \( v_d \) is the Darcy velocity, \( f \) is the mass flux, \( k_r \) is the relative permeability, \( P \) is pressure and \( g \) is gravity.

**Parent Block(s)**

ARIA MATERIAL

#### 4.18.1 INTRINSIC PERMEABILITY = CONSTANT

**Parameters**

- \( [XX = REAL] \)
- \( [XY = REAL] \)
- \( [XZ = REAL] \)
- \( [YX = REAL] \)
- \( [YY = REAL] \)
- \( [YZ = REAL] \)
- \( [ZX = REAL] \)
- \( [ZY = REAL] \)
- \( [ZZ = REAL] \)

**Example**

Intrinsic Permeability = Constant XX=1 YY=2 ZZ=1

**Description**

All components default to zero and all values are constant in space and time.

### 4.19 INTERNAL ENERGY

**INTERNAL ENERGY = MODEL[param\_1 = val\_1, param\_2 = val\_2 ...]**

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Description  Specifies a model for the internal energy of a material.

Summary  Specifies a model for the internal of a material.

Parent Block(s)  ARIA MATERIAL

4.19.1  INTERNAL ENERGY = ENCORE_FUNCTION

Parameters  
- NAME = STRING
  - [EVAL_TYPE = STRING]

Example  Internal Energy = Encore_Function Name=My_Function

Description  Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```
Begin String Function My_Function
  Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

4.19.2  INTERNAL ENERGY = GAS_PHASE

Parameters  (none)

Example  Internal Energy = Gas_Phase

Description  The internal energy $e$ is computed using thermodynamic relation

$$e = h - P/\rho$$  \hspace{1cm}(4.40)$$

where $h$ is the enthalpy, $P$ is the (partial) pressure and $\rho$ is the density.

4.20  LEVEL SET HEAVISIDE

LEVEL SET HEAVISIDE = MODEL[param1 = val1, param2 = val2 ...]
Description  Specifies the functional form of the Heaviside function used with level set algorithms. This also implies the Dirac delta function used for level set algorithms.

Summary  Specifies the functional form of the Heaviside function used with level set algorithms. This also implies the Dirac delta function used for level set algorithms.

Parent Block(s)  ARIA MATERIAL

4.20.1 LEVEL SET HEAVISIDE = SMOOTH

Parameters  (none)

Example  LEVEL SET HEAVISIDE = SMOOTH

Description  The Heaviside function in this case is given as

\[ H(f) = \frac{1}{2} [1 + f/w + \sin(\pi f/w) / \pi] \]  (4.41)

Here \( f \) is the level set distance function and \( w \) is half of the level set width (see 4.21).

4.21 LEVEL SET WIDTH

LEVEL SET WIDTH = MODEL[param_1 = val_1, param_2 = val_2 ...]

Description  Specifies the total width of the level set interface. Half of this width falls on the positive side of the zero level set and half falls on the negative side.

Summary  Specifies the total width of the level set interface. Half of this width falls on the positive side of the zero level set and half falls on the negative side.

Parent Block(s)  ARIA MATERIAL

4.21.1 LEVEL SET WIDTH = CONSTANT

Parameters  WIDTH = REAL

Example  LEVEL SET WIDTH = CONSTANT WIDTH=0.1

Description  This is what you’d expect it to be – a uniform constant everywhere for all time.
4.22  LUBRICATION HEIGHT LOWER

LUBRICATION HEIGHT LOWER = MODEL[\text{param}_1 = \text{val}_1, \text{param}_2 = \text{val}_2 \ldots]

Description  Specifies the material model for the lower height function for the lubrication equation.

Summary  Specifies the material model for the lower height function for the lubrication equation.

Parent Block(s)  ARIA MATERIAL

4.22.1  LUBRICATION HEIGHT LOWER = CONSTANT

Parameters  \texttt{H\_LOWER} = \texttt{REAL}

Example  LUBRICATION HEIGHT LOWER = CONSTANT \texttt{H\_LOWER} = 1.0

Description  \texttt{H\_LOWER} is the constant value.

4.23  LUBRICATION HEIGHT UPPER

LUBRICATION HEIGHT UPPER = MODEL[\text{param}_1 = \text{val}_1, \text{param}_2 = \text{val}_2 \ldots]

Description  Specifies the material model for the upper height function for the lubrication equation.

Summary  Specifies the material model for the upper height function for the lubrication equation.

Parent Block(s)  ARIA MATERIAL

4.23.1  LUBRICATION HEIGHT UPPER = CONSTANT

Parameters  \texttt{H\_UPPER} = \texttt{REAL}

Example  LUBRICATION HEIGHT UPPER = CONSTANT \texttt{H\_UPPER} = 1.0

Description  \texttt{H\_UPPER} is the constant value.
4.24 LUBRICATION K

LUBRICATION K = MODEL [param_1 = val_1, param_2 = val_2 ...]

Description Specifies the “k” model for the lubrication equation.

Summary In the standard derivation of Reynolds’ lubrication theory, the flow is assumed to be laminar and the constant “12” appears in the equation. However, there are extensions to the standard lubrication theory that allow studying turbulent flow by replacing the “12” with a model for turbulent mixing, a scalar that we call $K$.

If this line is omitted, the value of $K = 12$ is assumed. However, other forms for this constant $K$ may be used.

Parent Block(s) ARIA MATERIAL

4.24.1 LUBRICATION K = CONSTANT

Parameters $K = \text{REAL}$

Example LUBRICATION K = CONSTANT $K = 12.0$

Description $K$ is the constant value. Defaults to $K = 12$ if this line is omitted.

4.24.2 LUBRICATION K = PRANDTL_MIXING

Parameters (none)

Example LUBRICATION K = PRANDTL_MIXING

Description This model calculates the turbulent mixing viscosity based on the standard Prandtl mixing theory. First, the Reynolds’ number is calculated using

$$\text{Re} = \frac{h \rho \sqrt{V_{\text{UPPER}} - V_{\text{LOWER}}}}{\mu}.$$  \hspace{1cm} (4.42)

Then, the value of $K$ is calculated with

$$K = \begin{cases} 12 & \text{Re} < 2000 \\ 0.03(2\text{Re})^{0.75} & \text{Re} > 2000 \end{cases}.$$ \hspace{1cm} (4.43)

4.25 LUBRICATION VELOCITY LOWER

LUBRICATION VELOCITY LOWER = MODEL [param_1 = val_1, param_2 = val_2 ...]
Description Specifies the material model for the velocity of the lower boundary for the lubrication equation.

Summary Specifies the material model for the velocity of the lower boundary for the lubrication equation.

Parent Block(s) ARIA MATERIAL

### 4.25.1 LUBRICATION VELOCITY LOWER = CONSTANT

Parameters \( V_{\text{LOWER}} = \text{REAL} \)

Example \( \text{LUBRICATION VELOCITY LOWER} = \text{CONSTANT} \ V_{\text{LOWER}} = 0.0 \)

Description \( V_{\text{LOWER}} \) is the constant value.

### 4.26 LUBRICATION VELOCITY UPPER

\[
\text{LUBRICATION VELOCITY UPPER} = \text{MODEL}[\text{param}_1 = \text{val}_1, \text{param}_2 = \text{val}_2 \ldots]
\]

Description Specifies the material model for the velocity of the upper boundary for the lubrication equation.

Summary Specifies the material model for the velocity of the upper boundary for the lubrication equation.

Parent Block(s) ARIA MATERIAL

### 4.26.1 LUBRICATION VELOCITY UPPER = CONSTANT

Parameters \( V_{\text{UPPER}} = \text{REAL} \)

Example \( \text{LUBRICATION VELOCITY UPPER} = \text{CONSTANT} \ V_{\text{UPPER}} = 0.0 \)

Description \( V_{\text{UPPER}} \) is the constant value.

### 4.27 MASS FLUX

\[
\text{MASS FLUX} = \text{MODEL}[\text{param}_1 = \text{val}_1, \text{param}_2 = \text{val}_2 \ldots]
\]
Description Specifies a constitutive model for the mass flux for porous flow applications.

Summary Specifies a constitutive model for the mass flux for porous flow applications.

Parent Block(s) ARIA MATERIAL

### 4.27.1 MASS FLUX = DARCY

**Parameters**

- \[GX = \text{REAL}\]
- \[GY = \text{REAL}\]
- \[GZ = \text{REAL}\]

**Example**

Mass Flux = Darcy \( GY=-9.8 \)

**Description**

The macroscopic, convective mass flux in phase \( \beta \), \( \rho \nu \) is obtained from the extended Darcy’s Law,

\[
F = \rho f = \frac{\rho k_r}{\mu} \cdot (\nabla P - \rho g)
\]  
(4.44)

### 4.28 MESH LAMBDA

**MESH LAMBDA = MODEL** \([\text{param}_1 = \text{val}_1, \text{param}_2 = \text{val}_2 \ldots]\)

**Description**

Specifies the material model for the \( \lambda \) Lamé coefficient for the mesh (psuedo-solid) stress tensor.

**Summary**

The solid stress \( T \) is given by

\[
T = \lambda E_{kk} I + 2\mu E - \beta (T - T_{ref}) I
\]  
(4.45)

where \( \lambda \) and \( \mu \) are the Lamé coefficients, \( E = \frac{1}{2} \left( \nabla d + \nabla d^T \right) \) is the deformation tensor, \( \beta \) is the coefficient of thermal stress, \( T \) is temperature and \( T_{ref} \) is the thermal strain reference temperature.

These Lamé coefficients are related to the more standard Young’s modulus, Poisson’s ratio and CTE \( (\alpha) \) as follows:

\[
2\mu = \frac{E}{(1 + \nu)}
\]  
(4.46)

\[
\lambda = \frac{\nu E}{(1 + \nu)(1 - 2\nu)} = 2\mu \frac{\nu}{(1 - 2\nu)}
\]  
(4.47)

\[
\beta = \frac{\alpha E}{(1 - 2\nu)} = \alpha (3\lambda + 2\mu)
\]  
(4.48)

When a user supplies the Young’s modulus, Poisson’s ratio and CTE properties Aria internally converts them into the Lamé coefficients.

Parent Block(s) ARIA MATERIAL
4.28.1 MESH LAMBDA = CONSTANT

Parameters

\[ L = \text{REAL} \]

Example

MESH LAMBDA = CONSTANT \( L = 1.0 \)

Description

\( L \) is the value of the constant \( \lambda \).

4.28.2 MESH LAMBDA = CONVERTED

Parameters

(None)

Example

MESH LAMBDA = Converted

Description

Aria will use Young's modulus and Poisson ratio to compute the Lamé \( \lambda \) coefficient. Supplying the Lamé coefficients is more computationally efficient but perhaps less convenient, especially if the material properties are varying (e.g., temperature dependent in a non-isothermal problem).

4.28.3 MESH LAMBDA = ENCORE_FUNCTION

Parameters

\[ \text{NAME} = \text{STRING} \]
\[ [\text{EVAL_TYPE} = \text{STRING}] \]

Example

MESH LAMBDA = Encore_Function Name=My_Function

Description

Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```
Begin String Function My_Function
  Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument `EVAL_TYPE` can be used to select the particular function on the Encore function object. Valid options include `VALUE`, `DOT`, `GRADIENT`, `FLUX` and `STRESS`. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – `VALUE` for `NO_OP` expressions, `DOT` for `DT_OP` expressions and `GRADIENT` for `GRAD_OP` expressions.

4.29 MESH TWO MU

\[ \text{MESH TWO MU} = \text{MODEL} [\text{param}_1 = \text{val}_1, \text{param}_2 = \text{val}_2 \ldots] \]
Specifies the material model for twice the $\mu$ Lamé coefficient for the mesh (pseudo-solid) stress tensor.

The solid stress $T$ is given by

$$ T = \lambda E_{kk} I + 2\mu E - \beta (T - T_{ref}) I $$

(4.49)

where $\lambda$ and $\mu$ are the Lamé coefficients, $E = \frac{1}{2} (\nabla d + \nabla d^T)$ is the deformation tensor, $\beta$ is the coefficient of thermal stress, $T$ is temperature and $T_{ref}$ is the thermal strain reference temperature.

These Lamé coefficients are related to the more standard Young’s modulus, Poisson’s ratio and CTE ($\alpha$) as follows:

$$ 2\mu = \frac{E}{1 + \nu} $$

(4.50)

$$ \lambda = \frac{\nu E}{(1 + \nu)(1 - 2\nu)} = 2\mu \frac{\nu}{(1 - 2\nu)} $$

(4.51)

$$ \beta = \frac{\alpha E}{(1 - 2\nu)} = \alpha (3\lambda + 2\mu) $$

(4.52)

When a user supplies the Young’s modulus, Poisson’s ratio and CTE properties Aria internally converts them into the Lamé coefficients.

4.29.1 MESH TWO MU = CONSTANT

Parameters

 TWO_MU = REAL

Example

 MESH TWO MU = CONSTANT TWO_MU = 1.0

Description

TWO_MU is the value of $2\mu$.

4.29.2 MESH TWO MU = CONVERTED

Parameters

(None)

Example

 MESH TWO MU = Converted

Description

Aria will use Young’s modulus and Poisson ratio to compute the Lamé $\mu$ coefficient. Supplying the Lamé coefficients is more computationally efficient but perhaps less convenient, especially if the material properties are varying (e.g., temperature dependent in a non-isothermal problem).
### 4.29.3 MESH TWO MU = ENCORE_FUNCTION

**Parameters**

- NAME = *STRING*
  - [EVAL_TYPE = *STRING*]

**Example**

TWO_MU = Encore_Function Name=My Function

**Description**

Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```plaintext
Begin String Function My Function
  Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

### 4.30 MESH STRESS

MESH STRESS = MODEL [param₁ = val₁, param₂ = val₂ ...]

**Description**

Specifies a contribution to the mesh (pseudo-solid) stress tensor. Multiple stresses are combined additively and may be specified by using this line command multiple times.

**Summary**

Specifies a contribution to the mesh (pseudo-solid) stress tensor. The total stress \( T \) is given by

\[
T = \sum_j T_j
\]  \hspace{1cm} (4.53)

**Parent Block(s)**

ARIA MATERIAL

### 4.30.1 MESH STRESS = ISOTHERMAL

**Parameters**

- \( T = \text{REAL} \)
- \( T\_\text{REF} = \text{REAL} \)

**Example**

MESH STRESS = Isothermal \( T=500 \) \( T\_\text{REF}=325 \)
This stress accounts for the mechanical stresses due to thermally induced strains.

\[ T = -\beta (T - T_{\text{ref}}) I \]  \hspace{1cm} (4.54)

where \( \beta \) is the Lamé coefficient of thermal stress (related to the coefficient of thermal expansion, \( \alpha \)), \( T \) is the temperature and \( T_{\text{ref}} \) is the temperature of the undeformed reference state of the mesh (pseudo-solid). This is a specialization of the THERMAL model that uses uniform, fixed temperature and thermal strain reference temperature.

4.30.2 MESH STRESS = LINEAR_ELASTIC

Parameters

\[ \text{REFERENCE\_FRAME} = \text{"MOVING" | "UNDEFORMED"} \]

Example

MESH STRESS = Linear_Elastic Reference_Frame = undeformed

Description

Supplies the linear elasticity stress tensor,

\[ T = \lambda \text{trace} \, E I + 2\mu E \]  \hspace{1cm} (4.55)

where \( \lambda \) and \( \mu \) and the Lamé coefficients and \( E \) is the strain tensor. When the choice of reference frame is “UNDEFORMED” then the strain is computed in the undeformed reference state; this is commonly referred to as small strain theory. When the reference frame is “MOVING” then the strain is computed with respect to the deformed coordinates.

Specifically, the strain tensor is given by

\[ E = \frac{1}{2} (\nabla d + \nabla d^t) \]  \hspace{1cm} (4.56)

where \( d \) is the mesh displacement field. The choice of reference frame determines whether the \( \nabla \) operator is computed in the undeformed or moving reference frames.

4.30.3 MESH STRESS = NEOHOOKEAN_ELASTIC

Parameters

(none)

Example

MESH STRESS = Neohookean_Elastic

Description

Supplies a nonlinear hyperelastic stress of the form,

\[ T = \frac{\mu}{J} (b - I) + \frac{\lambda}{J} \ln J I \]  \hspace{1cm} (4.57)

where \( \lambda \) and \( \mu \) and the Lamé coefficients, \( b \equiv F \cdot F^t \) is the left Cauchy-Green tensor, \( F \) is the deformation gradient and \( J \equiv \det F \). See, e.g., [4] or [9].

4.30.4 MESH STRESS = NONLINEAR_ELASTIC

Parameters

(none)
**Example**  \( \text{MESH STRESS} = \text{Nonlinear\_Elastic} \)

**Description**  Supplies a nonlinear elastic stress,

\[
T = \lambda \text{trace} \ E I + 2\mu E
\]  \hspace{1cm} (4.58)

where \( \lambda \) and \( \mu \) and the Lamé coefficients and \( E \) is the strain tensor. The particular choice of strain tensor chosen depends on where the configuration (reference frame) which is set via the MESH MOTION command line. See section 5.16 for more information. When the MESH MOTION is set to \text{REFERENCE\_FRAME} then the Green strain is used. Otherwise, the Almansi strain is used.

### 4.30.5 MESH STRESS = RESIDUAL

**Parameters**

- \[ \text{SXX | SX = REAL} \]
- \[ \text{SYY | SY = REAL} \]
- \[ \text{SZZ | SZ = REAL} \]
- \[ \text{SXY = REAL} \]
- \[ \text{SXZ = REAL} \]
- \[ \text{SYZ = REAL} \]

**Example**  \( \text{SOLID STRESS} = \text{Residual Sxx}=0.02 \text{ Syy}=0.02 \)

**Description**  This stress accounts for the initial residual stress in a solid that is constant and uniform everywhere. The components of the residual stress tensor are supplied by the (up-to) six components \( \text{SXX}, \text{SYY}, \text{SZZ}, \text{SXY}, \text{SXZ}, \text{SYZ} \).

This is directly analogous to the \text{ISTRESS} condition in ANSYS. To that end, the diagonal components can be specified as either \text{SXX} or \text{SX} etc.

### 4.30.6 MESH STRESS = THERMAL

**Parameters**  (none)

**Example**  \( \text{MESH STRESS} = \text{Thermal} \)

**Description**  This stress accounts for the mechanical stresses due to thermally induced strains.

\[
T = -\beta (T - T_{ref}) I
\]  \hspace{1cm} (4.59)

where \( \beta \) is the Lamé coefficient of thermal stress (related to the coefficient of thermal expansion, \( \alpha \)), \( T \) is the temperature and \( T_{ref} \) is the temperature of the undeformed reference state of the mesh (pseudo-solid).

### 4.30.7 MESH STRESS = LAME

**Parameters**  \( \text{MODEL} = \text{STRING} \)
Example MESH STRESS = Lame Model = Neo_Hookean

Description This stress interfaces with the Lame constitutive model library used by Sierra/SolidMechanics. As such, it allows the use of many of Lame’s constitutive models. Sierra/Multimechanics prepares the kinematic quantities, and Lame returns the Cauchy stress, which is integrated in the MESH equation.

Material parameters for the constitutive models are handled by an input block consistent with the Sierra/SolidMechanics formatting. For documentation of the available constitutive models and the required parameter input, please see the Sierra/SolidMechanics documentation.

Currently this interface is instrumented to support hyperelastic constitutive models, and is restricted to the following models:

- NEO_HOOKEAN
- THERMALBATTERYSEPARATOR
- SOILFOAMPHASETRANSITION

4.31 MOLECULAR WEIGHT

MOLECULAR WEIGHT = MODEL [param1 = val1, param2 = val2 ...]

Description Specifies the molecular weight for a species.

Summary Specifies the molecular weight for a species.

Parent Block(s) ARIA MATERIAL

4.31.1 MOLECULAR WEIGHT = CONSTANT

Parameters M = REAL
[SUBINDEX = INT]

Example Molecular Weight = Constant Subindex=1 M = 17.0
Molecular Weight = Constant Subindex=2 M = 23.0
Molecular Weight = Constant Subindex=5 M = 34.0

Description M is the value of the molecular weight.
SUBINDEX is the species subindex.

4.31.2 MOLECULAR WEIGHT = ENCORE_FUNCTION

Parameters NAME = STRING
[EVAL_TYPE = STRING]

Example Molecular Weight = Encore_Function Name=My_Function
Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```
Begin String Function My_Function
  Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument `EVAL_TYPE` can be used to select the particular function on the Encore function object. Valid options include `VALUE`, `DOT`, `GRADIENT`, `FLUX` and `STRESS`. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – `VALUE` for `NO_OP` expressions, `DOT` for `DT_OP` expressions and `GRADIENT` for `GRAD_OP` expressions.

### 4.32 MOMENTUM STRESS

**MOMENTUM STRESS = ** `MODEL`[param1 = val1, param2 = val2 ...]

**Description**
Specifies a contribution to the fluid stress tensor. Multiple stresses are combined additively and may be specified by using this line command multiple times.

**Summary**
Specifies a contribution to the fluid momentum stress tensor. The total stress \( \mathbf{T} \) is given by

\[
\mathbf{T} = \sum_j T_j
\]  

(4.60)

**Parent Block(s)** ARIA MATERIAL

#### 4.32.1 MOMENTUM STRESS = LS_CAPILLARY

**Parameters** (none)

**Example**
MOMENTUM STRESS = Newtonian
MOMENTUM STRESS = LS_Capillary

**Description**
This adds the capillary boundary condition contributions in the vicinity of the level set interface.

\[
\mathbf{T} = \sigma \delta(F)(\mathbf{I} - \mathbf{N}\mathbf{N})
\]  

(4.61)

where \( \sigma \) is the surface tension, \( \delta(F) \) is the level set delta function, \( F \) is the level set distance function and \( \mathbf{N} \) is the level set normal field.
4.32.2 MOMENTUM STRESS = INCOMPRESSIBLE_NEWTONIAN

Parameters

[PRESSURE = STRING]

Example

MOMENTUM STRESS = Incompressible_Newtonian

Description

Supplies the incompressible Newtonian stress tensor,

\[ T = -pI + \mu (\nabla v + \nabla v^t) \]  \hspace{1cm} (4.62)

where \( \mu \) is the fluid viscosity, \( p \) is the pressure and \( v \) is the fluid velocity. The viscosity \( \mu \) is provided with the viscosity line command as described in section 4.56.

Note that this model does not include stress contributions that are proportional to the divergence of the velocity. To incorporate those contributions, use the NEWTONIAN_DILATIONAL stress model in addition to this model, or use the FORMAL_NEWTONIAN stress model instead of this model.

By default, the pressure field used by this model is PRESSURE with the same subindex as the model. The optional parameter PRESSURE can be used to specify an alternate pressure field. This is useful for running problems with a continuous velocity field but a discontinuous pressure field.

4.32.3 MOMENTUM STRESS = FORMAL_NEWTONIAN

Parameters

(none)

Example

MOMENTUM STRESS = Formal_Newtonian

Description

Supplies the complete Newtonian stress tensor,

\[ T = -pI + \mu (\nabla v + \nabla v^t) + \left( \kappa - \frac{2}{3} \mu \right) \nabla \cdot v I \]  \hspace{1cm} (4.63)

where \( \mu \) is the fluid viscosity, \( p \) is the pressure and \( v \) is the fluid velocity. The viscosity \( \mu \) is provided with the viscosity line command as described in section 4.56. The bulk viscosity \( \kappa \) is provided with the bulk viscosity line command as described in section 4.4.

4.32.4 MOMENTUM STRESS = NEWTONIAN_DILATIONAL

Parameters

(none)

Example

MOMENTUM STRESS = Newtonian_Dilational

Description

Adds the dilational stress contribution for Newtonian fluids,

\[ T = \left( \kappa - \frac{2}{3} \mu \right) \nabla \cdot v I \]  \hspace{1cm} (4.64)
where $\kappa$ is the bulk viscosity of the fluid, $\mu$ is the fluid (dynamic) viscosity and $v$ is the fluid velocity. The viscosity $\mu$ is provided with the viscosity line command as described in section 4.56. The bulk viscosity $\kappa$ is provided with the bulk viscosity line command as described in section 4.4.

See, also, the NEWTONIAN momentum stress model.

### 4.32.5 MOMENTUM STRESS = NEWTONIAN_VISCOUS

**Parameters**

(none)

**Example**

MOMENTUM STRESS = Newtonian_Viscous

**Description**

Supplies only the viscous contribution of the Newtonian stress tensor,

$$T = \mu (\nabla v + \nabla v^t)$$  \hspace{1cm} (4.65)

where $\mu$ is the fluid viscosity and $v$ is the fluid velocity. The viscosity $\mu$ is provided with the viscosity line command as described in section 4.56.

### 4.32.6 MOMENTUM STRESS = NEWTONIAN_PRESSURE

**Parameters**

(none)

**Example**

MOMENTUM STRESS = Newtonian_Pressure

**Description**

Supplies only the pressure contribution of the Newtonian stress tensor,

$$T = -pI$$  \hspace{1cm} (4.66)

where $p$ is the pressure.

### 4.32.7 MOMENTUM STRESS = MAXWELL

**Parameters**

(none)

**Example**

MOMENTUM STRESS = Maxwell

**Description**

Includes the force due to an electric field (neglecting magnetic effects) due to the Maxwell stress tensor,

$$T_M = \varepsilon \left( EE^T - \frac{1}{2}(E \cdot E)I \right)$$  \hspace{1cm} (4.67)

where $\varepsilon$ is the electrical permittivity and $E$ is the electric field. The divergence of the Maxwell stress tensor is a force applied to the momentum equation,

$$\nabla \cdot T_M = \rho_e E - \frac{1}{2}(E \cdot E) \nabla \varepsilon,$$  \hspace{1cm} (4.68)

where $\rho_e$ is the volumetric charge density.
4.33  POISSONS RATIO

POISSONS RATIO = MODEL [param1 = val1, param2 = val2 ...]

Description  Specifies the material model for the Poisson's ratio.

Summary  The solid stress \( T \) is given by

\[
T = \lambda E_{kk} I + 2\mu E - \beta (T - T_{ref}) I
\]

where \( \lambda \) and \( \mu \) are the Lamé coefficients, \( E = \frac{1}{2} (\nabla d + \nabla d^T) \) is the deformation tensor, \( \beta \) is the coefficient of thermal stress, \( T \) is temperature and \( T_{ref} \) is the thermal strain reference temperature.

These Lamé coefficients are related to the more standard Young’s modulus, Poisson’s ratio and CTE \( (\alpha) \) as follows:

\[
2\mu = \frac{E}{(1 + \nu)} \quad (4.70)
\]

\[
\lambda = \frac{\nu E}{(1 + \nu)(1 - 2\nu)} = 2\mu \frac{\nu}{(1 - 2\nu)} \quad (4.71)
\]

\[
\beta = \frac{\alpha E}{(1 - 2\nu)} = \alpha (3\lambda + 2\mu) \quad (4.72)
\]

When a user supplies the Young’s modulus, Poisson’s ratio and CTE properties Aria internally converts them into the Lamé coefficients.

Supplying the Lamé coefficients is more computationally efficient but perhaps less convenient, especially if the material properties are varying (e.g., temperature dependent in a non-isothermal problem).

Parent Block(s)  ARIA MATERIAL

4.33.1  POISSONS RATIO = CONSTANT

Parameters  PR = REAL

Example  POISSONS RATIO = CONSTANT PR = 1.0

Description  PR is the value of the constant Poisson’s ratio.

4.33.2  POISSONS RATIO = ENCORE_FUNCTION

Parameters  NAME = STRING
[EVAT_TYPE = STRING]

Example  POISSONS RATIO = Encore_Function Name=My_Function
Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```
Begin String Function My_Function
  Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument `EVAL_TYPE` can be used to select the particular function on the Encore function object. Valid options include `VALUE`, `DOT`, `GRADIENT`, `FLUX` and `STRESS`. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – `VALUE` for `NO_OP` expressions, `DOT` for `DT_OP` expressions and `GRADIENT` for `GRAD_OP` expressions.

### 4.34 POROSITY

POROSITY = *MODEL* [param\(_1\) = val\(_1\), param\(_2\) = val\(_2\) ...]

**Description**

Specifies the material model for the porosity for porous flow applications.

**Summary**

Specifies the material model for the porosity for porous flow applications.

**Parent Block(s)**

ARIA MATERIAL

#### 4.34.1 POROSITY = CONSTANT

**Parameters**

PHI = *REAL*

**Example**

Porosity = Constant PHI=0.1

**Description**

The value is constant in space and time.

#### 4.34.2 POROSITY = DEFORMING

**Parameters**

PHI\(_{INIT}\) = *REAL*

**Example**

Porosity = Deforming

**Description**

The porosity evolves from an initial value in accordance with a deforming media.

\[
\phi = 1 \left( 1 - \phi_0 \right) e^{\epsilon v}
\]  

(4.73)
where $\epsilon_v$ is the volume strain,

\[ \epsilon_v = \nabla \cdot d \]  \hfill (4.74)

where $d$ is the total mesh displacement field. Here $\phi_0$ is the initial porosity value given by the PHI_INIT parameter.

4.34.3 POROSITY = ENCORE_FUNCTION

Parameters

- NAME = STRING
- [EVAL_TYPE = STRING]

Example

Porosity = Encore_Function Name=My_Function

Description

Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```
Begin String Function My_Function
  Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

4.34.4 POROSITY = POLYNOMIAL

Parameters

- VARIABLE = STRING
- ORDER = INT
- [VARIABLE_OFFSET = REAL]
- [C0 = REAL]
- [C1 = REAL]
- ...  \[ CN = REAL \]

Example

Porosity= Polynomial Variable=Temperature Order=1 C0=401.0 C1=88.5

Description

Arbitrary order polynomial function of a specified scalar variable.

\[ \phi = \sum_{i=0}^{N} C_i (X + X_o)^i \]  \hfill (4.75)

Here, $N$ is the order of the polynomial provided by the ORDER parameter and $X$ is the variable supplied by the VARIABLE parameter, $X_o$ is an optional offset (VARIABLE_OFFSET, default is zero) and $C_i$ are the supplied coefficients. Coefficients that are not supplied default to a value of zero. The VARIABLE argument can be TIME or any internal Expression that evaluates to a scalar. For the latter case, the format of the VARIABLE argument is described in section 2.6.1.
4.34.5  POROSITY = ROCK_COMPRESSIBLE

Parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>REF_POROSITY</td>
<td>REAL</td>
</tr>
<tr>
<td>REF_PRESSURE</td>
<td>REAL</td>
</tr>
<tr>
<td>CR</td>
<td>REAL</td>
</tr>
</tbody>
</table>

Example

Porosity = Rock_Compressible Ref_Porosity=0.1 Ref_Pressure=101325 Cr=-1e-7

Description

The compressibility of a porous material is defined as

\[ C_r = \frac{1}{\phi} \frac{\partial \phi}{\partial P} \]  (4.76)

where \( \phi \) is the porosity and \( P \) is pressure. Using this definition, we define an approximate model for the porosity as a function of pressure,

\[ \phi = \phi_{ref} \left(1 + C_r (P - P_{ref})\right) \]  (4.77)

where \( \phi_{ref} \) is given by the REF_POROSITY parameter and is the porosity at the reference pressure \( P_{ref} \) which is given by the REF_PRESSURE parameter. The compressibility \( C_r \) is given by the CR parameter.

4.35  RADIATION FORM FACTOR

RADIATION FORM FACTOR = MODEL[param1 = val1, param2 = val2 ...]

Description

Specifies the material model for the radiative heat transfer form factor used in the GENERALIZED_RAD boundary condition.

Summary

The form factor represents the fraction of the radiation leaving a surface which interacts with a body at another temperature. Thus the form factor should be less than or equal to unity.

Parent Block(s)  ARIA MATERIAL

4.35.1  RADIATION FORM FACTOR = CALORE_USER_SUB

Parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>NAME</td>
<td>STRING</td>
</tr>
<tr>
<td>TYPE</td>
<td>STRING</td>
</tr>
<tr>
<td>[MULTIPLIER = REAL]</td>
<td></td>
</tr>
<tr>
<td>[NR = INT]</td>
<td></td>
</tr>
<tr>
<td>[R0 = INT]</td>
<td></td>
</tr>
<tr>
<td>[R1 = INT(etc.)]</td>
<td></td>
</tr>
<tr>
<td>[NI = INT]</td>
<td></td>
</tr>
<tr>
<td>[I0 = INT]</td>
<td></td>
</tr>
<tr>
<td>[I1 = INT(etc.)]</td>
<td></td>
</tr>
</tbody>
</table>

Example

Radiation Form Factor = Calore_User_Sub name=w80afftuser type=element NR=2 R0=1000 R1=0.7
NAME is the name of the user subroutine was registered with. NR and NI are the numbers of real and integer parameters, respectively. Each real (int) parameter is supplied by the \( R_n (I_n) \) parameter. Note: the parameters use zero based counting. The TYPE parameter denotes the API the user subroutine uses; currently Aria only supports types ELEMENT and NODE.

The optional MULTIPLIER parameter (defaults to 1.0) can be supplied to scale the output of the user subroutine, i.e., the flux \( f \) will be \( f = mf_{user} \) where \( m \) is the multiplier and \( f_{user} \) is the output of the user subroutine.

See section 9.2.2 for a more complete example of a Calore user subroutine.

### 4.35.2 RADIATION FORM FACTOR = CONSTANT

**Parameters**

\[ F = \text{REAL} \]

**Example**

Radiation Form Factor = Constant \( F = 0.8 \)

**Description**

\( F \) is the value of the constant radiation form factor.

### 4.35.3 RADIATION FORM FACTOR = POLYNOMIAL

**Parameters**

\[ \text{VARIABLE} = \text{STRING} \]
\[ \text{ORDER} = \text{INT} \]
\[ \text{[VARIABLE_OFFSET} = \text{REAL}] \]
\[ \text{[C0} = \text{REAL}] \]
\[ \text{[C1} = \text{REAL}] \]
\[ \ldots \]
\[ \text{[CN} = \text{REAL}] \]

**Example**

Radiation Form Factor = Polynomial

\( \text{Variable=} \text{Temperature} \)

\( \text{Order=} 1 \)

\( \text{C0=} 401.0 \)

\( \text{C1=} 88.5 \)

**Description**

Arbitrary order polynomial function of a specified scalar variable.

\[
F = \sum_{i=0}^{N} C_i (X + X_o)^i
\]

(4.78)

Here, \( N \) is the order of the polynomial provided by the ORDER parameter and \( X \) is the variable supplied by the VARIABLE parameter, \( X_o \) is an optional offset (VARIABLE_OFFSET, default is zero) and \( C_i \) are the supplied coefficients. Coefficients that are not supplied default to a value of zero. The VARIABLE argument can be TIME or any internal Expression that evaluates to a scalar. For the latter case, the format of the VARIABLE argument is described in section 2.6.1.

### 4.35.4 RADIATION FORM FACTOR = ENCORE_FUNCTION

**Parameters**

\[ \text{NAME} = \text{STRING} \]
\[ \text{[EVAL_TYPE} = \text{STRING}] \]

**Example**

Radiation Form Factor = Encore_Function

\( \text{Name=} \text{My\_Function} \)
Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```
Begin String Function My_Function
    Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

### 4.35.5 RADIATION FORM FACTOR = USER_FUNCTION

**Parameters**

- NAME = STRING
- X = STRING

**Example**

```plaintext
Begin Definition for Function My_Radiation_Form_Factor
    Type is Piecewise Linear
    Begin Values
        100 0.7
        200 0.6
        300 0.5
    End
End
```

```plaintext
Begin Aria Material Foo
    ... 
    Radiation Form Factor = User_Function Name=My_Radiation_Form_Factor X=Temperature 
    ... 
End Aria Material Foo
```

**Description**

A look-up function is used to compute the values of the form factor as a function of some other variable, i.e. \( f(x) \). The function type (“piecewise linear” in the example above) must support the differentiate() method for Newton’s method.

Here NAME is the name of the user-defined function (My_Radiation_Form_Factor in the example) and X is the Aria name of the abscissa variable (TEMPERATURE in the example). Note that X is not necessarily the same name as the abscissa variable identified in the user-defined function (T in the example).

### 4.36 RADIATIVE CONDUCTIVITY

**Description**

Specifies the material model for the radiative effective thermal conductivity.
Summary specifies the material model for the radiative portion of thermal conductivity that appears in the diffusion term of the energy equation for temperature.

Parent Block(s) ARIA MATERIAL

4.36.1 RADIATIVE CONDUCTIVITY = CONSTANT

Parameters \( K_{\text{rad}} = \text{REAL} \)

Example RADIATIVE CONDUCTIVITY = CONSTANT \( K_{\text{rad}} = 1.0 \)

Description \( K_{\text{rad}} \) is the value of the constant radiative thermal conductivity.

4.36.2 RADIATIVE CONDUCTIVITY = CHEMEQ_FOAM

Parameters \( K_{c0} = \text{REAL} \)
\( K_{c1} = \text{REAL} \)

Example Radiative Conductivity = ChemEq_Foam \( K_{c0} = 0.00268 \) \( K_{c1} = 0.0 \)

Description For a reacting foam defined in ChemEq, this model uses

\[
k_{rad} = \frac{K_{c0}}{K_{c1} + \sum Y_i \sigma T^3}
\]

where the denominator includes the sum of the mass fractions of the solid species of the foam. This uses the species phase definition from the ChemEq block to determine which species are solid. This model can only be used in a material block which also contains a ChemEq definition.

This uses expressions for the mass fractions, so its material block must also include “Mass_Fraction of X = From_ChemEq” for each solid species.

The GLOBAL CONSTANTS block must include include the STEFAN-BOLTZMANN CONSTANT entry to provide a value of \( \sigma \).

4.36.3 RADIATIVE CONDUCTIVITY = OPTICALLY_THICK

Parameters \( K = \text{REAL} \)
\( N = \text{REAL} \)
\( \beta_{\text{R}} = \text{REAL} \)

Example RADIATIVE CONDUCTIVITY = OPTICALLY_THICK \( N = 2.1 \) \( \beta_{\text{R}} = 1.0 \times 10^{-5} \)
Description
Thermal conductivity used to model effective radiative conductivity diffusion in the optically-thick limit. Constant values for index of refraction \( n \), and Rosseland-mean extinction coefficient \( \beta_R \) must be provided on the input line.

\[
k_{rad} = \frac{16\pi n^2}{3\beta_R} T^3
\] (4.80)

The GLOBAL CONSTANTS block must include include the STEFAN-BOLTZMANN CONSTANT entry to provide a value of \( \sigma \).

4.36.4 RADIATIVE CONDUCTIVITY = ENCORE_FUNCTION

Parameters
- NAME = STRING
  [EVAL_TYPE = STRING]

Example
Radiative Conductivity = Encore_Function Name=My_Function

Description
Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

Begin String Function My_Function
  Value is "200 + 1.0*t"
End

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

4.36.5 RADIATIVE CONDUCTIVITY = POLYNOMIAL

Parameters
- VARIABLE = STRING
- ORDER = INT
  [VARIABLE_OFFSET = REAL]
  [C0 = REAL]
  [C1 = REAL]
  ...
  [CN = REAL]

Example
Radiative Conductivity= Polynomial Variable=Temperature Order=1 C0=401.0 C1=88.5

Description
Arbitrary order polynomial function of a specified scalar variable.

\[
\kappa = \sum_{i=0}^{N} C_i (X + X_0)^i
\] (4.81)
Here, \( N \) is the order of the polynomial provided by the \texttt{ORDER} parameter and \( X \) is the variable supplied by the \texttt{VARIABLE} parameter, \( X_0 \) is an optional offset (\texttt{VARIABLE_OFFSET}, default is zero) and \( C_i \) are the supplied coefficients. Coefficients that are not supplied default to a value of zero. The \texttt{VARIABLE} argument can be \texttt{TIME} or any internal Expression that evaluates to a scalar. For the latter case, the format of the \texttt{VARIABLE} argument is described in section 2.6.1.

4.36.6 \textbf{RADIATIVE CONDUCTIVITY = USER\_FUNCTION}

**Parameters**
- \texttt{NAME = STRING}
- \texttt{X = STRING}

**Example**
```plaintext
begin definition for function SI_Kr
  type is piecewise linear
  begin values
    20.0  5.50e7
    100.0 4.60e7
    ...
    800.0 1.30e7
    2000.0 1.30e7
  end values
end definition for function SI_Kr

Begin Aria Material Foo
  ...
  Radiative Conductivity = User_Function Name=SI_Kr X=Temperature
  ...
End Aria Material Foo
```

**Description**
A look-up function is used to compute the values of the thermal conductivity as a function of some other variable, i.e. \( f(x) \). The function type ("piecewise linear" in the example above) must support the \texttt{differentiate()} method for Newton’s method.

Here \texttt{NAME} is the name of the user-defined function for (\texttt{CONDUCTIVITY\_DATA}, \texttt{SI\_K}, in the example) and \texttt{X} is the Aria name of the abscissa variable (\texttt{TEMPERATURE} in the example). Note that \texttt{X} is not necessarily the same name as the optional abscissa variable identified in the user-defined function.

4.37 \textbf{RELATIVE PERMEABILITY}

```plaintext
RELATIVE PERMEABILITY = MODEL[\texttt{param}_1 = \texttt{val}_1, \texttt{param}_2 = \texttt{val}_2 \ldots]
```

**Description**
Specifies the material model for the relative permeability (scalar) for porous flow in Darcy’s Law.

**Summary**
Specifies the material model for the relative permeability (scalar) for porous flow in Darcy’s Law. In general, the permeability may be nonisotropic in porous media. In that case, Darcy’s law may be written as,

\[
\rho w_d = f = \frac{\rho k_r}{\mu} K \cdot (\nabla P + \rho g)
\]  
(4.82)
where \( K \) is the intrinsic permeability tensor, \( \rho \) is the density, \( \bm{v}_d \) is the Darcy velocity, \( f \) is the mass flux, \( k_r \) is the relative permeability, \( \mu \) is the dynamic viscosity, \( P \) is pressure and \( g \) is gravity.

**Parent Block(s)** ARIA MATERIAL

### 4.37.1 RELATIVE PERMEABILITY = CONSTANT

**Parameters**

\[ K = \text{REAL} \]

**Example**

Relative Permeability = Constant \( K=1e^{-3} \)

**Description**

The value is constant in space and time.

### 4.37.2 RELATIVE PERMEABILITY = ENCORE_FUNCTION

**Parameters**

\[ \text{NAME} = \text{STRING} \]

\[ \text{EVAL_TYPE} = \text{STRING} \]

**Example**

Relative Permeability = Encore_Function \( \text{Name}=\text{My\_Function} \)

**Description**

Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```plaintext
Begin String Function My\_Function
  Value is "200 + 1.0\*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument \text{EVAL\_TYPE} can be used to select the particular function on the Encore function object. Valid options include \text{VALUE}, \text{DOT}, \text{GRADIENT}, \text{FLUX} and \text{STRESS}. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – \text{VALUE} for \text{NO\_OP} expressions, \text{DOT} for \text{DT\_OP} expressions and \text{GRADIENT} for \text{GRAD\_OP} expressions.

### 4.37.3 RELATIVE PERMEABILITY = POLYNOMIAL

**Parameters**

\[ \text{VARIABLE} = \text{STRING} \]

\[ \text{ORDER} = \text{INT} \]

\[ \text{VARIABLE\_OFFSET} = \text{REAL} \]

\[ \text{C0} = \text{REAL} \]

\[ \text{C1} = \text{REAL} \]

... 

\[ \text{CN} = \text{REAL} \]
Relative Permeability= Polynomial Variable=Temperature Order=1 C0=401.0 C1=88.5

Description
Arbitrary order polynomial function of a specified scalar variable.

\[ k_r = \sum_{i=0}^{N} C_i (X + X_o)^i \]  (4.83)

Here, \( N \) is the order of the polynomial provided by the ORDER parameter and \( X \) is the variable supplied by the VARIABLE parameter, \( X_o \) is an optional offset (VARIABLE_OFFSET, default is zero) and \( C_i \) are the supplied coefficients. Coefficients that are not supplied default to a value of zero. The VARIABLE argument can be TIME or any internal Expression that evaluates to a scalar. For the latter case, the format of the VARIABLE argument is described in section 2.6.1.

4.38 SEEBECK COEFFICIENT

SEEBECK COEFFICIENT = MODEL [param1 = val1, param2 = val2 ... ]

Description
Specifies the material model for Seebeck coefficient.

Summary
Specifies the material model for the Seebeck coefficient that appears in a thermoelectric contribution to the diffusion term of the energy equation for temperature. This command line must be accompanied the HEAT_CONDUCTION = THERMOELECTRIC 4.15.5 and CURRENT_DENSITY = THERMOELECTRIC 4.6.4 options which utilize isotropic thermal and electrical conductivity tensors. Additionally the Energy equation for temperature should have the SRC term active for JOULE HEATING 11.3.10.

4.38.1 SEEBECK COEFFICIENT = CONSTANT

Parameters
\( A = \text{REAL} \)

Example
SEEBECK COEFFICIENT = CONSTANT A = 1.0

Description
\( A \) is the value of the constant Seebeck coefficient. Must be accompanied by the HEAT_CONDUCTION = THERMOELECTRIC option.

4.38.2 SEEBECK COEFFICIENT = ENCORE_FUNCTION

Parameters
\( \text{NAME} = \text{STRING} \)
\( [\text{EVAL_TYPE} = \text{STRING}] \)

Example
Seebeck Coefficient = Encore_Function Name=My_Function
Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```
Begin String Function My_Function
  Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument `EVAL_TYPE` can be used to select the particular function on the Encore function object. Valid options include `VALUE`, `DOT`, `GRADIENT`, `FLUX` and `STRESS`. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – `VALUE` for `NO_OP` expressions, `DOT` for `DT_OP` expressions and `GRADIENT` for `GRAD_OP` expressions.

### 4.38.3 SEEBECK COEFFICIENT = POLYNOMIAL

**Parameters**
- `VARIABLE` = `STRING`
- `ORDER` = `INT`
- `[VARIABLE_OFFSET` = `REAL]`
- `[C0` = `REAL]`
- `[C1` = `REAL]`
- `...`
- `[CN` = `REAL]`

**Example**

Seebeck Coefficient= Polynomial Variable=Temperature Order=1 C0=401.0 C1=88.5

**Description**

Arbitrary order polynomial function of a specified scalar variable.

\[
\kappa = \sum_{i=0}^{N} C_i (X + X_o)^{i} 
\]

Here, \(N\) is the order of the polynomial provided by the `ORDER` parameter and \(X\) is the variable supplied by the `VARIABLE` parameter, \(X_o\) is an optional offset (`VARIABLE_OFFSET`, default is zero) and \(C_i\) are the supplied coefficients. Coefficients that are not supplied default to a value of zero. The `VARIABLE` argument can be `TIME` or any internal Expression that evaluates to a scalar. For the latter case, the format of the `VARIABLE` argument is described in section 2.6.1.

### 4.38.4 SEEBECK COEFFICIENT = USER_FUNCTION

**Parameters**
- `NAME` = `STRING`
- `X` = `STRING`
Example

begin definition for function SI_K
  type is piecewise linear
  begin values
    20.0  5.50e7
    100.0  4.60e7
    ...
    800.0  1.30e7
    2000.0  1.30e7
  end values
end definition for function SI_K

Begin Aria Material Foo
  ...
  Seebeck Coefficient = User_Function Name=SI_K X=Temperature
  ...
End Aria Material Foo

Description
A look-up function is used to compute the values of the thermal conductivity as a function of some other variable, i.e. $f(x)$. The function type (“piecewise linear” in the example above) must support the differentiate() method for Newton’s method.

Here `NAME` is the name of the user-defined function for (COEFFICIENT_DATA, SI_K, in the example) and `X` is the Aria name of the abscissa variable (TEMPERATURE in the example). Note that `X` is not necessarily the same name as the optional abscissa variable identified in the user-defined function. Must be accompanied by the HEAT_CONDUCTION = THERMOELECTRIC option. Full effect of this contribution can only be realized if Energy equation for temperature has the SRC term active for JOULE HEATING (11.3.10).

4.39 SHELL THICKNESS

SHELL THICKNESS = MODEL[param1 = val1, param2 = val2 ...]

Description
Specifies the material model for the shell thickness.

Summary
Specifies the material model and parameters for the shell thickness.

Parent Block(s)
ARIA MATERIAL

4.39.1 SHELL THICKNESS = CONSTANT

Parameters
  T = REAL

Example
  Shell Thickness = Constant T = 0.8

Description
  T is the value of the constant shell thickness.
4.39.2 SHELL THICKNESS = ELEMENT_ATTRIBUTE

Parameters

**MULTIPLIER** = *REAL*

Example

Shell Thickness = Element_Attribute multiplier = 1.0

Description

For shell elements the ExodusII mesh file can contain area attribute values for each element within an element block. This value can then be retrieved and used by the application. This model is used to scale the value of area attribute by **Multiplier** to obtain the shell thickness for a particular element.

4.40 SKELETON DENSITY

**SKELETON DENSITY** = *MODEL*[param1 = val1, param2 = val2 ...]

Description

Specifies a model for the porous skeleton density of a material.

Summary

Specifies a model for the porous skeleton density of a material.

Parent Block(s)

ARIA MATERIAL

4.40.1 SKELETON DENSITY = CONSTANT

Parameters

**RHO** = *REAL*

Example

Skeleton Density = Constant Rho=1e3

Description

The value is constant in space and time.

4.40.2 SKELETON DENSITY = ENCORE_FUNCTION

Parameters

**NAME** = *STRING*

[EVAL_TYPE = *STRING*

Example

Skeleton Density = Encore_Function Name=My_Function

Description

Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:
Then, this function can be used as shown in the example above.
The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

4.41 SKELETON INTERNAL ENERGY

SKELETON INTERNAL ENERGY = MODEL[param1 = val1, param2 = val2 ...]

Description  Specifies a model for the porous skeleton internal energy of a material.

Summary     Specifies a model for the porous skeleton internal energy of a material.

Parent Block(s)  ARIA MATERIAL

4.41.1 SKELETON INTERNAL ENERGY = CONSTANT

Parameters  E

Example      Skeleton Internal Energy = Constant E=2.3e-4

Description  The value is constant in space and time.

4.41.2 SKELETON INTERNAL ENERGY = ENCORE_FUNCTION

Parameters  NAME = STRING

Example      Skeleton Internal Energy = Encore_Function Name=My_Function

Description  Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```plaintext
Begin String Function My_Function
  Value is "200 + 1.0*t"
End
```
Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

### 4.41.3 SKELETON INTERNAL ENERGY = LINEAR

**Parameters**
- CP = REAL
- T_REF = REAL

**Example**
Skeleton Internal Energy = Linear Cp=13.7 T_ref=298.15

**Description**
The internal energy of the porous skeleton, $e_s$, is given by the simple relation,

$$ e_s = C_p (T - T_{ref}) $$

(4.85)

where $C_p$ is the specific heat supplied by the CP parameter and $T_{ref}$ is a reference temperature supplied by the T_REF parameter. This model also supplies the time derivative of $e_s$,

$$ \frac{\partial e_s}{\partial t} = C_p \frac{\partial T}{\partial t} $$

(4.86)

### 4.42 SKELETON SPECIFIC HEAT

**SKELETON SPECIFIC HEAT = MODEL[param1 = val1, param2 = val2 ...]**

**Description**
Specifies a model for the porous skeleton specific heat of a material.

**Summary**
Specifies a model for the porous skeleton specific heat of a material.

**Parent Block(s)**
ARIA MATERIAL

#### 4.42.1 SKELETON SPECIFIC HEAT = CONSTANT

**Parameters**
- CP = REAL

**Example**
Skeleton Specific Heat = Constant Cp=400

**Description**
The value is constant in space and time.
4.42.2 SKELETON SPECIFIC HEAT = ENCORE_FUNCTION

Parameters

- NAME = STRING
  - [EVAL_TYPE = STRING]

Example

Skeleton Specific Heat = Encore_Function Name=My_Function

Description

Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```
Begin String Function My_Function
  Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

4.42.3 SKELETON SPECIFIC HEAT = POLYNOMIAL

Parameters

- VARIABLE = STRING
- ORDER = INT
  - [VARIABLE_OFFSET = REAL]
  - [C0 = REAL]
  - [C1 = REAL]
  - ...
  - [CN = REAL]

Example

Skeleton Specific Heat= Polynomial Variable=Temperature Order=1 C0=401.0 C1=88.5

Description

Arbitrary order polynomial function of a specified scalar variable.

\[ C_p = \sum_{i=0}^{N} C_i (X + X_o)^i \]  \hspace{1cm} (4.87)

Here, \( N \) is the order of the polynomial provided by the ORDER parameter and \( X \) is the variable supplied by the VARIABLE parameter, \( X_o \) is an optional offset (VARIABLE_OFFSET, default is zero) and \( C_i \) are the supplied coefficients. Coefficients that are not supplied default to a value of zero. The VARIABLE argument can be TIME or any internal Expression that evaluates to a scalar. For the latter case, the format of the VARIABLE argument is described in section 2.6.1.

4.42.4 SKELETON SPECIFIC HEAT = USER_FUNCTION

Parameters

- NAME = STRING
- X = STRING
Example

Begin Definition for Function Table_Fcn
Type is Piecewise Linear
Begin Values
273.15 100
277.15 200
283.15 300
End
End
...

Begin Aria Material Foo
...
Skeleton Specific Heat = User_Function Name=Table_Fcn X=Temperature
...
End Aria Material Foo

Description
A look-up function is used to compute the values of the skeleton specific heat as a function of some other variable, i.e. \( f(x) \). The function type (“piecewise linear” in the example above) must support the \texttt{differentiate()} method for Newton’s method.

Here \texttt{NAME} is the name of the user-defined function (\texttt{Table_Fcn} in the example) and \texttt{X} is the Aria name of the abscissa variable (\texttt{TEMPERATURE} in the example). Note that \texttt{X} is not necessarily the same name as the abscissa variable identified in the user-defined function (\texttt{T} in the example).

4.43 SOLID LAMBDA

SOLID LAMBDA = \texttt{MODEL}[\texttt{param} \_1 = \texttt{val} \_1, \texttt{param} \_2 = \texttt{val} \_2 \ldots ]

Description
Specifies the material model for the \( \lambda \) Lamé coefficient for the solid stress tensor.

Summary
The solid stress \( \mathbf{T} \) is given by
\[
\mathbf{T} = \lambda E_{kk} \mathbf{I} + 2\mu \mathbf{E} - \beta (T - T_{\text{ref}}) \mathbf{I}
\]  
(4.88)

where \( \lambda \) and \( \mu \) are the Lamé coefficients, \( \mathbf{E} = \frac{1}{2} \left( \nabla \mathbf{d} + \nabla \mathbf{d}^T \right) \) is the deformation tensor, \( \beta \) is the coefficient of thermal stress, \( T \) is temperature and \( T_{\text{ref}} \) is the thermal strain reference temperature.

These Lamé coefficients are related to the more standard Young’s modulus, Poisson’s ratio and CTE (\( \alpha \)) as follows:

\[
2\mu = \frac{E}{(1 + \nu)}
\]  
(4.89)

\[
\lambda = \frac{\nu E}{(1 + \nu)(1 - 2\nu)} = 2\mu \frac{\nu}{(1 - 2\nu)}
\]  
(4.90)

\[
\beta = \frac{\alpha E}{(1 - 2\nu)} = \alpha (3\lambda + 2\mu)
\]  
(4.91)

When a user supplies the Young’s modulus, Poisson’s ratio and CTE properties Aria internally converts them into the Lamé coefficients.

Parent Block(s) ARIA MATERIAL
4.43.1 SOLID LAMBDA = CONSTANT

Parameters

LAMBDA = REAL

Example

SOLID LAMBDA = CONSTANT LAMBDA = 1.0

Description

LAMBDA is the value of the constant λ.

4.43.2 SOLID LAMBDA = CONVERTED

Parameters

(None)

Example

SOLID LAMBDA = Converted

Description

Aria will use Young's modulus and Poisson ratio to compute the Lamé λ coefficient. Supplying the Lamé coefficients is more computationally efficient but perhaps less convenient, especially if the material properties are varying (e.g., temperature dependent in a non-isothermal problem).

4.43.3 SOLID LAMBDA = ENCORE_FUNCTION

Parameters

NAME = STRING
[EVAL_TYPE = STRING]

Example

SOLID LAMBDA = Encore_Function Name=My_Function

Description

Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```
Begin String Function My_Function
  Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

4.44 SOLID TWO MU

SOLID TWO MU = MODEL[param1 = val1, param2 = val2 ...]
Specifies the material model for twice the \( \mu \) Lamé coefficient for the solid stress tensor.

The solid stress \( \mathbf{T} \) is given by

\[
\mathbf{T} = \lambda \mathbf{E}_{kk} \mathbf{I} + 2\mu \mathbf{E} - \beta (T - T_{\text{ref}}) \mathbf{I}
\]  \hspace{1cm} (4.92)

where \( \lambda \) and \( \mu \) are the Lamé coefficients, \( \mathbf{E} = \frac{1}{2} \left( \nabla \mathbf{d} + \nabla \mathbf{d}^T \right) \) is the deformation tensor, \( \beta \) is the coefficient of thermal stress, \( T \) is temperature and \( T_{\text{ref}} \) is the thermal strain reference temperature.

These Lamé coefficients are related to the more standard Young’s modulus, Poisson’s ratio and CTE (\( \alpha \)) as follows:

\[
2\mu = \frac{E}{(1 + \nu)} \hspace{1cm} (4.93)
\]

\[
\lambda = \frac{\nu E}{(1 + \nu)(1 - 2\nu)} = 2\mu \frac{\nu}{(1 - 2\nu)} \hspace{1cm} (4.94)
\]

\[
\beta = \frac{\alpha E}{(1 - 2\nu)} = \alpha (3\lambda + 2\mu) \hspace{1cm} (4.95)
\]

When a user supplies the Young’s modulus, Poisson’s ratio and CTE properties Aria internally converts them into the Lamé coefficients.

4.44.1 SOLID TWO MU = CONSTANT

Parameters \hspace{1cm} TWO_MU = REAL

Example \hspace{1cm} SOLID TWO MU = CONSTANT TWO_MU = 1.0

Description \hspace{1cm} TWO_MU is the value of \( 2\mu \).

4.44.2 SOLID TWO MU = CONVERTED

Parameters \hspace{1cm} (None)

Example \hspace{1cm} SOLID TWO MU = Converted

Description \hspace{1cm} Aria will use Young’s modulus and Poisson ratio to compute the Lamé \( \mu \) coefficient. Supplying the Lamé coefficients is more computationally efficient but perhaps less convenient, especially if the material properties are varying (e.g., temperature dependent in a non-isothermal problem).
4.44.3 SOLID TWO MU = ENCORE_FUNCTION

Parameters
- NAME = STRING
  [EVAL_TYPE = STRING]

Example
TWO_MU = Encore_Function Name=My_Function

Description
Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```
Begin String Function My_Function
  Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

4.45 SOLID STRESS

SOLID STRESS = MODEL [param1 = val1, param2 = val2 ...]

Description
Specifies a contribution to the solid stress tensor. Multiple stresses are combined additively and may be specified by using this line command multiple times.

Summary
Specifies a contribution to the solid stress tensor. The total stress $T$ is given by

$$T = \sum_j T_j$$

Parent Block(s) ARIA MATERIAL

4.45.1 SOLID STRESS = ISOTHERMAL

Parameters
- $T = \text{REAL}$
- $T_{\text{REF}} = \text{REAL}$

Example
SOLID STRESS = Isothermal $T=500$ $T_{\text{REF}}=325$

Description
This stress accounts for the mechanical stresses due to thermally induced strains.

$$T = -\beta (T - T_{\text{REF}}) I$$

(4.97)
where $\beta$ is the Lamé coefficient of thermal stress (related to the coefficient of thermal expansion, $\alpha$), $T$ is the temperature and $T_{\text{ref}}$ is the temperature of the undeformed reference state of the mesh (pseudo-solid). This is a specialization of the THERMAL model that uses uniform, fixed temperature and thermal strain reference temperature.

### 4.45.2 SOLID STRESS = LINEAR ELASTIC

**Parameters**

| REFERENCE_FRAME = ‘MOVING’ | ‘UNDEFORMED’ |

**Example**

SOLID STRESS = Linear_Elastic Reference_Frame=Moving

**Description**

Supplies the linear elasticity stress tensor,

$$ T = \lambda \text{trace} \, EI + 2\mu E $$

(4.98)

where $\lambda$ and $\mu$ and the Lamé coefficients and $E$ is the strain tensor. When the choice of reference frame is “UNDEFORMED” then the strain is computed in the undeformed reference state; this is commonly referred to as small strain theory. When the reference frame is “MOVING” then the strain is computed with respect to the deformed coordinates.

Specifically, the strain tensor is given by

$$ E = \frac{1}{2} (\nabla d + \nabla d^t) $$

(4.99)

where $d$ is the solid displacement field. The choice of reference frame determines whether the $\nabla$ operator is computed in the undeformed or moving reference frames.

### 4.45.3 SOLID STRESS = NEOHOOKEAN ELASTIC

**Parameters**

(none)

**Example**

SOLID STRESS = Neohookean_Elastic

**Description**

Supplies a nonlinear hyperelastic stress of the form,

$$ T = \frac{\mu}{J} (b - I) + \frac{\lambda}{J} \ln J $$

(4.100)

where $\lambda$ and $\mu$ and the Lamé coefficients, $b \equiv F \cdot F^t$ is the left Cauchy-Green tensor, $F$ is the deformation gradient and $J \equiv \det F$. See, e.g., [4] or [5].

### 4.45.4 SOLID STRESS = NONLINEAR ELASTIC

**Parameters**

(none)

**Example**

SOLID STRESS = Nonlinear_Elastic
Supplies a nonlinear elastic stress,
\[ T = \lambda \text{trace} \mathbf{E} I + 2\mu \mathbf{E} \] (4.101)

where \( \lambda \) and \( \mu \) and the Lamé coefficients and \( \mathbf{E} \) is the strain tensor. The particular choice of strain tensor chosen depends on where the configuration (reference frame) which is set via the MESH MOTION command line. See section 5.16 for more information. When the MESH MOTION is set to REFERENCE_FRAME then the Green strain is used. Otherwise, the Almansi strain is used.

### 4.45.5 SOLID STRESS = RESIDUAL

**Parameters**

- \([SXX | SX = \text{REAL}]\)
- \([SYY | SY = \text{REAL}]\)
- \([SZZ | SZ = \text{REAL}]\)
- \([SXY = \text{REAL}]\)
- \([SXZ = \text{REAL}]\)
- \([SYZ = \text{REAL}]\)

**Example**

SOLID STRESS = Residual Sxx=0.02 Syy=0.02

**Description**

This stress accounts for the initial residual stress in a solid that is constant and uniform everywhere. The components of the residual stress tensor are supplied by the (up-to) six components \( SXX, SYY, SZZ, SXY, SXZ, \) and \( SYZ \). This is directly analogous to the ISTRESS condition in ANSYS. To that end, the diagonal components can be specified as either \( SXX \) or \( SX \) etc.

### 4.45.6 SOLID STRESS = THERMAL

**Parameters**

-(none)-

**Example**

SOLID STRESS = Thermal

**Description**

This stress accounts for the mechanical stresses due to thermally induced strains.

\[ T = -\beta (T - T_{ref}) I \] (4.102)

where \( \beta \) is the Lamé coefficient of thermal stress (related to the coefficient of thermal expansion, \( \alpha \)), \( T \) is the temperature and \( T_{ref} \) is the temperature of the undeformed reference state of the solid.

### 4.45.7 SOLID STRESS = LAME

**Parameters**

- MODEL = STRING

**Example**

SOLID STRESS = Lame Model = Neo_Hookean
This stress interfaces with the Lame constitutive model library used by Sierra/SolidMechanics. As such, it allows the use of many of Lame’s constitutive models. Sierra/Multimechanics prepares the kinematic quantities, and Lame returns the Cauchy stress, which is integrated in the SOLID equation.

Material parameters for the constitutive models are handled by an input block consistent with the Sierra/SolidMechanics formatting. For documentation of the available constitutive models and the required parameter input, please see the Sierra/SolidMechanics documentation.

Currently this interface is instrumented to support hyperelastic constitutive models, and is restricted to the following models:

- **NEO_HOOKEAN**
- **THERMALBATTERYSEPARATOR**
- **SOILFOAMPHASETRANSITION**

### 4.46 SPECIES DIFFUSION

SPECIES DIFFUSION = *MODEL*[param₁ = val₁, param₂ = val₂ ...]

**Description**  
Specifies the material (constitutive) model for the species diffusion (diffusive flux) in the bulk.

**Summary**  
Specifies the material (constitutive) model for the species diffusion (diffusive flux) in the bulk.

**Parent Block(s)**  
ARIA MATERIAL

#### 4.46.1 SPECIES DIFFUSION = BASIC

This is an alias for FICKS_LAW.

**Example**  
Species Diffusion = Basic

#### 4.46.2 SPECIES DIFFUSION = FICKS_LAW

**Parameters**  
(none)

**Example**  
Species Diffusion = Ficks_Law

**Description**  
The diffusive species flux \( q \) is given by Fick’s Law,

\[
q = -D \nabla C
\]

where \( D \) is the species diffusivity and \( C \) is the species concentration.
4.47 SPECIES DIFFUSIVITY

Species diffusivity = Model[param1 = val1, param2 = val2 ...]

Description  Specifies the material model for the species diffusivity.

Summary  Specifies the material model for the species diffusivity.

Parent Block(s)  ARIA MATERIAL

4.47.1 SPECIES DIFFUSIVITY = ARRHENIUS

Parameters
[C = REAL]
D = REAL
Q = REAL
R = REAL

Example  SPECIES DIFFUSIVITY = Arrhenius D=2e-6 Q=6900 R=8.314

Description  The species diffusivity is given by the Arrhenius function

\[ D = C + D_0 e^{-Q/RT} \]  (4.104)

where \( C \) is given by the \( C \), \( D_0 \) is given by the \( D \) parameter, \( Q \) is given by the \( Q \) parameter and \( R \) is given by the \( R \) parameter.

4.47.2 SPECIES DIFFUSIVITY = CONSTANT

Parameters  D = REAL

Example  SPECIES DIFFUSIVITY = CONSTANT D = 1.0

Description  \( D \) is the value of the constant species diffusivity.

4.47.3 SPECIES DIFFUSIVITY = ENCORE_FUNCTION

Parameters
[NAME = STRING]
[EVAL_TYPE = STRING]

Example  SPECIES DIFFUSIVITY = Encore_Function Name=My_Function
Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```plaintext
Begin String Function My_Function
    Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument `EVAL_TYPE` can be used to select the particular function on the Encore function object. Valid options include `VALUE`, `DOT`, `GRADIENT`, `FLUX` and `STRESS`. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – `VALUE` for `NO_OP` expressions, `DOT` for `DT_OP` expressions and `GRADIENT` for `GRAD_OP` expressions.

### 4.48 SPECIFIC HEAT

**SPECIFIC HEAT = MODEL[par\_m\_1 = val\_1, par\_m\_2 = val\_2 ...]**

**Description**
Specifies the material model for the specific heat.

**Summary**
Specifies the material model for the specific heat.

**Parent Block(s)**
ARIA MATERIAL

#### 4.48.1 SPECIFIC HEAT = CONSTANT_EVALUATOR

**Parameters**
- `CP = REAL`
- `h0 = REAL`

**Example**
SPECIFIC HEAT = CONSTANT_EVALUATOR CP = 1.0 h0 = 500

**Description**
`CP` is the value of the constant specific heat, and `h0` (optional) is the value of the reference enthalpy. This also creates an expression for enthalpy, as \( h(T) = h_0 + C_pT \).

#### 4.48.2 SPECIFIC HEAT = POLYNOMIAL_EVALUATOR

**Parameters**
- `a0 = REAL`
- `a1 = REAL`
- `a2 = REAL`
- `a3 = REAL`
- `a4 = REAL`
- `a5 = REAL`
- `a6 = REAL`
- `a7 = REAL`
- `a8 = REAL`
- `R = REAL`

**Example**
SPECIFIC HEAT = POLYNOMIAL_EVALUATOR a0 = 50 a1 = 1000 a2 = 1 R = 8.314

**Description**
The coefficients `a0` to `a8` are the polynomial coefficients and `R` is the gas constant they are based on (defaults to 8.314). Only `a0` and `a1` are required.

Specific heat is evaluated using \( C_p(T)/R = \sum_{i=1}^{n} a_i T^{i-1} \) and enthalpy by \( h(T)/R = a_0 + \sum_{i=1}^{n} \frac{1}{2} a_i T^i \). The polynomial order is determined by how many non-zero entries are provided.
4.48.3 SPECIFIC HEAT = TABULAR_EVALUATOR

Parameters
Cp_function = STRING h0 = REAL

Example
SPECIFIC HEAT = TABULAR_EVALUATOR Cp_function = f_Cp

Description
The function specified here is the same syntax as for USER_FUNCTION, but the function must be with respect to temperature. The reference enthalpy, h0 is optional. This also provides an expression for enthalpy that is based on an integration of the tabular values of specific heat.

4.48.4 SPECIFIC HEAT = T_EXPONENT

Parameters
Cp_ref = REAL T_ref = REAL n = REAL h0 = REAL

Example
SPECIFIC HEAT = T_EVALUATOR Cp_ref = 1000 T_ref = 300 n = 2

Description
This provides specific heat as \( C_p(T) = C_{p,\text{ref}} \left( \frac{T}{T_{\text{ref}}} \right)^n \) and enthalpy as \( h(T) = h_0 + \frac{C_{p,\text{ref}}}{T_{\text{ref}}^n} \frac{T^{n+1}}{n+1} \).

4.48.5 SPECIFIC HEAT = CONSTANT

Parameters
CP = REAL

Example
SPECIFIC HEAT = CONSTANT CP = 1.0

Description
CP is the value of the constant specific heat.

4.48.6 SPECIFIC HEAT = CURING_FOAM

Parameters
VFRAC_SUBINDEX = INT [CP_FL = REAL] [CP_FG = REAL] [CP_E = REAL] [PHI_ZERO = REAL]

Example
Specific Heat = Curing_Foam Vfrac_Subindex=1 Cp_fl=1 Cp_fg=1 Cp_e=1 phi_zero=0.2

Description
For a curing epoxy with volume fraction \( \phi \) the specific heat is given by

\[
C_p = C_{p,\text{fl}} \phi + C_{p,\text{fg}} (\phi_o - \phi) + C_{p,e} (1 - \phi_o) \tag{4.105}
\]

\[
= a + b\phi \tag{4.106}
\]
where $C_{p,f_l}$ is the specific heat of the liquid phase flourinert, $C_{p,f_g}$ is the specific heat of the gas phase flourinert, $C_{p,e}$ is the specific heat of the epoxy and $\phi_o$ is the reference volume fraction in the flourinert. In the latter form of this relationship

\[
\begin{align*}
  a &= C_{p,f_l} - C_{p,f_g} \\
  b &= C_{p,f_g}\phi_o + C_{p,e}(1 - \phi_o).
\end{align*}
\]

(4.107) (4.108)

**NOTE:** The volume fraction is assumed to be a SPECIES field with the subindex provided by the VFRAC_SUBINDEX parameter.

### 4.48.7 SPECIFIC HEAT = ENCORE_FUNCTION

**Parameters**

- **NAME** = STRING
- [EVAL_TYPE = STRING]

**Example**

```
SPECIFIC HEAT = Encore_Function Name=My_Function
```

**Description**

Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```
Begin String Function My_Function
  Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

### 4.48.8 SPECIFIC HEAT = EXPONENTIAL

**Parameters**

- **VARIABLE** = STRING
- [CONSTANT = REAL]
- [MULTIPLIER = REAL]
- **EXPONENT** = REAL

**Example**

```
Specific Heat = Exponential Variable=Temperature Multiplier=1.0
Exponent=-0.3
```

**Description**

Exponential function of in specified scalar variable. The specific heat is computed as

\[
C_p = C + M e^{EX}
\]

(4.109)

Here, $C$ is the constant term supplied by the CONSTANT parameter which defaults to zero, $M$ is the value supplied by the MULTIPLIER parameter which defaults to unity, $X$ is the variable supplied by the VARIABLE parameter and $E$ is the exponential multiplier provided by the EXPONENT parameter.
4.48.9 SPECIFIC HEAT = POLYNOMIAL

Parameters
- VARIABLE = STRING
- ORDER = INT
- [VARIABLE_OFFSET = REAL]
- [C0 = REAL]
- [C1 = REAL]
- ...
- [CN = REAL]

Example
Specific Heat= Polynomial Variable=Temperature Order=1 C0=401.0 C1=88.5

Description
Arbitrary order polynomial function of a specified scalar variable.

\[ C_p = \sum_{i=0}^{N} C_i (X + X_o)^i \] (4.110)

Here, \( N \) is the order of the polynomial provided by the ORDER parameter and \( X \) is the variable supplied by the VARIABLE parameter, \( X_o \) is an optional offset (VARIABLE_OFFSET, default is zero) and \( C_i \) are the supplied coefficients. Coefficients that are not supplied default to a value of zero. The VARIABLE argument can be TIME or any internal Expression that evaluates to a scalar. For the latter case, the format of the VARIABLE argument is described in section 2.6.1.

4.48.10 SPECIFIC HEAT = USER_FUNCTION

Parameters
- NAME = STRING
- X = STRING
**Example**

```plaintext
begin definition for function Water_Heat_Capacity
    # Source Appendix 2 from "Transport Processes and
    # Unit Operations" by C. J. Geankoplis
    type is piecewise linear
    begin values
        # K J / kg K
        273.15 4220
        283.15 4195
        293.15 4185
        298.15 4182
        303.15 4181
        313.15 4181
        323.15 4183
        333.15 4187
        343.15 4192
        353.15 4199
        363.15 4208
        373.15 4219
    end
end
...

Begin Aria Material Foo
...
    Specific Heat = User_Function X=Temperature Name=Water_Heat_Capacity
...
End Aria Material Foo
```

**Description**

A look-up function is used to compute the values of the specific heat as a function of some other variable, i.e. \( f(x) \). The function type ("piecewise linear" in the example above) must support the `differentiate()` method for Newton’s method.

Here `NAME` is the name of the user-defined function (`Water_Heat_Capacity` in the example) and `X` is the Aria name of the abscissa variable (`TEMPERATURE` in the example). Note that `X` is not necessarily the same name as the abscissa variable identified in the user-defined function (`T` in the example).

### 4.48.11 SPECIFIC HEAT = USE_PHASE_CHANGE

```plaintext
SPECIFIC_HEAT = USE_PHASE_CHANGE FLH=LH_VAL TS=T_SOLIDUS TL=T_LIQUIDUS
```

**Description**

Defines a specialized material model to account for change of phase during melting or solidification. Must be accompanied by the PHASE CHANGE SPECIFIC HEAT command line.

**Summary**

Phase change is being modeled by treating the phase transition temperature region as a mushy-zone about the melt temperature. Here the energy associated with phase change in the mushy-zone is accounted for by modifying the specific heat to include the latent heat of fusion as an additional contribution to the energy equation.

\[
\text{Specific Heat} = \begin{cases} 
C_p(T) & T_s < T \\
C_p(T) + \frac{FLH}{T_l - T_s} & T_s < T < T_l \\
C_p(T) & T > T_l
\end{cases}
\]

317
where $T$ is the current temperature, $C_p(T)$ is the specific heat of the material, $FLH$ is the latent heat of fusion, $T_s$ is a solidus temperature, $T_l$ is a liquidus temperature and $T_l - T_s$ is the temperature range of the mushy zone.

Material outside of the mushy-zone, $T < T_s$ or $T > T_l$, will employ a standard specific heat model, $C_p(T)$.

While a material is undergoing phase change, the time stepping will be restricted so that the modeled phase change behavior will be captured. Use of this model will often require additional restrictions be placed upon the adaptive time stepping 16.4, 16.4.1.

### 4.49 PHASE CHANGE SPECIFIC HEAT

**PHASE CHANGE SPECIFIC HEAT = SPECIFIC_HEAT_MODEL [param1 = val1, param2 = val2 …]**

**Description**
Defines the specific heat material model used in conjunction with phase change. This command can request any valid SPECIFIC HEAT model aside from USE_PHASE_CHANGE and must be accompanied by a SPECIFIC HEAT = USE_PHASE_CHANGE command line.

**Summary**
Includes the parameters for the specific heat material model.

**Parent Block(s)**
ARIA MATERIAL

**Parameters**
The optional parameters are those appropriate to the SPECIFIC_HEAT_MODEL.

| Example | Specific Heat = USE_PHASE_CHANGE FLH = 2.3E+04 TS = 599 TL = 600.5 |

### 4.50 SURFACE TENSION

**SURFACE TENSION = MODEL [param1 = val1, param2 = val2 …]**

**Description**
Specifies the model to use for the surface (interfacial) tension.

**Summary**
Specifies the model to use for the surface (interfacial) tension.

**Parent Block(s)**
ARIA MATERIAL
4.50.1 SURFACE TENSION = CONSTANT

Parameters

SIGMA = REAL

Example

Surface Tension = Constant Sigma = 72.0

Description

SIGMA is the value of the surface tension.

4.50.2 SURFACE TENSION = ENCORE_FUNCTION

Parameters

NAME = STRING

[EVAL_TYPE = STRING]

Example

Surface Tension = Encore_Function Name=My_Function

Description

Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```
Begin String Function My_Function
  Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

4.50.3 SURFACE TENSION = LINEAR_T

Parameters

SIGMA0 = REAL

DSIGMADT = REAL

T_REF = REAL

Example

Surface Tension = Linear_T sigma0=72. dsigmadt = -.15 T_ref = 298.

Description

SIGMA0 is the value of the surface tension at the reference temperature T_REF and DSIGMADT is the derivative of the surface temperature with respect to temperature, i.e.,

\[ \sigma = \sigma_0 + m(T - T_{ref}) \tag{4.111} \]

where \( m \) is DSIGMADT.
4.51 SUSPENSION FLUX

SUSPENSION FLUX = MODEL[param1 = val1, param2 = val2 ...]

Description Specifies the parameters for the suspension flux model.

Summary Specifies the suspension flux model and its parameters for this material.

Parent Block(s) ARIA MATERIAL

4.51.1 SUSPENSION FLUX = PHILLIPS

Parameters

\[ \begin{align*}
K_\mu & = REAL \\
K_c & = REAL \\
\phi_{max} & = REAL \\
\beta & = REAL \\
particle\_radius & = REAL
\end{align*} \]

Example

SUSPENSION FLUX = Phillips K_\mu=0.62 K_c=0.41 \phi_{max}=0.68 \beta=-1.82 particle\_radius=0.01

Description The Phillips diffusive flux model is intended to be used in conjunction with the Krieger viscosity model (4.56.10). Here, the flux is given by

\[ q = \left( K_c a^2 - K_\mu a^2 \beta \frac{\phi}{\phi_m - \phi} \right) \dot{\gamma} \phi \nabla \phi + K_c a^2 \phi^2 \nabla \dot{\gamma} \]

(4.112)

where \( \dot{\gamma} \) is the shear rate, \( \phi \) is the suspension concentration, \( \phi_m \) is the maximum suspension concentration and \( a \) is the particle radius.

4.52 THERMAL CONDUCTIVITY

THERMAL CONDUCTIVITY = MODEL[param1 = val1, param2 = val2 ...]

Description Specifies the material model for the thermal conductivity.

Summary Specifies the material model for the thermal conductivity that appears in the diffusion term of the energy equation for temperature. The command line must be accompanied by either the HEAT_CONDUCTION = BASIC option for an isotropic conductivity tensor or the HEAT_CONDUCTION = GENERALIZED option for the anisotropic case.

Parent Block(s) ARIA MATERIAL
4.52.1 THERMAL CONDUCTIVITY = CALORE_USER_SUB

Parameters

- NAME = STRING
- TYPE = STRING
- [MULTIPLIER = REAL]
- [NR = INT]
- [R0 = INT]
- [R1 = INT(etc.)]
- [NI = INT]
- [I0 = INT]
- [I1 = INT(etc.)]

Example

Thermal Conductivity = Calore_User_Sub name=w80afftuser type=element NR=2 R0=1000 R1=0.7

Description

The Calore_User_Sub model allows the user to explicitly evaluate thermal conductivity as a function of temperature and position within an element. Additionally, a user_query feature allows evaluation of thermal conductivity based upon other problem criteria. NAME is the name of the user subroutine was registered with. NR and NI are the numbers of real and integer parameters, respectively. Each real (int) parameter is supplied by the Rn (In) parameter. Note: the parameters use zero based counting. The TYPE parameter denotes the API the user subroutine uses; currently Aria supports type ELEMENT for scalar valued thermal conductivity.

The optional MULTIPLIER parameter (defaults to 1.0) can be supplied to scale the output of the user subroutine, i.e., the flux \( f \) will be \( f = m f_{user} \) where \( m \) is the multiplier and \( f_{user} \) is the output of the user subroutine.

See section 9.2.2 for a more complete example of a Calore user subroutine.

Must be accompanied by the HEAT_CONDUCTION = BASIC option.

4.52.2 THERMAL CONDUCTIVITY = CONSTANT

Parameters

- \( K = REAL \)

Example

THERMAL CONDUCTIVITY = CONSTANT K = 1.0

Description

\( K \) is the value of the constant thermal conductivity. Must be accompanied by the HEAT_CONDUCTION = BASIC option.

4.52.3 THERMAL CONDUCTIVITY = CURING_FOAM

Parameters

- RHO_E = REAL
- K_F = REAL
- K_E = REAL

Example

Thermal Conductivity = Curing_Foam rho_e=1.3 k_e=14 k_f=2.7
Description For a curing epoxy with mixture density $\rho$

\[
\kappa = \frac{2}{3} \left( \frac{\rho}{\rho_e} \right) \kappa_e + \left( 1 - \frac{\rho}{\rho_e} \right) \kappa_f
\]

(4.113)

\[
= a + b\rho
\]

(4.114)

where $\rho_e$ is the density of the epoxy, $\kappa_e$ is the thermal conductivity of the epoxy and $\kappa_f$ is the thermal conductivity of the flourinert. In the latter form of this relationship

\[
a = \kappa_f
\]

(4.115)

\[
b = \frac{1}{\rho_e} \left( \frac{2}{3} \kappa_e - \kappa_f \right)
\]

(4.116)

Must be accompanied by the HEAT_CONDUCTION = BASIC option.

4.52.4 THERMAL CONDUCTIVITY = ENCORE_FUNCTION

Parameters

NAME = STRING

[EVAL_TYPE = STRING]

Example

THERMAL CONDUCTIVITY = Encore_Function Name=My_Function

Description

Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

\[
\text{Begin String Function My\_Function} \\
\text{Value is "200 + 1.0\_t"} \\
\text{End}
\]

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO\_OP expressions, DOT for DT\_OP expressions and GRADIENT for GRAD\_OP expressions.

4.52.5 THERMAL CONDUCTIVITY = MESH_INPUT

Parameters

NAME = STRING

AXIS1 = STRING

AXIS2 = STRING

[TYPE = STRING]

[MULTIPLIER = REAL]

[TOLERANCE = REAL]

Example

THERMAL CONDUCTIVITY = MESH_INPUT NAME = COND AXIS1 = A1 AXIS2 = A2 TOLERANCE = 1.0E-4 TYPE = SYMMETRIC MULTIPLIER = 2.0
Thermal conductivity is being provided from the input mesh file.

The TYPE parameter defines what tensor components are supplied in the input file. The default TYPE = PRINCIPAL will cause the code to expect principal components only. Supplying TYPE = SYMMETRIC will require that the file contain corresponding the symmetric tensor components.

The NAME parameter specifies the base name of the tensor component values present in the mesh file. Component names are internally generated using the base name and the component subscripts. For example for TYPE = SYMMETRIC in the two-dimensional a base name of COND would generate COND11, COND22 and COND12, and for TYPE = PRINCIPAL would generate COND11 and COND22.

The tensor components provided are assumed to be defined in an orientation compactly described with unit vectors. For two-dimensional problems one unit vector must be provided and for three-dimensional problems one must provide two unit vectors. In 3D the AXIS1 and AXIS2 parameters define the variable names which specify orientation vectors associated with the components. Default value of AXIS1 = FIBER and default value of AXIS2 = NORMAL. Orientation vectors are given relative to the fixed Cartesian (xyz) frame. Internal to the code AXIS1 and AXIS2 will be used to generate the remaining orthogonal axis needed to define a complete orthonormal transformation into the computational reference frame. Internal checks are done to guarantee that orientation vectors are of nonzero length. For 2D only the AXIS1 variable need be provided and AXIS2, if provided, will be ignored.

The TOLERANCE parameter defines a desired tolerance for the dot product of vectors defined by AXIS1 and AXIS2. An error message will be generated when the dot product is greater than the specified tolerance. Default value for TOLERANCE = 1.0E9

Use of the MULTIPLIER parameter results in scaling all components of the conductivity tensor.

**4.52.6 THERMAL CONDUCTIVITY = OPTICALLY_THICK**

**Parameters**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>K</td>
<td>REAL</td>
</tr>
<tr>
<td>N</td>
<td>REAL</td>
</tr>
<tr>
<td>BETA_R</td>
<td>REAL</td>
</tr>
</tbody>
</table>

**Example**

THERMAL CONDUCTIVITY = OPTICALLY_THICK K = 3.0 N = 2.1 BETA_R = 1.0e-5

**Description**

Thermal conductivity used to model combined pure diffusion and radiative diffusion in the optically-thick limit. Constant values for the pure diffusion thermal conductivity $K$, index of refraction $n$, and Rosseland-mean extinction coefficient $\beta_R$ must be provided on the input line.

$$\kappa = K + \frac{16\pi n^2}{3\beta_R}T^3$$

(4.117)

Must be accompanied by the HEAT_CONDUCTION = BASIC option. Additionally the GLOBAL CONSTANTS block must include include the STEFAN-BOLTZMANN CONSTANT entry to provide a value of $\sigma$.

**4.52.7 THERMAL CONDUCTIVITY = POWER_LAW**

**Parameters**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>REAL</td>
</tr>
<tr>
<td>GAMMA</td>
<td>REAL</td>
</tr>
</tbody>
</table>

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Example

**THERMAL CONDUCTIVITY = POWER_LAW**

**A** = 3.8  **GAMMA** = -1.2

Description

Thermal conductivity is a power law function of temperature

\[ \kappa = A T^\gamma \] (4.118)

where \( A \) is a scale factor and \( \gamma \) is the power law exponent. For values of \( \gamma = 1 \) the THERMAL model 4.52.9 should be used instead. Must be accompanied by the HEAT_CONDUCTION = BASIC option.

### 4.52.8 THERMAL CONDUCTIVITY = POLYNOMIAL

**Parameters**

- **VARIABLE** = STRING
- **ORDER** = INT
- **[VARIABLE_OFFSET]** = REAL
- **[C0]** = REAL
- **[C1]** = REAL
- ...  
- **[CN]** = REAL

**Example**

Thermal Conductivity= Polynomial 
Variable=Temperature 
Order=1 
C0=401.0 
C1=88.5

Description

Arbitrary order polynomial function of a specified scalar variable.

\[ \kappa = \sum_{i=0}^{N} C_i (X + X_o)^i \] (4.119)

Here, \( N \) is the order of the polynomial provided by the **ORDER** parameter and \( X \) is the variable supplied by the **VARIABLE** parameter, \( X_o \) is an optional offset (**VARIABLE_OFFSET**, default is zero) and \( C_i \) are the supplied coefficients. Coefficients that are not supplied default to a value of zero. The **VARIABLE** argument can be **TIME** or any internal Expression that evaluates to a scalar. For the latter case, the format of the **VARIABLE** argument is described in section 2.6.1.

### 4.52.9 THERMAL CONDUCTIVITY = THERMAL

**Parameters**

- **[A]** = REAL
- **[B]** = REAL
- **[C]** = REAL
- **[D]** = REAL

**Example**

**THERMAL CONDUCTIVITY = THERMAL**  
**A** = 401.0  **B** = 88.5

Description

Cubic polynomial function of temperature for the conductivity.

\[ \kappa = A + BT + CT^2 + DT^3 \] (4.120)

Must be accompanied by the HEAT_CONDUCTION = BASIC option.
4.52.10 THERMAL CONDUCTIVITY = USER_FUNCTION

Parameters
- **NAME** = STRING
- **X** = STRING

Example
```
begin definition for function SI_K
    type is piecewise linear
    begin values
        20.0 5.50e7
        100.0 4.60e7
        ...
        800.0 1.30e7
        2000.0 1.30e7
    end values
end definition for function SI_K
```

Description
A look-up function is used to compute the values of the thermal conductivity as a function of some other variable, i.e. \( f(x) \). The function type ("piecewise linear" in the example above) must support the `differentiate()` method for Newton’s method.

Here **NAME** is the name of the user-defined function for (CONDUCTIVITY_DATA, SI_K, in the example) and **X** is the Aria name of the abscissa variable (TEMPERATURE in the example). Note that **X** is not necessarily the same name as the optional abscissa variable identified in the user-defined function. Must be accompanied by the `HEAT_CONDUCTION = BASIC` option.

4.52.11 THERMAL CONDUCTIVITY = CONSTANT (TENSOR)

Parameters
```
XX = REAL
[XY = REAL]
[XZ = REAL]
[YX = REAL]
YY = REAL
[YZ = REAL]
[ZX = REAL]
[ZY = REAL]
ZZ = REAL
```

Example
```
TENSOR THERMAL CONDUCTIVITY = Constant XX=1.0 YY=2.0 ZZ=1.0
TENSOR THERMAL CONDUCTIVITY = Constant 11=1.0 22=2.0 33=1.0
```

Description
All components default to zero and all values are constant in space and time. Must be accompanied by the `HEAT_CONDUCTION = GENERALIZED` option.
4.52.12 THERMAL CONDUCTIVITY = USER_FUNCTION (TENSOR)

Parameters

- NAME_xx = STRING
- NAME_yy = STRING
- [NAME_zz = STRING]
- X = STRING

Example

begin definition for function SI_K11
  type is piecewise linear
  begin values
    20.0  5.50e7
    100.0 4.60e7
    ...
    800.0 1.30e7
    2000.0 1.30e7
  end values
end definition for function SI_K11

begin definition for function SI_K22
  type is piecewise linear
  begin values
    20.0  8.50e7
    100.0 7.60e7
    ...
    800.0  0.30e7
    2000.0 0.30e7
  end values
end definition for function SI_K22

... 

Begin Aria Material Foo

... 

  Tensor Thermal Conductivity = User Function Name_xx=SI_K11 Name_xx=SI_K22 X=Temperature

... 

End Aria Material Foo

Description

A look-up function is used to compute the values of the thermal conductivity as a function of some other variable, i.e. \( f(x) \). The function type (“piecewise linear” in the example above) must support the \texttt{differentiate()} method for Newton’s method.

Here \texttt{NAME_xx}, \texttt{NAME_yy}, \texttt{NAME_zz}, are the names of user-defined functions for (CONDUCTIVITY_DATA, SI_K11, SI_K22 in the example) and \texttt{X} is the Aria name of the abscissa variable (TEMPERATURE in the example). \texttt{NAME_zz} is required only for three-dimensional analysis. Note that \texttt{X} is not necessarily the same name as the optional abscissa variable identified in the user-defined function. Must be accompanied by the \texttt{HEAT_CONDUCTION = GENERALIZED} option.
4.52.13 THERMAL CONDUCTIVITY = CALORE_USER_SUB (TENSOR)

Parameters

- NAME = STRING
- TYPE = STRING
  - [MULTIPLIER = REAL]
  - [NR = INT]
  - [R0 = INT]
  - [R1 = INT(etc.)]
  - [NI = INT]
  - [I0 = INT]
  - [I1 = INT(etc.)]

Example

Tensor Thermal Conductivity = Calore_User_Sub name=w80afftuser
type=element_tensor NR=2 R0=1000 R1=0.7

Description

The Calore_User_Sub Tensor model allows the user to explicitly evaluate thermal conductivity as a function of temperature and position within an element. Additionally, a user_query feature allows evaluation of thermal conductivity based upon other problem criteria. NAME is the name of the user subroutine was registered with. NR and NI are the numbers of real and integer parameters, respectively. Each real (int) parameter is supplied by the Rn (In) parameter. Note: the parameters use zero based counting. The TYPE parameter denotes the API the user subroutine uses; currently Aria supports type ELEMENT_TENSOR for tensor valued thermal conductivity.

The optional MULTIPLIER parameter (defaults to 1.0) can be supplied to scale the output of the user subroutine, i.e., the flux \( f = mf_{user} \) where \( m \) is the multiplier and \( f_{user} \) is the output of the user subroutine.

See section 9.2.2 for a more complete example of a Calore user subroutine.

Must be accompanied by the HEAT_CONDUCTION = GENERALIZED option.

4.53 THERMAL DIFFUSIVITY

THERMAL DIFFUSIVITY = MODEL[param = val1, param2 = val2 ...]

Description

Specifies the material model for the thermal diffusivity.

Summary

Specifies the material model for the thermal diffusivity.

Parent Block(s) ARIA MATERIAL
4.53.1 THERMAL DIFFUSIVITY = CALORE_USER_SUB

Parameters

- NAME = STRING
- TYPE = STRING
- [MULTIPLIER = REAL]
- [NR = INT]
- [R0 = INT]
- [R1 = INT(etc.)]
- [NI = INT]
- [I0 = INT]
- [I1 = INT(etc.)]

Example

Thermal Diffusivity = Calore_User_Sub name=w80afftuser type=element NR=2 R0=1000 R1=0.7

Description

NAME is the name of the user subroutine was registered with. NR and NI are the numbers of real and integer parameters, respectively. Each real (int) parameter is supplied by the Rn (In) parameter. Note: the parameters use zero based counting. The TYPE parameter denotes the API the user subroutine uses; currently Aria only supports types ELEMENT and NODE.

The optional MULTIPLIER parameter (defaults to 1.0) can be supplied to scale the output of the user subroutine, i.e., the flux \( f \) will be \( f = mf_{user} \) where \( m \) is the multiplier and \( f_{user} \) is the output of the user subroutine.

See section 9.2.2 for a more complete example of a Calore user subroutine.

4.53.2 THERMAL DIFFUSIVITY = CONSTANT

Parameters

- D = REAL

Example

THERMAL DIFFUSIVITY = CONSTANT D = 1.0

Description

D is the value of the constant thermal diffusivity.

4.53.3 THERMAL DIFFUSIVITY = ENCORE_FUNCTION

Parameters

- NAME = STRING
- [EVAL_TYPE = STRING]

Example

THERMAL DIFFUSIVITY = Encore_Function Name=My_Function

Description

Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```plaintext
Begin String Function My_Function
    Value is "200 + 1.0*t"
End
```
Then, this function can be used as shown in the example above. The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

4.54 TOTAL INTERNAL ENERGY

TOTAL INTERNAL ENERGY = MODEL [param1 = val1, param2 = val2 ...]

Description Specifies a model for the total internal energy of a material.

Summary Specifies a model for the total internal of a material.

Parent Block(s) ARIA MATERIAL

4.54.1 TOTAL INTERNAL ENERGY = ENCORE_FUNCTION

Parameters NAME = STRING
[EVAL_TYPE = STRING]

Example Total Internal Energy = Encore_Function Name=My_Function

Description Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

Begin String Function My_Function
  Value is "200 + 1.0*t"
End

Then, this function can be used as shown in the example above. The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

4.54.2 TOTAL INTERNAL ENERGY = POROUS

Parameters (none)

Example Total Internal Energy = Porous
The total internal energy $e$ is computed as

$$ e = (1 - \phi)\rho_s e_s + \phi \rho_f e_f $$

(4.121)

where $\phi$ is the porosity, $\rho_s$ is the density of the solid porous skeleton, $e_s$ is the internal energy of the solid porous skeleton, $\rho_f$ is the density of the fluid phase and $e_f$ is the internal energy of the fluid phase.

### 4.55 VALENCE

VALENCE = MODEL[param1 = val1, param2 = val2 ...]

**Description**
Specifies the valence (net charge) for a species.

**Summary**
Specifies the valence (net charge) for a species.

**Parent Block(s)**
ARIA MATERIAL

#### 4.55.1 VALENCE = CONSTANT

**Parameters**

$Z = \text{REAL}$

[SUBINDEX = INT]

**Example**

Valence = Constant Subindex=1 $Z = 1$
Valence = Constant Subindex=2 $Z = -1$
Valence = Constant Subindex=5 $Z = -2$

**Description**

$Z$ is the value of the species valence.

SUBINDEX is the species subindex.

#### 4.55.2 VALENCE = ENCORE_FUNCTION

**Parameters**

NAME = STRING

[EVAL_TYPE = STRING]

**Example**

Valence = Encore_Function Name=My_Function

**Description**

Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```
Begin String Function My_Function
    Value is "200 + 1.0*t"
End
```
Then, this function can be used as shown in the example above.
The optional argument `EVAL_TYPE` can be used to select the particular function on the Encore function object. Valid options include `VALUE`, `DOT`, `GRADIENT`, `FLUX` and `STRESS`. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – `VALUE` for `NO_OP` expressions, `DOT` for `DT_OP` expressions and `GRADIENT` for `GRAD_OP` expressions.

4.56 VISCOSITY

VISCOSITY = `MODEL`[param = val1, param2 = val2 ...]

Description  Specifies the material model for the fluid viscosity.

Summary  Specifies the material model for the fluid viscosity.

Parent Block(s)  ARIA MATERIAL

4.56.1 VISCOSITY = ARRHENIUS

Parameters  

- `mu0` = `REAL`
- `E` = `REAL`

Example  

VISCOSITY = Arrhenius mu0=16.4 E=5000.

Description  This model provides a viscosity with an Arrhenius temperature dependence:

\[ \mu = \mu_0 e^{-E/T} \]  

where \( T \) is the temperature.

4.56.2 VISCOSITY = ARRHENIUS_CARREAU

Parameters  

- `MU_ZERO` = `REAL`
- `[MU_INF = `REAL`]`
- `[A = `REAL`]`
- `N` = `REAL`
- `[LAMBDA = `REAL`]`
- `K` = `REAL`
- `T_REF` = `REAL`
- `[SKIP_SENSITIVITIES = `INT`]`

Example  

Viscosity = Arrhenius_Carreau ...

Description  This viscosity model is a function of the shear rate (Carreau model) and temperature (Arrhenius model). The two contributions are combined multiplicatively.

\[ \mu = \left[ \mu_\infty + (\mu_0 - \mu_\infty) \left(1 + (\lambda \dot{\gamma})^a\right)^{\frac{n-1}{a}} \right] e^\frac{\lambda}{\lambda_{\text{ref}}} \]  

where \( \lambda \) is the shear rate and \( \lambda_{\text{ref}} \) is the reference shear rate.

(4.123)
where $\mu_\infty$ is the infinite shear viscosity ($\text{MU\_INF}$, defaults to zero), $\mu_0$ is the zero shear viscosity ($\text{MU\_ZERO}$), $n$ ($\text{N}$) and $a$ ($\text{A}$, defaults to 2) are model parameters, $\dot{\gamma}$ is the shear rate, $\lambda$ ($\text{LAMBDA}$) is a time constant (defaults to 1), $T_{\text{ref}}$ is a reference temperature and $K$ is an Arrhenius constant.

If the optional flag \texttt{SKIP\_SENSITIVITIES} is nonzero then no Newton sensitivities will be included.

### 4.56.3 \texttt{VISCOSITY = BINGHAM\_WLF}

**Parameters**

- \texttt{MU\_ZERO = REAL}
- \texttt{MU\_INF = REAL}
- \texttt{F = REAL}
- \texttt{N = REAL}
- \texttt{A = REAL}
- \texttt{LAMBDA = REAL}
- \texttt{TAU\_Y = REAL}
- \texttt{[SKIP\_SENSITIVITIES = INT]}

**Example**

\texttt{VISCOSITY = Bingham\_WLF ...}

**Description**

\[
\mu = \mu_\infty + \left(\mu_0 - \mu_\infty + \tau_y \frac{1 - e^{-\frac{\dot{\gamma} F}{\dot{\gamma}}}}{\dot{\gamma}}\right) \left(1 + (\lambda \dot{\gamma})^a\right)^{\frac{n-1}{a}}
\]

(4.124)

where $\dot{\gamma}$ is the shear rate.

If the optional flag \texttt{SKIP\_SENSITIVITIES} is nonzero then no Newton sensitivities will be included.

### 4.56.4 \texttt{VISCOSITY = BINGHAM\_WLFT}

**Parameters**

- \texttt{MU\_ZERO = REAL}
- \texttt{MU\_INF = REAL}
- \texttt{F = REAL}
- \texttt{N = REAL}
- \texttt{A = REAL}
- \texttt{LAMBDA = REAL}
- \texttt{TAU\_Y = REAL}
- \texttt{C\_1 = REAL}
- \texttt{C\_2 = REAL}
- \texttt{T\_REF = REAL}
- \texttt{[SKIP\_SENSITIVITIES = INT]}

**Example**

\texttt{VISCOSITY = Bingham\_WLFT ...}

**Description**

\[
\mu = a_T \left(\mu_\infty + \left(\mu_0 - \mu_\infty + \tau_y \frac{1 - e^{-a_T \dot{\gamma} F}}{a_T \dot{\gamma}}\right) \left(1 + (a_T \lambda \dot{\gamma})^a\right)^{\frac{n-1}{a}}\right)
\]

(4.125)

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where

\[ \alpha_T = e^{\frac{c_1(T - T_0)}{T - T_0}} \] (4.126)

and \( T \) is the temperature and \( \dot{\gamma} \) is the shear rate.

If the optional flag SKIP_SENSITIVITIES is nonzero then no Newton sensitivities will be included.

### 4.56.5 VISCOSITY = CARREAU

**Parameters**

- \texttt{MU_ZERO = REAL}
- \texttt{[MU_INF = REAL]}
- \texttt{[A = REAL]}
- \texttt{N = REAL}
- \texttt{LAMBDA = REAL}
- \texttt{[SKIP_SENSITIVITIES = INT]}

**Example**

Viscosity = Carreau ...

**Description**

\[
\frac{\mu - \mu_\infty}{\mu_0 - \mu_\infty} = (1 + (\lambda \dot{\gamma})^a)^{\frac{n-1}{n}}
\] (4.127)

or

\[
\mu = \mu_\infty + (\mu_0 - \mu_\infty)(1 + (\lambda \dot{\gamma})^a)^{\frac{n-1}{n}}
\] (4.128)

where \( \mu_\infty \) is the infinite shear viscosity (\texttt{MU_INF}, defaults to zero), \( \mu_0 \) is the zero shear viscosity (\texttt{MU_ZERO}), \( n \) (\texttt{N}) and \( a \) (\texttt{A}, defaults to 2) are model parameters, \( \dot{\gamma} \) is the shear rate and \( \lambda \) (\texttt{LAMBDA}) is a time constant.

If the optional flag SKIP_SENSITIVITIES is nonzero then no Newton sensitivities will be included.

### 4.56.6 VISCOSITY = CARREAU_T

**Parameters**

- \texttt{MU_ZERO = REAL}
- \texttt{[MU_INF = REAL]}
- \texttt{[A = REAL]}
- \texttt{N = REAL}
- \texttt{K = REAL}
- \texttt{[SKIP_SENSITIVITIES = INT]}

**Example**

Viscosity = Carreau_T ...

**Description**

\[
\frac{\mu - \mu_\infty}{\mu_0 - \mu_\infty} = \left(1 + \left(e^{k/T \dot{\gamma}}\right)^a\right)^{\frac{n-1}{n}}
\] (4.129)

or

\[
\mu = \mu_\infty + (\mu_0 - \mu_\infty)\left(1 + \left(e^{k/T \dot{\gamma}}\right)^a\right)^{\frac{n-1}{n}}
\] (4.130)

where \( \mu_\infty \) is the infinite shear viscosity (\texttt{MU_INF}, defaults to zero), \( \mu_0 \) is the zero shear viscosity (\texttt{MU_ZERO}), \( n \) (\texttt{N}) and \( a \) (\texttt{A}, defaults to 2) are model parameters and \( \dot{\gamma} \) is the shear rate. The
quantity $e^{k/T}$, where $T$ is temperature and $k$ (K) is a reference temperature, is a temperature dependent time scale; it takes the place of the constant $\lambda$ time scale in the CARREAU model.

If the optional flag SKIP_SENSITIVITIES is nonzero then no Newton sensitivities will be included.

### 4.56.7 VISCOSITY = CONSTANT

**Parameters**

$\mu = \text{REAL}$

**Example**

$\text{VISCOSITY = CONSTANT } \mu = 1.0$

**Description**

$\mu$ is the value of the constant fluid viscosity.

### 4.56.8 VISCOSITY = CURING_FOAM

**Parameters**

$\text{VFRAC_SUBINDEX} = \text{INT}$

$\text{EXTENT_SUBINDEX} = \text{INT}$

$\text{PHI_ZERO} = \text{REAL}$

$[A = \text{REAL}]$

$[B = \text{REAL}]$

$[C = \text{REAL}]$

$[\text{KSI}_C = \text{REAL}]$

**Example**

Viscosity = Curing_Foam Vfrac_Subindex=1 Extent_Subindex=2 Phi_Zero=0.45

**Description**

For a curing epoxy with volume fraction $\phi$ and extent of reaction $\xi$ the viscosity is given by

$$\mu = \mu_o \exp \frac{\phi_o - \phi}{1 - \phi_o + \phi}$$

(4.131)

where $\mu_o$ is given by

$$\mu_o = (a - bT) \left( \frac{\xi^2}{\xi_i^2} \right)^c$$

(4.132)

where $T$ is the temperature. The remaining parameters $a$, $b$, $c$ and $\xi_c$ have default values of $a = 20$, $b = 0.22$, $c = -4/3$ and $\xi_c = 0.45$ though they can be overridden with the optional model parameters.

**NOTE:** The volume fraction is assumed to be a SPECIES field with the subindex provided by the VFRAC_SUBINDEX parameter. Likewise, the extent of reaction field is assumed to be a SPECIES field with the subindex provided by the EXTENT_SUBINDEX parameter.

### 4.56.9 VISCOSITY = ENCORE_FUNCTION

**Parameters**

$\text{NAME} = \text{STRING}$

$[\text{EVAL_TYPE} = \text{STRING}]$
Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```
Begin String Function My_Function
  Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

### 4.56.10 VISCOSITY = KRIEGER

**Parameters**

- BETA = *REAL*
- PHI_MAX = *REAL*
- MU_S = *REAL*

**Example**

```
VISCOSITY = KRIEGER BETA = -1.65, PHI_MAX = 1.0, MU_S = 1.0
```

**Description**

In the viscosity model of [6]

\[
\mu = \mu_s \left(1 - \frac{\phi}{\phi_m}\right)^\beta
\]

(4.133)

BETA is the Krieger exponent, PHI_MAX is the maximum suspension concentration and MU_S is the solvent viscosity.

### 4.56.11 VISCOSITY = POLYNOMIAL

**Parameters**

- VARIABLE = *STRING*
- ORDER = *INT*
- [VARIABLE_OFFSET = *REAL*]
- [C0 = *REAL*]
- [C1 = *REAL*]
  ...
- [CN = *REAL*]

**Example**

```
Viscosity= Polynomial Variable=Temperature Order=1 C0=401.0 C1=88.5
```

**Description**

Arbitrary order polynomial function of a specified scalar variable.

\[
\mu = \sum_{i=0}^{N} C_i (X + X_0)^i
\]

(4.134)
Here, $N$ is the order of the polynomial provided by the ORDER parameter and $X$ is the variable supplied by the VARIABLE parameter, $X_o$ is an optional offset (VARIABLE_OFFSET, default is zero) and $C_i$ are the supplied coefficients. Coefficients that are not supplied default to a value of zero. The VARIABLE argument can be TIME or any internal Expression that evaluates to a scalar. For the latter case, the format of the VARIABLE argument is described in section 2.6.1.

4.56.12 VISCOSITY = POWER_LAW

Parameters

$K = \text{REAL}$

$N = \text{REAL}$

Example

Viscosity = Power_Law $K=0.8$ $N=0.5$

Description

The viscosity is proportional to the shear rate, $\dot{\gamma}$ raised to some power, e.g.,

$$\mu = k^{n}$$

where $k$ ($K$) and $n$ ($N$) are model parameters.

4.56.13 VISCOSITY = THERMAL

Parameters

$[A = \text{REAL}]$

$[B = \text{REAL}]$

$[C = \text{REAL}]$

$[D = \text{REAL}]$

Example

VISCOSITY = THERMAL $A=1750$ $C=0.12$ $D=0$

Description

This model is simply a cubic polynomial in temperature where the viscosity is given by

$$\mu = A + BT + CT^2 + DT^3$$

where $T$ is the temperature.

4.56.14 VISCOSITY = USER_FUNCTION

Parameters

$\text{NAME} = \text{STRING}$

$X = \text{STRING}$
begin definition for function Water_Viscosity
# Source Appendix 2 from "Transport Processes and
# Unit Operations" by C. J. Geankoplis
# type is piecewise linear
begin values
# K Pa*s (or cP)
273.15 1.7921
275.15 1.6728
277.15 1.5674
279.15 1.4728
281.15 1.3860
283.15 1.3077
285.15 1.2363
287.15 1.1709
289.15 1.1111
291.15 1.0559
293.15 1.0050
293.35 1.0000
295.15 0.9579
297.15 0.9142
299.15 0.8737
301.15 0.8360
303.15 0.8007
305.15 0.7679
307.15 0.7371
309.15 0.7085
311.15 0.6814
313.15 0.6560
315.15 0.6321
317.15 0.6097
319.15 0.5883
321.15 0.5683
323.15 0.5494
325.15 0.5315
327.15 0.5146
329.15 0.4985
331.15 0.4832
333.15 0.4688
335.15 0.4550
337.15 0.4418
339.15 0.4293
341.15 0.4174
343.15 0.4061
345.15 0.3952
347.15 0.3849
349.15 0.3750
351.15 0.3665
353.15 0.3565
355.15 0.3478
357.15 0.3395
359.15 0.3315
361.15 0.3239
363.15 0.3165
365.15 0.3095
367.15 0.3027
369.15 0.2962
371.15 0.2899
373.15 0.2838
A look-up function is used to compute the values of the viscosity as a function of some other variable, i.e. \( f(x) \). The function type (“piecewise linear” in the example above) must support the `differentiate()` method for Newton’s method.

Here `NAME` is the name of the user-defined function (`Water_Viscosity` in the example) and `X` is the Aria name of the abscissa variable (`TEMPERATURE` in the example). Note that `X` is not necessarily the same name as the abscissa variable identified in the user-defined function (`T` in the example).

### 4.56.15 VISCOSITY = WELD

**Parameters**

- \( \text{BETA} = \text{REAL} \)
- \( C0 = \text{REAL} \)
- \( C1 = \text{REAL} \)
- \( C2 = \text{REAL} \)
- \( C3 = \text{REAL} \)
- \( T\_LIQ = \text{REAL\_T\_90} = \text{REAL\_T\_MAX} = \text{REAL} \)

**Example**

Viscosity = Weld C0=1 C1=-1e-2 C2=0 C3=0 T\_LIQ=920 T\_MAX=1400 T\_90=1000

**Description**

This is an empirical model that emulates the melting of a solid metal during the laser welding process.

\[
\mu = \begin{cases} 
\mu_90 + (\mu_{liq} - \mu_90) \frac{T - T_{liq}}{T_{liq} - T_{90}} & : T < T_{liq} \\
 c_0 + c_1 \hat{T} + c_2 \hat{T}^2 + c_3 \hat{T}^3 & : T \geq T_{liq}
\end{cases}
\]  

(4.137)

where \( \mu_{liq} \) is given by

\[
\mu_{liq} = c_0 + c_1 T_{liq} + c_2 T_{liq}^2 + c_3 T_{liq}^3,
\]  

(4.138)

\( \mu_90 = \beta \mu_{liq} \) and \( \hat{T} = \min(T, T_{max}) \). The default value of \( \text{BETA} \) is \( 10^{11} \).

### 4.57 FLOWING LIQUID VISCOSITY

FLOWING LIQUID VISCOSITY = MODEL[\( \text{param}_1 = \text{val}_1, \text{param}_2 = \text{val}_2 \ldots \)]

**Description**

Specifies the material model for the fluid viscosity.

**Summary**

Specifies the material model for the fluid viscosity in the Brinkman momentum equation.

**Parent Block(s)**

ARIA MATERIAL
4.57.1 FLOWING LIQUID VISCOSITY = CONSTANT

Parameters \( \mu = \text{REAL} \)

Example FLOWING LIQUID VISCOSITY = CONSTANT \( \mu = 1.0 \)

Description \( \mu \) is the value of the constant fluid viscosity.

4.58 VOLUME FRACTION GAS

\[ \text{VOLUME FRACTION GAS} = \text{MODEL}[\text{param}_1 = \text{val}_1, \text{param}_2 = \text{val}_2 \ldots] \]

Description Specifies the material model for the volume fraction of gas.

Summary Specifies the material model for the volume fraction of gas.

Parent Block(s) ARIA MATERIAL

4.58.1 VOLUME FRACTION GAS = CONSTANT

Parameters \( \text{value} = \text{REAL} \)

Example Volume Fraction Gas = Constant \( \text{value} = 1.0 \)

Description \( \text{value} \) is the volume fraction of gas.

4.58.2 VOLUME FRACTION GAS = FROM_MASS_FRACTIONS

Parameters None

Example Volume Fraction Gas = From_Mass_Fractions

Description For a multiphase ChemEq model, this model uses

\[ \phi = 1 - \rho \sum_{i} Y_i \rho_{b,i} \]  \hspace{1cm} (4.139)

where \( \phi \) is the gas volume fraction, \( \rho \) is the overall density, \( \rho_{b,i} \) is the bulk density of the \( i^{th} \) solid species, and \( Y_i \) is the mass fraction of the \( i^{th} \) solid species. This model uses the species phase definitions from the ChemEq definition to determine which species are solid. As such, the model can only be used in a material block which also contains a ChemEq definition.
This uses expressions for the mass fractions, so its material block must also include “Mass_Fraction of X = From_ChemEq” for each solid species, or some comparable definition for mass fraction.

You must also provide an expression for the bulk density of each solid species.

4.58.3 VOLUME FRACTION GAS = ENCORE_FUNCTION

Parameters

- **NAME** = STRING
  - [EVAL_TYPE = STRING]

Example

 volume_fraction_gas = encore_function name=my_function

Description

Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```
Begin String Function My_Function
  Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

4.58.4 VOLUME FRACTION GAS = POLYNOMIAL

Parameters

- **VARIABLE** = STRING
- **ORDER** = INT
  - [VARIABLE_OFFSET = REAL]
  - [C0 = REAL]
  - [C1 = REAL]
  - ...
  - [CN = REAL]

Example

 volume_fraction_gas = polynomial variable=temperature order=1 C0=401.0 C1=88.5

Description

Arbitrary order polynomial function of a specified scalar variable.

\[
\kappa = \sum_{i=0}^{N} C_i (X + X_0)^i \quad (4.140)
\]

Here, \( N \) is the order of the polynomial provided by the ORDER parameter and \( X \) is the variable supplied by the VARIABLE parameter, \( X_0 \) is an optional offset (VARIABLE_OFFSET, default is zero) and \( C_i \) are the supplied coefficients. Coefficients that are not supplied default to a value of zero. The VARIABLE argument can be TIME or any internal Expression that evaluates to a scalar. For the latter case, the format of the VARIABLE argument is described in section 2.6.1.
4.58.5 VOLUME FRACTION GAS = USER_FUNCTION

Parameters

NAME = STRING
X = STRING

Example
begin definition for function SI_phi
  type is piecewise linear
  begin values
    20.0 5.50e7
    100.0 4.60e7
    ...
    800.0 1.30e7
    2000.0 1.30e7
  end values
end definition for function SI_phi

Begin Aria Material Foo
  ...
  Volume Fraction Gas = User_Function Name=SI_phi X=Temperature
  ...
End Aria Material Foo

Description

A look-up function is used to compute the values of the thermal conductivity as a function of
some other variable, i.e. \( f(x) \). The function type ("piecewise linear" in the example above)
must support the \texttt{differentiate}() method for Newton’s method.

Here \texttt{NAME} is the name of the user-defined function (\texttt{SI\_phi}, in the example) and \texttt{X} is the Aria
name of the abscissa variable (\texttt{TEMPERATURE} in the example). Note that \texttt{X} is not necessarily
the same name as the optional abscissa variable identified in the user-defined function.

4.59 YOUNGS MODULUS

YOUNGS MODULUS = \texttt{MODEL[\texttt{param}_1 = \texttt{val}_1, \texttt{param}_2 = \texttt{val}_2 \ldots]} 

Description

Specifies the material model for the Young’s modulus.

Summary

The solid stress \( \mathbf{T} \) is given by

\[
\mathbf{T} = \lambda E_{kk} \mathbf{I} + 2\mu \mathbf{E} - \beta (T - T_{\text{ref}}) \mathbf{I}
\]  

(4.141)

where \( \lambda \) and \( \mu \) are the Lamé coefficients, \( \mathbf{E} = \frac{1}{2} \left( \nabla \mathbf{d} + \nabla \mathbf{d}^T \right) \) is the deformation tensor, \( \beta \) is the coefficient of thermal stress, \( T \) is temperature and \( T_{\text{ref}} \) is the thermal strain reference temperature.

These Lamé coefficients are related to the more standard Young’s modulus, Poisson’s ratio
and CTE \((\alpha)\) as follows:

\[
\begin{align*}
2\mu &= \frac{E}{1 + \nu} \quad (4.142) \\
\lambda &= \frac{\nu E}{(1 + \nu)(1 - 2\nu)} = 2\mu \frac{\nu}{(1 - 2\nu)} \quad (4.143) \\
\beta &= \frac{\alpha E}{(1 - 2\nu)} = \alpha \frac{3\lambda + 2\mu}{(1 + \nu)(1 - 2\nu)} \quad (4.144)
\end{align*}
\]

When a user supplies the Young’s modulus, Poisson’s ratio and CTE properties Aria internally converts them into the Lamé coefficients.

Supplying the Lamé coefficients is more computationally efficient but perhaps less convenient, especially if the material properties are varying (e.g., temperature dependent in a non-isothermal problem).

Parent Block(s)  \text{ARIA MATERIAL}

### 4.59.1 YOUNG'S MODULUS = CONSTANT

**Parameters**  
YM = \text{REAL}

**Example**  
YOUNG'S MODULUS = CONSTANT YM = 1.0

**Description**  
YM is the value of the constant Young’s modulus.

### 4.59.2 YOUNG'S MODULUS = ENCORE_FUNCTION

**Parameters**  
NAME = \text{STRING}  
[EVAL_TYPE = \text{STRING}]

**Example**  
YOUNG'S MODULUS = Encore_Function Name=My_Function

**Description**  
Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```
Begin String Function My_Function
  Value is "200 + 1.0*t"
End
```

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include \text{VALUE}, \text{DOT}, \text{GRADIENT}, \text{FLUX} and \text{STRESS}. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – \text{VALUE} for \text{NO_OP} expressions, \text{DOT} for \text{DT_OP} expressions and \text{GRADIENT} for \text{GRAD_OP} expressions.
Chapter 5

Equations Aria Solves

5.1 Generalized Conservation Equation

We first introduce a general conservation equation, as a model for the specific equations that Aria solves, demonstrating how the Galerkin finite element method is applied to it, and how the integration by parts is carried out on its individual terms. Following [7], the conservation of a general scalar quantity $b(x,t)$, with units of amount-per-unit-volume, at a point $x$ and time $t$ can be expressed as

$$\frac{\partial b}{\partial t} + \nabla \cdot (bv) = -\nabla \cdot f + B_V$$

(5.1)

where $v$ is the mass average velocity, $f$ is the diffusive flux of $b$, and $B_V$ is the volumetric source of $b$.

The Galerkin FEM (G/FEM) residual form of 5.1 is formed by bringing the right hand side terms to the left, multiplying by the FEM weight function $\phi^i$ and integrating over the volume $V$,

$$R_b^i = \int_V \left( \frac{\partial b}{\partial t} + v \cdot \nabla b + b \nabla \cdot v + \nabla \cdot f - B_V \right) \phi^i \, dV = 0.$$  

(5.2)

In many applications $\nabla \cdot v = 0$ so we ignore that term from here on. However, it is straightforward to account for this term via the source term $B_V$. Using the vector identity $(\nabla \cdot f)\phi^i = \nabla \cdot (f \phi^i) - \nabla \phi^i \cdot f$ and using the divergence theorem, 5.2 becomes

$$R_b^i = \int_V \left[ \left( \frac{\partial b}{\partial t} + v \cdot \nabla b - B_V \right) \phi^i - \nabla \phi^i \cdot f \right] \, dV + \int_S n \cdot f \phi^i \, dS = 0.$$  

(5.3)

Here $n$ is a unit normal along the boundary $S$, pointing out of the volume $V$.

Equation 5.3 embodies the sign convention for sources, fluxes and equation terms used within Aria. For example, scalar flux expressions in Aria provide values for $f_n \equiv n \cdot f$ and should be positive for a flux of $b$ leaving the volume $V$.

Note also that we have not assigned a units convention to the equation. Any unit system may be employed in the specification of the individual terms in 5.1. However, each term in 5.1 must have overall units of $[b] / [time]$, and the overall residual expression has units of $[b] \cdot [L]^3 / [time]$, where $[b]$ are units of the conserved quantity, $b$, $[L]$ is the unit of the length scale, and $[time]$ is the unit for time.

5.2 Conservation of Mass

For a material with density $\rho$, letting $b = \rho$ results in the conservation of mass. Since there is no net flow relative to the mass average velocity $f = 0$. Although there are no sources of mass, having such a source
can be convenient in modeling and simulation; so, we let the mass source be $B_V = q_m$. Thus, (5.1) becomes

$$\frac{\partial \rho}{\partial t} + \rho \nabla \cdot v + v \cdot \nabla \rho = q_m.$$  \hfill (5.4)

For the special but common case of constant density, this reduces to

$$\nabla \cdot v = 0.$$  \hfill (5.5)

Using equation 5.3, the G/FEM residual form is

$$R_P^i = \int_V \left( -\frac{\partial \rho}{\partial t} - \rho \nabla \cdot v - v \cdot \nabla \rho + q_m \right) \phi^i \, dV = 0. \quad (5.6)$$

**Important Note:** Equation 5.6 has been multiplied by -1 because this form results in a better linear system for the special case of incompressible flow. This is important to remember when defining mass source terms.

In Aria, each term in 5.6 is specified separately as identified in equation 5.7.

$$R_P^i = \int_V -\frac{\partial \rho}{\partial t} \phi^i \, dV + \int_V -(v \cdot \nabla \rho + \rho \nabla \cdot v) \phi^i \, dV + \int_V q_m \phi^i \, dV = 0 \quad (5.7)$$

For a purely incompressible form, Aria offers the alternative form given in 5.8;

$$R_P^i + \int_V -\nabla \cdot v \phi^i \, dV + \int_V q_m \phi^i \, dV = 0 \quad (5.8)$$

### 5.3 Conservation of Energy

For a material with constant density and specific heat $C_p$, temperature $T$, heat flux $q$ and volumetric energy source $H_V$, letting $b = \rho C_p T$, $f = q$ and $B_V = H_V$ results in the conservation of energy.

$$\rho C_p \frac{\partial T}{\partial t} + \rho C_p v \cdot \nabla T = -\nabla \cdot q + H_V. \quad (5.9)$$

A common constitutive relationship for $q$ is Fourier’s law, $q = -\kappa \nabla T$ where $\kappa$ is the thermal conductivity. However, we leave the heat flux as an option to be specified as part of the material properties (see section 4.15). Using equation 5.3, the G/FEM residual form is

$$R_T^i = \int_V \left[ (\rho C_p \frac{\partial T}{\partial t} + \rho C_p v \cdot \nabla T - H_V) \phi^i - \nabla \phi^i \cdot q \right] \, dV + \int_S q_n \phi^i \, dS = 0 \quad (5.10)$$

where $q_n$ is the heat flux at the boundary. For example, the natural convection boundary condition gives $q_n = h(T - T_\infty)$ where $h$ is the heat transfer coefficient and $T_\infty$ is the bulk temperature away from the surface.
In Aria, each term in 5.10 is specified separately as identified in equation 5.11.

$$R^i_T = \int_V \rho C_p \frac{\partial T}{\partial t} \phi^i \, dV + \int_V \rho C_p \mathbf{v} \cdot \nabla T \phi^i \, dV - \int_V H_T \phi^i \, dV - \int_V \nabla \phi^i \cdot \mathbf{q} \, dV + \int_S q_n \phi^i \, dS = 0 \quad (5.11)$$

More and more often we need to account for variable density problems and so we need to bring back some of the terms we threw away because we were going to assume $\nabla \cdot \mathbf{v} \equiv 0$. Here’s a do-over of equation 5.11 that accommodates a variable density through the DIV term:

$$R^i_T = \int_V \rho C_p \frac{\partial T}{\partial t} \phi^i \, dV + \int_V \rho C_p \mathbf{v} \cdot \nabla T \phi^i \, dV + \int_V \rho C_p T \nabla \cdot \mathbf{v} \phi^i \, dV - \int_V H_T \phi^i \, dV - \int_V \nabla \phi^i \cdot \mathbf{q} \, dV + \int_S q_n \phi^i \, dS = 0 \quad (5.12)$$

Note, however, that equation 5.12 still assumes a constant specific heat $C_p$.

### 5.3.1 SUPG

SUPG stabilization can be activated by added the SUPG term on the input file line. This capability is still under development.

### 5.4 Conservation of Chemical Species

For a material with species $k$ with molar concentration $C_k$, molar flux $J_k$ relative to the mass average velocity and volumetric reaction rate $R_{V,k}$, letting $b = y_k$, $f = J_k$ and $B_V = R_{V,k}$ in (5.1) results in the conservation equation for species $k$,

$$\frac{\partial C_k}{\partial t} + \mathbf{v} \cdot \nabla C_k = -\nabla \cdot \mathbf{J}_k + R_{V,k}. \quad (5.13)$$

For liquid mixtures which are dilute in all species except one, Fick’s law is often used to approximate $J_k$. In this approximation, $D_k$ represents the diffusion coefficient of species $k$ with respect to the concentrated species and it is assumed that the interactions between dilute species is assumed negligible. Again, however, we choose to leave the governing equation in the more general form and require the particular diffusive flux model as user input (see section 4.46). Using equation 5.3, the G/FEM residual form is

$$R_{C_k} = \int_V \left[ \left( \frac{\partial C_k}{\partial t} + \mathbf{v} \cdot \nabla C_k - R_{V,k} \right) \phi^i - \nabla \phi^i \cdot \mathbf{J}_k \right] \, dV + \int_S q_{n,k} \phi^i \, dS = 0 \quad (5.14)$$

where $q_{n,k}$ is the mass flux at the boundary. For example, the natural convection boundary condition gives $q_n = k(C_k - C_{\infty,k})$ where $k$ is the mass transfer coefficient and $C_{\infty,k}$ is the bulk concentration away from the surface.
In Aria, each term in 5.14 is specified separately as identified in equation 5.15.

\[
R_{C_k}^i = \int_V \frac{\partial C_k}{\partial t} \phi_i^j \, dV + \int_V v \cdot \nabla C_k \phi_i^j \, dV - \int_V R_{V,k} \phi_i^j \, dV
\]

\[
\text{MASS} \quad \text{ADV} \quad \text{SRC}
\]

\[ - \int_V \nabla \phi_i^j \cdot J_k \, dV + \int_S q_{n,k} \phi_i^j \, dS = 0 \quad (5.15)
\]

More and more often we need to account for variable density problems and so we need to bring back some of the terms we threw away because we were going to assume \( \nabla \cdot v \equiv 0 \). Here’s a do-over of equation 5.15 that accommodates a variable density through the DIV term:

\[
R_{C_k}^i = \int_V \frac{\partial C_k}{\partial t} \phi_i^j \, dV + \int_V v \cdot \nabla C_k \phi_i^j \, dV + \int_V C \nabla \cdot v \phi_i^j \, dV
\]

\[
\text{MASS} \quad \text{ADV} \quad \text{DIV}
\]

\[ - \int_V R_{V,k} \phi_i^j \, dV - \int_V \nabla \phi_i^j \cdot J_k \, dV + \int_S q_{n,k} \phi_i^j \, dS = 0 \quad (5.16)
\]

Often times it is useful to solve for mass, weight or volume fractions of each species rather than for the concentration directly. In that case, an additional condition exists,

\[ \sum_k C_k = 1 \quad (5.17) \]

Using this condition, it is only necessary to solve for \( N - 1 \) species fractions where \( N \) is the total number of species present in the problem. The final species, then, is simply given as

\[ C_j = 1 - \sum_{k \neq j} C_k \quad (5.18) \]

This method can be triggered in Aria by specifying the equation term FRACBAL. In this case, the equation for \( C_j \) is not included in the system of unknowns but is instead post-processed on the fly. Aria will automatically detect all other species equations and include them in the fraction balance.

### 5.5 Conservation of Fluid Momentum

The Cauchy momentum equation is given by

\[ \rho \frac{\partial v}{\partial t} + \rho v \cdot \nabla v - g - \nabla \cdot T = 0 \quad (5.19) \]

where \( T \) is the fluid stress tensor and \( g \) is a body force. We construct the G/FEM residual form of 5.19 by contracting with the unit coordinate vector in the \( k \)-direction, \( e_k \), multiplying by the weight function \( \phi_i^j \) and integrating over the volume. Using the vector identity \( (\nabla \cdot T) \cdot e_k \phi^i = \nabla \cdot (T \cdot e_k \phi^i) - T^i \cdot \nabla (e_k \phi^i) \) and integrating by parts gives

\[
R_{m,k}^i = \int_V \left[ \left( \frac{\partial v}{\partial t} + \rho v \cdot \nabla v - g \right) \cdot e_k \phi^i + T^i \cdot \nabla (e_k \phi^i) \right] \, dV - \int_S n \cdot T \cdot e_k \phi^i \, dS = 0 \quad (5.20)
\]
In Aria, each term in 5.20 is specified separately as identified in equation 5.15.

\[ R_{m,k}^i = \int_V \rho \frac{\partial \mathbf{v}}{\partial t} \cdot \mathbf{e}_k \phi^i \, dV + \int_V \rho \mathbf{v} \cdot \nabla \mathbf{v} \cdot \mathbf{e}_k \phi^i \, dV - \int_V g \cdot \mathbf{e}_k \phi^i \, dV + \int_V T^i : \nabla (\mathbf{e}_k \phi^i) \, dV - \int_S \mathbf{n} \cdot \mathbf{T} \cdot \mathbf{e}_k \phi^i \, dS = 0 \]  

(5.21)

### 5.6 Conservation of Solid Momentum

Aria currently solves the quasistatic form of the solid momentum equations. Furthermore, the solid stress is treated as a linear elastic material. In this limit, the Cauchy momentum equation is given by

\[ \nabla \cdot \mathbf{T} = 0 \]  

(5.22)

where \( \mathbf{T} \) is the solid stress tensor. We construct the G/FEM residual form of 5.19 by contracting with the unit coordinate vector in the \( k \)-direction, \( \mathbf{e}_k \), multiplying by the weight function \( \phi^i \) and integrating over the volume. Using the vector identity \( (\nabla \cdot \mathbf{T}) \cdot \mathbf{e}_k \phi^i = \nabla \cdot (\mathbf{T} \cdot \mathbf{e}_k \phi^i) - \mathbf{T}^i : \nabla (\mathbf{e}_k \phi^i) \) and integrating by parts gives

\[ R_{m,k}^i = \int_V T^i : \nabla (\mathbf{e}_k \phi^i) \, dV = 0 \]  

(5.23)

Here, the surface contribution, \( \int_S \mathbf{n} \cdot \mathbf{T} \cdot \mathbf{e}_k \phi^i \, dS \), has been dropped because Aria currently only supports Dirichlet and natural (homogeneous) boundary conditions for the solid equation.

In Aria, each term in 5.23 is specified separately as identified in equation 5.24.

\[ R_{m,k}^i = \int_V T^i : \nabla (\mathbf{e}_k \phi^i) \, dV = 0 \]  

(5.24)

Currently, Aria does not support direct specification of the more popular stress-strain parameterization that utilizes Young’s modulus \( E \), Poisson’s ratio \( \nu \) and coefficient of thermal expansion \( \alpha \) (note, the shear modulus \( G = \mu \)). The relationship between these two parametrizations is summarized here for convenience.

\[ 2\mu = \frac{E}{(1 + \nu)} \]  

(5.25)

\[ \lambda = \frac{\nu E}{(1 + \nu) (1 - 2\nu)} = 2\mu \frac{\nu}{(1 - 2\nu)} \]  

(5.26)

\[ \beta = \frac{\alpha E}{(1 - 2\nu)} = \alpha (3\lambda + 2\mu) \]  

(5.27)

When the solid momentum equation is solved, the physical coordinates of the region, used to calculate integration quantities, is deformed by the displacement vector. The current configuration, \( \mathbf{x} \), is calculated from the reference configuration, \( \mathbf{X} \), by

\[ \mathbf{x} = \mathbf{X} + \mathbf{d}, \]  

(5.28)
where \( \mathbf{d} \) is the displacement field. The displacement field is chosen automatically by the code, but may be overridden by setting a solution option. These displaced, physical coordinates are used for integration of all of the equations.

Additionally, the code supports the integration of the solid momentum equations in the reference configuration, rather than in the current configuration. This is activated through the use of the `REF_DIFF` equation term rather than the `DIFF` term. When this is used, the Cauchy stress tensor is rotated into the Second Piola-Kirchhoff stress tensor. Integration quantities are also calculated in the reference configuration. Care should be taken to ensure that the constitutive model used also changes its strain measure to one which is work conjugate with the Second Piola-Kirchhoff stress tensor.

If both the MESH and SOLID equations are defined in the same region, then the TALE mesh motion algorithm is activated automatically. Here, the reference frame for the SOLID equation is changed to the TALE reference frame, which is defined as

\[
\mathbf{x}_r = \mathbf{X} + \mathbf{d}_m - \mathbf{d}_s
\]  

(5.29)

where \( \mathbf{x}_r \) is the new TALE reference frame, \( \mathbf{d}_m \) is MESH_DISPLACEMENTS, and \( \mathbf{d}_s \) is SOLID_DISPLACEMENTS. This affects the calculation of strain measures, such as \( \mathbf{F} \), and integration quantities, such as \( \det(\mathbf{J}) \), among others. More details of the TALE method will be available soon in a pending report.

### 5.7 Voltage Equation

The electric potential or voltage \( V \) is frequently used in determining the electric field, \( \mathbf{E} = -\nabla V \). While (5.1) cannot be applied to the voltage, the equation governing the voltage – Gauss’ law from Maxwell’s equations – has a similar form. Writing the electric displacement \( \mathbf{D} \) as \( \mathbf{D} = \varepsilon \mathbf{E} \), where \( \varepsilon \) is the electric permittivity, Gauss’ law is

\[
- \nabla \cdot \varepsilon \nabla V = \rho_e
\]  

(5.30)

where the permittivity is taken to be a constant and \( \rho_e \) is the volumetric free charge density.

Using equation 5.3, the G/FEM residual form is

\[
R^i_V = \int_V (-\rho_e \phi^i + \nabla \phi^i \cdot \varepsilon \nabla V) \, \mathrm{d}V + \int_S q_n \phi^i \, \mathrm{d}S = 0
\]  

(5.31)

In Aria, each term in 5.31 is specified separately as identified in equation 5.32.

\[
R^i_V = -\int_V \rho_e \phi^i \, \mathrm{d}V + \underbrace{\int_V \nabla \phi^i \cdot \varepsilon \nabla V \, \mathrm{d}V}_\text{DIFF} + \int_S q_n \phi^i \, \mathrm{d}S = 0
\]  

(5.32)

### 5.8 Current Equation

An alternate formulation for solving for the electrical potential (see section 5.7) is to solve the “current” equation which is a conservation equation for electrical charge. The electrical current \( \mathbf{J} \) is frequently related to the electric field \( \mathbf{E} \) using Ohm’s law as \( \mathbf{J} = \sigma_e \mathbf{E} \) where \( \sigma_e \) is the electrical conductivity. The electric potential or voltage \( V \) is used in determining the electric field, \( \mathbf{E} = -\nabla V \). However, we choose to leave the electrical current as a more general constitutive model to be provided as a material model input (see
section 4.6). More complex version for charged ion fluxes (Nerst Planck, etc. can be utilized with the species equation).

\[- \nabla \cdot J = S\]  \hspace{1cm} (5.33)

Here \(S\) is a current source term, for example from electrochemical reactions in a battery simulation.

Using equation 5.3, the G/FEM residual form is

\[R^i_V = \int_V (-S\phi^i - \nabla \phi^i \cdot J) \, dV + \int_S q_n \phi^i \, dS = 0\]  \hspace{1cm} (5.34)

In Aria, each term in 5.34 is specified separately as identified in equation 5.35.

\[R^i_V = - \int_V S\phi^i \, dV - \int_V \nabla \phi^i \cdot J \, dV + \int_S q_n \phi^i \, dS = 0\]  \hspace{1cm} (5.35)

### 5.9 Charge Density Equation

This equation solves for the volumetric free charge density, \(\rho_c\), and is meant to be coupled with the voltage equation in section 5.7). This equation can also be used in place of the current equation 5.33, since this equation has the correct form (the current equation has the correct form with \(\rho_c\) when electric displacement is activated) and the charge density is treated as a variable. The charge density evolves as

\[\frac{\partial \rho_c}{\partial t} + v \cdot \nabla \rho_c = - \nabla \cdot J = \nabla \cdot (\sigma \nabla V)\]  \hspace{1cm} (5.36)

where \(J = -\sigma \nabla V\) is assumed to obey Ohm’s law and \(\sigma\) represents the conductivity.

The G/FEM residual form of this equation is

\[R^i_V = \int_V \left( \frac{\partial \rho_c}{\partial t} \phi^i + \nabla \phi^i \cdot v + \nabla \phi^i \cdot \sigma \nabla V \right) \, dV + \int_S q_n \phi^i \, dS = 0\]  \hspace{1cm} (5.37)

In Aria, each term in 5.37 is specified separately as identified in equation 5.38.

\[R^i_V = \int_V \frac{\partial \rho_c}{\partial t} \phi^i \, dV + \int_V \nabla \phi^i \cdot v \, dV + \int_V \nabla \phi^i \cdot \sigma \nabla V \, dV + \int_S q_n \phi^i \, dS = 0.\]  \hspace{1cm} (5.38)

### 5.10 Suspension Equation

In treating the suspension as a continuum, we introduce an evolution equation for particle volume fraction, \(\phi\), as

\[\frac{\partial \phi}{\partial t} + v \cdot \nabla \phi + \nabla \cdot N = 0.\]  \hspace{1cm} (5.39)

The particle volume fraction is defined as the total summed volume of particles per volume of the particle medium. 5.39 represents a balance between the stored particles, the convected particle flux, and the diffusive particle flux, \(N\). Several mechanisms which include Brownian motion, sedimentation, hydrodynamic particle interactions, and gradients in suspension viscosity may contribute to the diffusive particle flux. Specification of the appropriate flux model must then be carried out to close the definition of the conservation equation.
5.11 Porous Flow Equations

In what follows we use a subscript $\alpha$ to denote a component index and $N_{\alpha}$ is the total number of components. We use a subscript $\beta$ to denote a phase and the total number of phases is $N_{\beta}$. The subscript $s$ is used to refer to the solid phase of the porous skeleton; this phase is not included in $N_{\beta}$. Summations over $\alpha$ and $\beta$ are assumed to range from one to $N_{\alpha}$ and $N_{\beta}$, respectively, unless otherwise specified.

5.11.1 Generalized Equations

This write up follows, directly, the reports of [8], [9] and [10].

Porosity and Void Ratio

In modeling of porous materials, one often needs to know the relative volume of pore space ($V_p$) and the solid porous skeleton ($V_s$). The most common measure of this is the porosity, $\phi$, which is defined as the ratio of the volume of pore space ($V_p$) per unit total volume ($V_t = V_p + V_s$), viz.,

$$\phi = \frac{V_p}{V_t} = \frac{V_p}{V_p + V_s}. \quad (5.40)$$

The void ratio ($r$) is sometimes used as well and is defined as the ratio of the pore volume to the solid volume, viz.,

$$r = \frac{V_p}{V_s}. \quad (5.41)$$

Clearly the porosity and void ratio are related,

$$r = \frac{\phi}{1 - \phi} \quad (5.42)$$

$$\phi = \frac{r}{1 + r} \quad (5.43)$$

Component Mass Balances

The mass balance for each component across all phases can be written

$$\frac{\partial d_{\alpha}}{\partial t} + \nabla \cdot F_{\alpha} = Q_{\alpha} \quad \alpha = 1, \ldots, N_{\alpha} \quad (5.44)$$

where $d_{\alpha}$ is mass concentration of component $\alpha$ across all phases, $t$ is time, $F_{\alpha}$ is the total mass flux of component $\alpha$ relative to a fixed frame of reference and $Q_{\alpha}$ is the volumetric source of component $\alpha$.

The mass concentration of component $\alpha$ can be written as

$$d_{\alpha} = \phi \sum_{\beta} Y_{\alpha\beta} \rho_{\beta} S_{\beta} \quad (5.45)$$

where $\phi$ is the interstitial volume fraction, a.k.a. the porosity, that ranges from zero to one, $Y$ denotes mass fraction so that $Y_{\alpha\beta}$ is the mass fraction of component $\alpha$ in phase $\beta$, $\rho_{\beta}$ is mixture density of phase $\beta$ and $S_{\beta}$ is saturation which is defined as the fraction of the interstitial pore space occupied by phase $\beta$. 

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For completeness it is noted here that the mass fractions sum to one in each phase and that the saturations of all phases sum to unity,

\[ \sum_{\alpha} Y_{\alpha\beta} = 1 \]  \hspace{1cm} (5.46)

\[ \sum_{\beta} S_{\beta} = 1. \]  \hspace{1cm} (5.47)

The macroscopic mass flux in phase \( \beta \), \( \rho_{\beta}v_{\beta} \) is obtained from the extended Darcy's Law,

\[ \rho_{\beta}v_{\beta} = f_{\beta} = -\frac{\rho_{\beta}\kappa_{\beta}}{\mu_{\beta}} K \cdot (\nabla P_{\beta} + \rho_{\beta}g) \]  \hspace{1cm} (5.48)

where \( \kappa_{\beta} \) is the relative permeability of phase \( \beta \) and \( K \) is the permeability tensor which are both intrinsic properties of the porous material and so may vary spatially. The mixture viscosity of phase \( \beta \) is \( \mu_{\beta} \), \( P_{\beta} \) is the pressure in phase \( \beta \) and \( g \) is the gravitational acceleration.

The component mass flux, \( F_\alpha \) can be written as the sum of the component mass fluxes across each phase,

\[ F_\alpha = \sum_{\beta} F_{\alpha\beta}. \]  \hspace{1cm} (5.49)

Within each phase, the component mass flux can be decomposed in its convective and diffusive components,

\[ F_{\alpha\beta} = Y_{\alpha\beta} f_{\beta} + J_{\alpha\beta} \]  \hspace{1cm} (5.50)

where \( J_{\alpha\beta} \) is the diffusive flux of component \( \alpha \) in phase \( \beta \). For example, for a gaseous, dilute binary mixture the diffusive flux may be modeled using Fick's Law,

\[ J_{\alpha\beta} = -\rho_{\beta}D_{\alpha\beta}\nabla Y_{\alpha\beta} \]  \hspace{1cm} (dilute binary gas mixture) (5.51)

where \( D_{\alpha\beta} \) is the diffusion coefficient of component \( \alpha \) in phase \( \beta \).

**Porous Species**

The porous species equation is identical to the mass balance equation, with all quantities being molar concentrations. The porous species equation is written as

\[ \frac{\partial c_\alpha}{\partial t} + \nabla \cdot \Pi_\alpha = R_\alpha \quad \alpha = 1, \ldots, N_\alpha, \]  \hspace{1cm} (5.52)

where \( c_\alpha \) is the molar concentration, defined analogously to (5.45), \( \Pi_\alpha \) is the molar flux, and \( R_\alpha \) is the volumetric molar source of component \( \alpha \). The porous species equation exists to facilitate using expressions derived for use with the non porous-flow species equation. An analogous set of terms to those found in the mass balance equation can be derived on a molar concentration basis.

**Energy Balance**

In order to accommodate multicomponent transport problems we employ an enthalpy based energy conservation law,

\[ \frac{\partial e}{\partial t} + \nabla \cdot q = Q_e \]  \hspace{1cm} (5.53)
where $e$ is the total internal energy, $q$ is the total heat flux relative to a stationary frame of reference and $Q_e$ is the volumetric source of energy. The total internal energy is written as a sum of the energies of all components across all phases as well as the internal energy of the solid porous structure,

$$e = (1 - \phi) \rho_s e_s + \phi \sum_{\beta} S_{\beta} \rho_{\beta} e_{\beta}$$  \hspace{1cm} (5.54)

where $\rho_{\beta}$ and $e_{\beta}$ are the mixture density and internal energy of phase $\beta$. Generally, thermodynamic relations are required to relate these quantities to the component specific quantities.

The total heat flux can be decomposed into different modes of transport. Typically,

$$q = -\lambda_T \nabla T + \sum_{\beta} \sum_{\alpha} h_{\alpha \beta} F_{\alpha \beta}$$  \hspace{1cm} (5.55)

where $\lambda_T$ is the effective thermal conductivity, $T$ is the temperature and $h_{\alpha \beta}$ is the partial enthalpy of component $\alpha$ in phase $\beta$ which is interpreted as the enthalpy of pure component $\alpha$ at the temperature and pressure of phase $\beta$. Using (5.50), we can expand (5.55) such that

$$\sum_{\alpha} h_{\alpha \beta} F_{\alpha \beta} = \sum_{\alpha} h_{\alpha \beta} (Y_{\alpha \beta} f_{\beta} + J_{\alpha \beta})$$

$$= h_{\beta} f_{\beta} + \sum_{\alpha} h_{\alpha \beta} J_{\alpha \beta}$$  \hspace{1cm} (5.56)

where $h_{\beta} = \sum_{\alpha} h_{\alpha \beta} Y_{\alpha \beta}$ is the mixture enthalpy of phase $\beta$. Thus, the total heat flux can be expressed as the sum of conductive, convective and diffusive contributions,

$$q = -\lambda_T \nabla T + \sum_{\beta} \rho_{\beta} v_{\beta} h_{\beta} + \sum_{\beta} \sum_{\alpha} h_{\alpha \beta} J_{\alpha \beta}$$  \hspace{1cm} (5.57)

### 5.11.2 Porous Flow Systems

To facilitate various flavors of single phase and multiphase flow in Aria, the porous flow system model was created to manage expression creation. In addition to the single phase model (sections 5.11.2 and 5.11.2), there are several others available including: mixed single phase, CO2-salt-brine (section 5.11.2), two-phase-immiscible (sections 5.11.2 and 5.11.2), air-water (section 5.11.2), and organic-material-decomposition (section 5.11.2). The basic purpose of the porous flow system model is to create the necessary expressions for each system. There are examples of each flow system in the regression tests, which are the best place to start when developing an input for a similar system. The input syntax for a porous flow system is given in section 4.1 in addition to the required parameters for each porous flow system.

#### Single Phase, Single Component, Isothermal Flow

The simplest possible case is for a single phase flow ($N_\beta = 1$, $S = 1$) with a single component ($N_\alpha = 1$, $Y = 1$) at a fixed temperature. This is the single phase porous flow system. For a single component flow, there is no diffusive mass flux so $J = 0$ and $F = f$ where

$$F = f = \rho v = -\frac{\rho \kappa_r}{\mu} K \cdot (\nabla P + \rho g)$$  \hspace{1cm} (5.58)

In this case, the mass balance equation is all that is needed to solve for the unknown pressure field $P$.

$$\frac{\partial d}{\partial t} + \nabla \cdot f = Q$$  \hspace{1cm} (5.59)

Since the density is most likely a function of temperature, the constant uniform temperature will have to be specified. Finally, in this special case, (5.45) reduces to $d = \phi \rho$. 

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Single Phase, Single Component, Non-isothermal Flow

Building on the simple case developed in section 5.11.2, the next step is to accommodate a non-isothermal flow. Here, we still have a single phase flow ($N_\beta = 1$, $S = 1$) with a single component ($N_\alpha = 1$, $Y = 1$). For a single component flow, there is no diffusive mass flux so $J = 0$ and $F = f$ where

$$ F = f = \rho v = -\frac{\rho k_r}{\mu} K \cdot (\nabla P + \rho g) $$

(5.60)

Again, the mass balance equation is all that is needed to solve for the unknown pressure field $P$.

$$ \frac{\partial d}{\partial t} + \nabla \cdot f = Q $$

(5.61)

Again, in this special case, (5.45) reduces to $d = \phi \rho$.

In order to solve for the unknown temperature, $T$, we solve an energy transport equation in the enthalpy form as described in section 5.11.1. Here, the energy flux in equation (5.55) reduces to

$$ q = -\lambda T \nabla T + \rho v h $$

(5.62)

where $h$ is the pure component enthalpy which may be a function of $P$ and $T$.

Two-Phase Immiscible

In this section, we define a formulation for two-phase flow of immiscible fluids, that is, without mass transfer between the phases. Common examples include oil/water, under pressure and temperature conditions where phase transitions are negligible, and systems involving non-aqueous phase liquids (NAPL) occurring in subsurface contamination by organic liquids, e.g., solvents, fuels, oils. If we disallow partitioning of components across phases, then we get the following system of mass balance equations for a wetting and non-wetting phase,

$$ \frac{\partial}{\partial t} (\rho_w \phi (1 - S_n)) = \nabla \cdot \left( \rho_w \frac{k_{rw}}{\mu_w} K \cdot (\nabla p - \rho_w g) \right) + Q_w $$

(5.63)

$$ \frac{\partial}{\partial t} (\rho_n \phi S_n) = \nabla \cdot \left( \rho_n \frac{k_{rn}}{\mu_n} K \cdot (\nabla p + \nabla p_c - \rho_n g) \right) + Q_n $$

(5.64)

In these equations, the subscript w represents wetting phase properties and the subscript n nonwetting phase properties. We have also incorporated the Darcy flux terms, including pressure and gravitational forces, as well as capillary pressure:

$$ p_c = p_n - p_w $$

(5.65)

The symbol $p$ in the equations is used in place of wetting phase pressure, $p_w$. Also, the pore space is assumed saturated with fluids,

$$ S_w + S_n = 1 $$

(5.66)

This constraint has been incorporated in the wetting phase mass balance. In addition to the density models described in the previous section, models for density and viscosity can be specified as constant, polynomial with respect to other primary or secondary variables, or by tabular functions, or a user-defined plug-in.

Both the capillary pressure and relative permeability are functions of phase saturations. Either of these can be specified by the user in a tabular form, as a function of either phase saturation. A plug-in can also be written by the user for either of these functions. Several commonly used functions are built-in to the code.
Pressure-pressure Formulation for Two-phase Immiscible

The \( P_w - S_n \) formulation is generally known to have superior numerical behavior for \( S_w \to 0 \), however, the capillary pressure in heterogeneous media creates a problem for a vertex-centered (e.g. FEM) numerical scheme. In FEM, because a node straddles a material interface, which separates two materials with different texture and therefore different wetting properties, the capillary pressure is continuous across the interface, but the saturation is not. So, a \((P_w, S_n)\) formulation in FEM would require dealing with discontinuous DOF for \( S_n \). One alternative is to instead solve for two pressures. Two pairs of pressure-based DOFs available in Aria include \((P_w, P_c)\) and \((P_w, P_n)\). (Note that in single phase regions, where \( S_w = 0 \) or 1, then \( P_w \) or \( P_n \) are undefined. The formulation should work for the thermal battery problem so long as \( S_w > 0 \).

The following system of mass balance equations for a wetting and non-wetting phase describes an immiscible flow,

\[
\frac{\partial}{\partial t} (\rho_w \phi S_w) = \nabla \cdot \left( \rho_w \left( \frac{k_{rw}}{\mu_w} \boldsymbol{K} \cdot \nabla P_w - \rho_w \mathbf{g} \right) \right) + Q_w \tag{5.67}
\]

\[
\frac{\partial}{\partial t} (\rho_n \phi S_n) = \nabla \cdot \left( \rho_n \left( \frac{k_{rn}}{\mu_n} \boldsymbol{K} \cdot \nabla P_w + \nabla P_c - \rho_n \mathbf{g} \right) \right) + Q_n \tag{5.68}
\]

In these equations, the subscript \( w \) represents wetting phase properties and the subscript \( n \) non-wetting phase properties. We have also incorporated the Darcy flux terms, including pressure and gravitational forces, as well as capillary pressure:

\[
p_c = p_n - p_w
\]

Also, the pore space is assumed saturated with fluids,

\[
S_w + S_n = 1
\]

The current implementation in Aria uses a van Genuchten function to model the relationship between the capillary pressure and the saturation. This relationship is inverted for satisfaction given the capillary pressure in the pressure-pressure formulation.

Two-phase Air-Water

This model describes the nonisothermal two-component, two-phase transport of water and air in a porous medium. The air is treated as a non-condensible gas, but it can dissolve in the liquid phase. The water can exist as water vapor or liquid. The model allows the fluids to partition between the two phases, depending on pressure and temperature conditions. Two single phase states (all liquid or all gas) and one two-phase state are allowed.

The component mass balance equations describing two-phase transport of water, air and energy are:

\[
\frac{\partial}{\partial t} (\phi (S_l Y_{wl} \rho_l + S_g Y_{wg} \rho_g)) + \nabla \cdot (S_l \rho_l \mathbf{v}_l + S_g \rho_g \mathbf{v}_g + \mathbf{J}_{wg}) + Q_w = 0 \tag{5.71}
\]

\[
\frac{\partial}{\partial t} (\phi (S_l Y_{al} \rho_l + S_g Y_{ag} \rho_g)) + \nabla \cdot (S_l \rho_l \mathbf{v}_l + S_g \rho_g \mathbf{v}_g + \mathbf{J}_{ag}) + Q_a = 0 \tag{5.72}
\]

\[
\frac{\partial}{\partial t} \left[ (1 - \phi) \rho_s \mathbf{e}_s + \phi (\rho_l \mathbf{e}_l + \rho_g S_g \mathbf{e}_g) \right] + \nabla \cdot (-\lambda_T \nabla T + \rho_l \mathbf{v}_l h_l + \rho_g \mathbf{v}_g h_g + h_{wg} \mathbf{J}_{wg} + h_{ag} \mathbf{J}_{ag}) + Q_e = 0 \tag{5.73}
\]

where

\[
\mathbf{J}_{ag} = -\rho_g D_{wa} \nabla Y_{ag} \tag{5.74}
\]

is the gas-phase binary diffusion flux for water vapor through air. In these equations, subscript \( l \) refers to the liquid phase and \( g \) to the gas phase. In addition to previously defined variables: \( Y_{\alpha\beta} \) is the mass fraction of component \( \alpha \) (\( w \) for water, \( a \) for air) in phase \( \beta \), \( S_\beta \) is the saturation of phase \( \beta \), \( \rho_\beta \) is the phase density,
$e_\beta$ is the internal energy of phase $\beta$, $\lambda_T$ is the effective thermal heat conduction, $h_\beta$ is the enthalpy of phase $\beta$ and $h_{ag}$ is the component enthalpy in the gas phase.

The liquid and gas phase Darcy velocities are

$$v_l = -\frac{k_{rl}}{\mu_l} \mathbf{K} \cdot (\nabla P_l + \rho_l \mathbf{g})$$  \hspace{1cm} (5.75)$$

$$v_g = -\frac{k_{rg}}{\mu_g} \mathbf{K} \cdot (\nabla P_g + \rho_g \mathbf{g})$$  \hspace{1cm} (5.76)$$

In addition, the following constraints hold:

$$S_l + S_g = 1 ; \quad Y_{w,\beta} + Y_{a,\beta} = 1$$  \hspace{1cm} (5.77)$$

**CO2-salt-brine**

The CO2-salt-brine system is intended to model the interaction of CO2 gas, solid salt, and liquid brine. In this system there are three mass balance equations that must be solved for the saturations of each. This formulation describes a 3-component 2-phase flow for the water-CO2-NaCl system. The two phases are brine, denoted by subscript $w$, a liquid phase of water with salt and CO2 dissolved in it, and “gas” denoted by subscript $g$, a CO2-rich phase, with some water dissolved in it, but no salt. The formulation also deals with precipitation of salt from the liquid phase.

To deal with the precipitation of NaCl from the liquid to form solid salt, the fluid system is treated as a three-phase system, with two flowing phases composed of the brine and CO2 phases, and one non-flowing solid phase, which is the solid salt precipitate. Obviously, the formulation holds for other chloride salts, e.g. CaCl, KCl.

In the equations above, we have used the following nomenclature, $S_s$ is the saturation of solid salt precipitate, $S_w$ is the brine phase saturation, $S_g$ is the CO2-rich phase saturation, $x_w$ is the mass fraction of water in the brine phase, $x_{CO2}$ is the mass fraction of CO2 in the brine phase, $x_{NaCl}$ is the mass fraction of salt dissolved in the brine phase, $y_{CO2}$ is the mass fraction of CO2 in the CO2-rich (“gas”) phase, $y_w$ is the mass fraction of water in the gas phase, $\rho_s(T, p)$ is the density of solid precipitate, and $J^a_\beta$ is the diffusion flux of component $\alpha$ in phase $\beta$ which can also include dispersion. The subscript ‘w’ refers to the brine phase, ‘g’ refers to the CO2 phase and ‘s’ refers to the solid salt phase.

This model treats all the separate-phase CO2 as a single CO2-rich phase. The phase diagram for the CO2-H2O system displays a saturation line where liquid water, liquid CO2 and a vapor phase can co-exist. The saturated water vapor pressure on this line is small, so the vapor phase is mostly CO2.

$$S_w + S_g + S_s = 1 ; \quad x_w + x_{CO2} + x_{NaCl} = 1 ; \quad y_w + y_{CO2} + y_{NaCl} = 1$$  \hspace{1cm} (5.81)$$

We also have the constraint on the over-all composition, which can replace either of the last two constraints above,
\[ z_w + z_{CO2} + z_{NaCl} = 1 \] (5.82)

To solve for the pressure, in all compositional situations, we can solve a pressure equation, obtained by summing all component balances. Since the resulting equation is not independent, it replaces one of the three mass balance equations presented earlier.

\[ \frac{\partial}{\partial t} \phi(S_w \rho_w + S_g \rho_g + S_s \rho_s) + \nabla \cdot (\rho_w \mathbf{v}_w + \rho_g \mathbf{v}_g) + Q^{H2O} + Q^{CO2} + Q^{NaCl} = 0 \] (5.83)

In the numerical implementation, this equation could be obtained after all BCs have been applied to the discrete equations, by summing the three residual equations together.

Recall, the primary variables for this system are defined as a set of variables that allow one to compute all the remaining (secondary) variables appearing in the mass and energy balance equations. For this problem, the primary solution vector (assuming we will also solve an energy balance for \( T \)) is \([P, z_{CO2}, \rho_{NaCl}, T]\), where

\[ z_{CO2} = \frac{S_w \rho_w x_{CO2} + S_g \rho_g y_{CO2}}{S_w \rho_w + S_g \rho_g} = (1 - \nu)x_{CO2} + \nu y_{CO2} \] (5.84)

is the over-all mass fraction of CO2 in the flowing phases. We have introduced the fraction of gas, defined as,

\[ \nu = \frac{\rho_g S_g}{\rho_w S_w + \rho_g S_g} \] (5.85)

Similarly, the overall density of salt is defined by,

\[ \rho_{NaCl} = \rho_w S_w x_{NaCl} + \rho_s S_s = (1 - \nu)x_{NaCl} F + \rho_s S_s \] (5.86)

where we have introduced, for later convenience, the following mass density,

\[ F = \rho_w S_w + \rho_g S_g \] (5.87)

Note that the accumulation term in the CO2 balance can be written as \( \phi F z_{CO2} \).

In this formulation, \( P \) is solved for from the pressure equation, \( z_{CO2} \) from the CO2 mass balance equation, \( \rho_{NaCl} \) from the NaCl mass balance and \( T \) from the energy balance equation. We assume the phase equilibrium for this system is done ahead of time.

The thermodynamics algorithm for this system is still in development and has not yet been completed.

**Organic Material Decomposition (OMD)**

Organic material decomposition (OMD) models currently define two-phase (solid and gas) problems in which the solid (or porous) species react to produce additional gas species that pressurize a system and volumetric heat transfer between phases. The notation used above does not include the porous solid as a phase, but since for OMD we are solving equations for the solid phase including reactions, we set \( N_\beta = 2 \), with it being understood that one of these phases is the solid phase. OMD problems will extend to include a liquid phase in the future. The OMD equations described here are based on the combustion model [11]. Further details about coupling boundary conditions between fluid-only and porous regions are provided in section 9.4.
There are separate mass balance equations for the solid phase and gas phase. The solid phase mass balance equation is used to solve for solid phase density:

\[
\frac{\partial \bar{\rho}}{\partial t}_{\text{MASS}} = -\frac{\dot{\omega}_{f,g}^m}{S_{RC}},
\]

(5.88)

where \( \bar{\rho} \) is the solid phase density and \( \dot{\omega}_{f,g}^m \) is the rate of formation of gases due to heterogeneous reaction kinetics. The mass balance equation for the gas phase solves for the pressure (from the gas phase density via the ideal gas law),

\[
\frac{\partial (\psi \rho_g)}{\partial t}_{\text{MASS}} + \frac{\partial (\rho_g u_{j,g})}{\partial x_j}_{\text{ADV}} = \frac{\dot{\omega}_{f,g}^m}{S_{RC}},
\]

(5.89)

where \( \rho_g \) is the gas-phase density and \( \psi \) is the mixture-averaged solid-phase porosity. The porosity is the void space not taken up by the volume fractions of each solid phase species, typically defined as a function of the volume fractions of each solid species,

\[
\psi = 1 - \sum_{\alpha} X_{\alpha}.
\]

(5.90)

The volume fraction of each solid species is calculated as

\[
X_{\alpha} = \frac{\bar{\rho} Y_{\alpha}}{\rho_{\text{so,}\alpha}},
\]

(5.91)

where \( \rho_{\text{so,}\alpha} \) is the solid (non-porous) density for each species. In equation 5.89, \( u_{j,g} \) is the gas-phase velocity and gets computed from Darcy’s approximation as

\[
\psi v_{j,g} = u_{j,g} = -\frac{\hat{K}_{ij}}{\mu_g} \left( \frac{\partial \rho_g}{\partial x_i} + \rho_g g_i \right),
\]

(5.92)

where \( \hat{K}_{ij} \) is the mixture-averaged solid-phase permeability tensor defined as the normalized volume average of the solid phase species’ permeabilities,

\[
\hat{K}_{ij} = \frac{\sum_{\alpha} X_{\alpha} K_{\alpha j}}{\sum_{\alpha} X_{\alpha}},
\]

(5.93)

where

\[
K_{\alpha j} = K_{\text{ref}} \left( \frac{T}{T_{\text{ref}}} \right)^n \delta_{\alpha j}
\]

(5.94)

The intrinsic permeability is the product of the intrinsic permeability scaling and the intrinsic permeability. The reason it is separated in this way is that scalar models have access to many generic functions, whereas tensor entries can only be constants. A more complex function for the intrinsic permeability tensor would have to be written as a user subroutine.

In equation 5.92 the parameter \( \mu_g \) is the gas-phase viscosity and \( g_j \) is the gravity vector. The ideal gas law gets used for the gas density in the pressure form of equation 5.89

\[
\rho_g = \frac{PM}{RT_g}.
\]

(5.95)

Therefore the actual equation for pressure being solved here is

\[
\frac{\partial}{\partial t} \left( \frac{\rho_g \bar{M} \psi}{RT_g} \right) + \frac{\partial}{\partial x_j} \left[ \rho_g \bar{M} K_{\alpha j} \left( \frac{\partial \rho_g}{\partial x_j} + \rho_g \frac{\bar{M}}{RT_g} g_j \right) \right] = 0.
\]

(5.96)
where \( M \) is the mass averaged molecular weight,
\[
M = \sum_{\alpha} Y_{\alpha} M_{\alpha},
\]
(5.97)
where \( Y_{\alpha} \) are the gas phase mass fractions and \( M_{\alpha} \) are the molecular weights of the gaseous species.

The solid phase species equations solve for the solid phase mass fractions, \( Y_{\alpha,s} \), of species \( \alpha \)
\[
\frac{\partial (\bar{\rho} Y_{\alpha})}{\partial t}_{\text{MASS}} + \frac{\partial (\bar{\rho}_g u_{j,g} Y_{\alpha,g})}{\partial x_j}_{\text{ADV}} + \frac{\partial q_{\alpha,j}^{Y,g}}{\partial x_j}_{\text{DIFF}} = \omega_{\alpha,g}^{Y,g}
\]
where \( \omega_{\alpha,g}^{Y,g} \) is the formation rate of species \( \alpha \) and \( \omega_{\alpha,g}^{Y,g} \) is its destruction rate. The gas-phase species equations solve for the gas-phase mass fraction \( Y_{\alpha,g} \) of species \( \alpha \)
\[
\frac{\partial (\bar{\rho} Y_{\alpha,g})}{\partial t}_{\text{MASS}} + \frac{\partial (\bar{\rho}_g u_{j,g} Y_{\alpha,g})}{\partial x_j}_{\text{ADV}} + \frac{\partial q_{\alpha,j}^{Y,g}}{\partial x_j}_{\text{DIFF}} = \omega_{\alpha,g}^{Y,g}
\]
(5.99)
where \( D_{\alpha,g} \) is the gas-phase mass diffusivity for species \( \alpha \)

Enthalpy equations are used to solve for temperature, and OMD type problems allow for different temperature fields in the gas and solid phases, so these are described separately here.

The solid phase enthalpy equation for the solid phase temperature \( \bar{T} \) is given by
\[
\frac{\partial (\bar{\rho} c_p \bar{T})}{\partial t}_{\text{MASS}} = - \frac{\partial q_{j}^{h}}{\partial x_j}_{\text{DIFF}} + h_{cv} (T_g - \bar{T})
\]
(5.101)
where \( \kappa \) is the thermal conductivity.

The gas-phase enthalpy equation within a porous region, to be solved for the gas-phase enthalpy (and corresponding temperature) \( h_g = c_p T_g \), is
\[
\frac{\partial (\bar{\rho} p_g h_g)}{\partial t}_{\text{MASS}} + \frac{\partial (\bar{\rho} p_g u_{j,g} h_g)}{\partial x_j}_{\text{ADV}} = - \frac{\partial q_{j}^{h,g}}{\partial x_j}_{\text{DIFF}} + \frac{\partial (\bar{\rho} p_g)}{\partial t}_{\text{SRC}} + h_{cv} (\bar{T} - T_g)
\]
(5.103)
where \( h_g \) is the mixture-averaged gas-phase enthalpy, \( h_{cv} \) is the volumetric heat transfer coefficient which gets set outside of the solid/gas phases of the material definition, \( \bar{T} \) is the porous solid-phase temperature and \( T_g \) is the gas phase temperature. The gas-phase energy diffusive flux vector \( q_{j}^{h,g} \) is modeled as
\[
q_{j}^{h,g} = - \psi p_g D_g \frac{\partial h_g}{\partial x_j}
\]
(5.104)
5.11.3 Porosity Models

There are several models for porosity in the porous flow capabilities in Aria. The following list provides a brief description for each.

Mesh Deforming

In the mesh (or solid) deforming porosity model, the expression for the porosity is given as

\[
\phi = 1.0 - \frac{b(1.0 - \phi_0)}{\det(F)}
\]

(5.105)

where \(\phi_0\) is the initial porosity, \(b\) is the Biot coefficient, and \(F\) is the mesh deformation gradient.

Rock Compressible

For the Rock Compressible porosity model, the porosity is given as

\[
\phi = \phi_0 (1.0 + C_r (p - p_{ref}))
\]

(5.106)

where again, \(\phi_0\) is the initial porosity, \(C_r\) is a rock compressibility coefficient, \(p\) is the pore pressure, and \(p_{ref}\) is a reference pore pressure defined in the input parameters.

Coussy

For the Coussy porosity model, the expression for porosity is

\[
\phi = \phi_0 + \frac{1}{M} (p - p_{ref}) + b \epsilon_u
\]

(5.107)

where \(M\) represents the formation compressibility, \(\epsilon_u = \text{div}(u)\) is the divergence of the solid deformation vector, \(b\) is the Biot coefficient, \(p\) is the pore pressure, and \(p_{ref}\) is a reference pore pressure.

5.12 Brinkman Momentum

The Brinkman (Brinkman-Forchheimer) momentum equation is a porous media specialization of the fluid momentum equation 5.19 given by

\[
\frac{\partial}{\partial t} \left( \frac{\rho}{\varphi} v \right) + \frac{\rho}{\varphi^2} v \cdot \nabla v - \nabla \cdot T + \left[ \frac{\rho \hat{c}}{\sqrt{k}} \|v\| + \frac{\mu}{k} \right] v - g = 0
\]

(5.108)

where \(\varphi\) is the matrix porosity, \(T\) is the fluid stress tensor \(g\) is a body force and \(\mu\) is termed the flowing liquid viscosity. The bracketed terms are often referred to as the Forchheimer or inertial flow contribution and the Darcy term respectively. In the limit of low velocities the equation reduces to a balance of pressure and a scaled velocity and Darcy’s law for a single component fluid is recovered [12]. This equation is most useful in cases where a clear fluid domain is bounded by a porous media. In most applications of this equation the hydrostatic contribution will be negligible but is retained here for completeness. We construct the G/FEM residual form of 5.108 by contracting with the unit coordinate vector in the \(k\)-direction, \(e_k\), multiplying
by the weight function $\phi^i$ and integrating over the volume. Using the vector identity $\nabla \cdot (T \cdot e_k \phi^i) = \nabla \cdot (T \cdot e_k \phi^i) - T^i : \nabla (e_k \phi^i)$ and integrating by parts gives

$$R_{m,k}^i = \int \frac{\partial}{\partial t} \left( \frac{\rho}{\varphi} v + \frac{\rho}{\varphi^2} v \cdot \nabla v + \left[ \frac{\rho_c}{\sqrt{k}} \|v\| + \frac{\mu}{k} \right] - g \right) \cdot e_k \phi^i + T^i : \nabla (e_k \phi^i) \right) \, dV - \int_S n \cdot T \cdot e_k \phi^i \, dS = 0 \quad (5.109)$$

Here we note that the viscosity associated with the stress tensor $T$ is not the viscosity of the fluid but instead a so-called Brinkman viscosity $\mu_B$. In Aria, each term in 5.109 is specified separately as identified in equation 5.15.

$$R_{m,k}^i = \int \frac{\partial}{\partial t} \left( \frac{\rho}{\varphi} v \right) \cdot e_k \phi^i \, dV + \int \frac{\rho}{\varphi^2} v \cdot \nabla v \cdot e_k \phi^i \, dV - \int \frac{\rho_c}{\sqrt{k}} \|v\| + \frac{\mu}{k} v \cdot e_k \phi^i \, dV$$

$$+ \int T^i : \nabla (e_k \phi^i) \, dV - \int_S n \cdot T \cdot e_k \phi^i \, dS = 0 \quad (5.110)$$

### 5.13 Lubrication Equation

Reynolds’ lubrication equation is a reduced-order model for fluid flow in thin, confined regions. This equation is commonly used for manufacturing applications, such as bearings, coating processes, and nanomanufacturing processes. Lubricating flows also occur in high-speed machinery, such as pumps. It is only defined for reduced-order elements, such as shells.

The lubrication equation implemented here is given by

$$- \frac{\partial h}{\partial t} + \nabla \cdot q + B_u \cdot \nabla h_u - B_l \cdot \nabla h_l = 0 \quad (5.111)$$

where $q$ is the lubrication flow rate, given by

$$q = - \frac{h^3}{k \mu} \nabla p_{lub} + \frac{h}{2} (B_u + B_l). \quad (5.112)$$

In these equations, the independent variable is the lubrication pressure, $p_{lub}$. Other variables are the lubrication region height (thickness) $h$, velocity of the upper confining surface $B_u$, velocity of the lower confining surface $B_l$, the upper wall height $h_u$, the lower wall height $h_l$, turbulent parameter $k$, and viscosity $\mu$. The lubrication height is defined as $h = h_u - h_l + n \cdot d$, where $n$ is the shell normal vector and $d$ is the mesh displacement. This equation is detailed in [13].

Using equation 5.111, the G/FEM residual form is

$$R_V^i = - \int \frac{\partial h}{\partial t} \phi^i \, dV + \int q \cdot \nabla \phi^i \, dV + \int (B_u \cdot \nabla h_u - B_l \cdot \nabla h_l) \phi^i \, dV + \int n \cdot q \phi^i \, dS = 0 \quad (5.113)$$

### 5.14 Stress Tensor Projection Equation

A projection equation is defined as an equation where a derived quantity at the interior Gauss points is evaluated and projected to be a solution unknown at the nodal points. The stress tensor projection
equation projects the momentum stresses, \(\tau\), without the pressure term, to the nodal points. Projecting the momentum stress smooths out the momentum stress tensor and allows for a dot product to be carried out on the projected field, which is needed for least squares stabilization schemes. The solution variable, \(\tau_{ab}\), is calculated from 5.114.

\[
R_V^i = -\int_V (\tau_{ab} - src_{\cdot\tau_{ab}})\phi^i \, dV = 0
\]  

(5.114)

\(\tau_{ab}\) is a tensor variable. For 2D problems, \(ab\) stands for XX, XY, YX, and YY. For 3D problems \(ab\) stands for XX, XY, XZ, YX, YY, YZ, ZX, ZY, and ZZ. The source term in the equation \(src_{\cdot\tau_{ab}}\) refers to the momentum stress without the pressure diagonal term.

\[
src_{\cdot\tau_{xx}} = 2\mu \frac{du}{dx} - \frac{2}{3}(\mu - \lambda)(\nabla \cdot \mathbf{v})
\]

\[
src_{\cdot\tau_{yy}} = 2\mu \frac{dv}{dy} - \frac{2}{3}(\mu - \lambda)(\nabla \cdot \mathbf{v})
\]

\[
src_{\cdot\tau_{zz}} = 2\mu \frac{dw}{dz} - \frac{2}{3}(\mu - \lambda)(\nabla \cdot \mathbf{v})
\]

\[
src_{\cdot\tau_{xy}} = src_{\cdot\tau_{yx}} = \mu \left( \frac{du}{dy} + \frac{dv}{dx} \right)
\]

\[
src_{\cdot\tau_{xz}} = src_{\cdot\tau_{zx}} = \mu \left( \frac{du}{dz} + \frac{dw}{dx} \right)
\]

\[
src_{\cdot\tau_{yz}} = src_{\cdot\tau_{zy}} = \mu \left( \frac{dv}{dz} + \frac{dw}{dy} \right)
\]

### 5.15 Potential Projection Equation

This equation is used to create a potential field which minimizes difference between the potential, \(\psi\), and the potential source, \(S\), in the least-squares sense

\[
(\nabla \psi - S)^2 = 0.
\]  

(5.115)

This equation was derived to create a hydrostatic potential field, so the most common usage is where \(S = \rho g\).

Equation 5.115 is integrated by parts, resulting in the residual

\[
R_V^i = \int_V \nabla \psi \cdot \nabla (e_k \phi^i) \, dV - \int_V \mathbf{S} \cdot \nabla (e_k \phi^i) \, dV
\]  

(5.116)

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5.16 Notes on Solid Mechanics

Some of the standard references on solid mechanics include [14], [15], [4] and [5]. As is often the case, the mathematical notion used throughout these texts is different in many cases and this is often a source of confusion. Here, we’ll lay out some basic definitions in our notation and, when possible, give the notation used in these other texts.

In what follows, \( \mathbf{x} \) is the position vector of a material particle in the deformed or current spatial configuration and \( \mathbf{X} \) is the position vector of a material particle in the undeformed or initial or reference configuration. The displacement vector \( \mathbf{d} \) is the difference between these to states\(^1 \) viz. \( \mathbf{x} = \mathbf{X} + \mathbf{d} \).

We will make extensive use of the gradients of these fields and so it is important to distinguish between gradients in the reference configuration and the current configuration. Gradients in the current configuration are denoted \( \nabla \) in Gibbs notation or \( \partial / \partial x_i \) in index notation. Gradients in the reference configuration are denoted \( \nabla_{\mathbf{X}} \) in Gibbs notation or \( \partial / \partial X_i \) in index notation\(^2 \).

Next, we define the deformation gradient \( \mathbf{F} \)

\[
\mathbf{F} \equiv \nabla_{\mathbf{X}} \mathbf{x}^t
\]

\[
= \nabla_{\mathbf{X}} \mathbf{X}^t + \nabla_{\mathbf{X}} \mathbf{d}^t
\]

\[
= \mathbf{I} + \nabla_{\mathbf{X}} \mathbf{d}^t
\]

where the superscript \(^t\) denotes the transpose operator\(^3 \). The inverse deformation gradient\(^4 \), \( \mathbf{F}^{-1} \), is also useful and can be computed as

\[
\mathbf{F}^{-1} \equiv \nabla_{\mathbf{X}}^t
\]

\[
= \nabla \mathbf{x}^t - \nabla \mathbf{d}^t
\]

\[
= \mathbf{I} - \nabla \mathbf{d}^t
\]

The determinants of \( \mathbf{F} \) and \( \mathbf{F}^{-1} \) are denoted \( J \) and \( J^{-1} \) respectively and are often used in transformations between different stress definitions\(^5 \).

It’s worth noting at this point that in Aria both gradient operators, \( \nabla \) and \( \nabla_{\mathbf{X}} \), are available as Expression objects as are \( \mathbf{F}, \mathbf{F}^{-1}, J \) and \( J^{-1} \).

The Green or Green-Lagrange strain tensor \( \mathbf{E} \) is defined\(^6 \) as

\[
\mathbf{E} \equiv \frac{1}{2} \left( \mathbf{F}^t \cdot \mathbf{F} - \mathbf{I} \right)
\]

\[
= \frac{1}{2} \left( \nabla_{\mathbf{X}} \mathbf{d} + \nabla_{\mathbf{X}} \mathbf{d}^t + \nabla_{\mathbf{X}} \mathbf{d} \cdot \nabla_{\mathbf{X}} \mathbf{d}^t \right)
\]

The Green strain is a strain measure in the reference configuration and is suitable for large deformations.

\(^1\) \( \mathbf{d} \) is denoted \( \mathbf{u} \) in [14], [15], [4], and [5].
\(^2\) In [5] \( \nabla_{\mathbf{X}} \) is denoted \( \nabla_0 \).
\(^3\) In [15] this is called the conjugate dyadic and is denoted with a subscript \( c \). In [14] the quantity \( \nabla_{\mathbf{X}} \mathbf{x}^t \) is denoted \( \mathbf{x} \nabla_{\mathbf{X}} \) where the arrow over the gradient operator denotes the direction of the operation.
\(^4\) In [15] \( \mathbf{F}^{-1} \) is denoted \( \mathbf{H} \).
\(^5\) Sometimes \( J \) is expressed as a ratio of the densities between the reference and current configurations, \( \rho_0 / \rho \).
\(^6\) In [15] \( \mathbf{E} \) is denoted \( \mathbf{L}_G \) and is called the Lagrangian strain.
and large rotations. The analogous Eulerian (or Almansi’s) strain tensor $E^*$ is defined\(^7\) as

\[
E^* = \frac{1}{2} \left( I - F^{-t} \cdot F^{-1} \right)
\]  
\[= \frac{1}{2} \left( \nabla d + \nabla d^t - \nabla d \cdot \nabla d^t \right). \tag{5.125}
\]

The Eulerian strain is also a suitable strain measure for large deformations and rotations but is defined in the current configuration.

The Cauchy stress, $\sigma$, is a stress measure defined in the current configuration as

\[
\sigma = \lambda E_{kk} I + 2\mu E \tag{5.127}
\]

where $E_{kk}$ denotes the trace of $E$ and $\lambda$ and $\mu$ are the Lamé coefficients. This constitutive equation may also be augmented with some initial residual stress or a thermal stress,

\[
\sigma = \lambda E_{kk} I + 2\mu E - \beta (T - T_\circ) I + \sigma_r. \tag{5.128}
\]

Here $\beta$ is related to the coefficient of thermal expansion, $T$ is the temperature, $T_\circ$ is the reference temperature of the solid and $\sigma_r$ is the residual stress.

For large deformations and large rotations Aria uses the second Piola-Kirchhoff stress which is defined in the reference configuration and is related to the Cauchy stress as

\[
S = J F^{-1} \cdot \sigma \cdot F^{-t}. \tag{5.129}
\]

The reverse transformation is readily given by

\[
\sigma = J^{-1} F \cdot S \cdot F^t. \tag{5.130}
\]

Mathematically, the Cauchy stress $\sigma$ is most conveniently expressed in terms of the Lamé coefficients $\lambda$, $\mu$ and $\beta$. In practice, however, it is more common to measure and report a different but related set of parameters: the Young’s modulus $E$, the Poisson’s ratio $\nu$ and the coefficient of thermal expansion $\alpha$. (Note, the shear modulus $G = \mu$.) The relationship between these two sets of parameters is

\[
2\mu = \frac{E}{(1 + \nu)} \tag{5.131}
\]

\[
\lambda = \frac{\nu E}{(1 + \nu) (1 - 2\nu)} = 2\mu \frac{\nu}{(1 - 2\nu)} \tag{5.132}
\]

\[
\beta = \frac{\alpha E}{(1 - 2\nu)} = \alpha (3\lambda + 2\mu) \tag{5.133}
\]

In Aria, there are separate Expression Names for the Cauchy stress and the second Piola-Kirchhoff stress expressions. In the input files, users provide their choice of constitutive relations in the material model specification, e.g.,

\begin{verbatim}
Begin Aria Material The_Material
   Density  = Constant rho = 2.33e-15
   Mesh Lambda = Constant lambda = 52810.30445
   Mesh Two Mu = Constant two_mu = 134426.2295
   Mesh Stress = Nonlinear_Elastic Reference_Frame=Moving
   Mesh Stress = Residual Sx=-11 Sy=-11
   Mesh Stress = Isothermal T=800 T_ref=450
End
\end{verbatim}

\(^7\)In [15] $E^*$ is denoted $E_A$. 

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In this Example, there will be three contributions to the stress in the mesh stress: nonlinear elasticity, a planar residual stress and an isotropic linear thermal stress. Internally, Aria contains a separate expression for transforming these Cauchy stresses into Piola-Kirchhoff stresses, viz.

\[ S = JF^{-1} \cdot \left( \sum_{i} \sigma_i \right) \cdot F^{-t}. \]  

(5.134)

Note that second Piola-Kirchhoff stresses are not specified in the input file – only Cauchy stresses are. Aria automatically creates an Expression to compute the transform in (5.134).

### 5.17 Units and Unit Conversions

Aria makes no a priori specification concerning the units of each term. However, as with all engineering codes, errors associated with unit conversions are quite easy to do. In some situations, with just one or two driving forces playing a role in a calculation, nondimensionalization of the equations can lead to a simplification of the problem statement (and to increased solution robustness due to proper scaling of the terms in the equations). However, in complicated cases with multiple competing forces and rate constants, sticking to unit systems to specify all quantities frequently leads to less errors in engineering calculations, and also leads to the ability to incorporate third party library packages for specification of source terms and transport properties which necessarily presume to employ units systems in their application programming interfaces (API). The next section discusses the SI units system, and its application for reacting flow and electromagnetic applications.

#### 5.17.1 SI Units

<table>
<thead>
<tr>
<th>Quantity</th>
<th>Name</th>
<th>Abbreviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>length</td>
<td>meter (metre)</td>
<td>m</td>
</tr>
<tr>
<td>mass</td>
<td>kilogram</td>
<td>kg</td>
</tr>
<tr>
<td>time</td>
<td>second</td>
<td>s</td>
</tr>
<tr>
<td>electric current</td>
<td>ampere</td>
<td>A</td>
</tr>
<tr>
<td>thermodynamic temperature</td>
<td>kelvin</td>
<td>K</td>
</tr>
<tr>
<td>amount of substance</td>
<td>kmole</td>
<td>kmol</td>
</tr>
<tr>
<td>luminous intensity</td>
<td>candela</td>
<td>cd</td>
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</tbody>
</table>

Table 5.1. Fundamental SI units.

discuss electromagnetic unit specification, e.g., Gauss’s law.
<table>
<thead>
<tr>
<th>Factor</th>
<th>Prefix</th>
<th>Abbreviation</th>
</tr>
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<tbody>
<tr>
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<td>Y</td>
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**Table 5.2.** SI magnitude prefixes.
Table 5.3. SI derived units and their definitions.
### Table 5.4. CGS derived units and their definitions.

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<tr>
<th>Quantity</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>erg</td>
<td>1 erg = $10^{-7}$ J</td>
</tr>
<tr>
<td>dyne</td>
<td>1 dyn = $10^{-5}$ N</td>
</tr>
<tr>
<td>poise</td>
<td>1 P = 1 dyn·s/cm² = 0.1 Pa·s</td>
</tr>
<tr>
<td>stokes</td>
<td>1 St = 1 cm²/s = $10^{-4}$ m²/s</td>
</tr>
<tr>
<td>gauss</td>
<td>1 G = $10^{-4}$ T</td>
</tr>
<tr>
<td>oersted</td>
<td>1 Oe = (1000/(4 pi)) A/m</td>
</tr>
<tr>
<td>maxwell</td>
<td>1 Mx = $10^{-8}$ Wb</td>
</tr>
<tr>
<td>stilb</td>
<td>1 sb = 1 cd/cm² = $10^4$ cd/m²</td>
</tr>
<tr>
<td>phot</td>
<td>1 ph = $10^4$ lx</td>
</tr>
</tbody>
</table>

### 5.17.2 Common Units and Conversion Factors

\[
1 \text{ g} \cdot \text{s}^2 = 1 \text{ dyn} = 10^{-5} \text{ kg} \cdot \text{m/s}^2 = 10^{-5} \text{ N}
\]
\[
1 \text{ g} \cdot \text{s}^2 = 7.2330 \times 10^{-5} \text{ lbm} \cdot \text{ft/s}^2 \text{ (poundal)}
\]
\[
1 \text{ lbf} = 4.4482 \text{ N}
\]
\[
1 \text{ g} \cdot \text{s}^2 = 2.2481 \times 10^{-6} \text{ lbf}
\]

### Table 5.5. Units and conversion factors for force.

<table>
<thead>
<tr>
<th>Pressure</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 bar</td>
</tr>
<tr>
<td>1 psia</td>
</tr>
<tr>
<td>1 psia</td>
</tr>
<tr>
<td>1 psia</td>
</tr>
<tr>
<td>1 psia</td>
</tr>
<tr>
<td>1 atm</td>
</tr>
<tr>
<td>1 atm</td>
</tr>
<tr>
<td>1 atm</td>
</tr>
<tr>
<td>1 atm</td>
</tr>
</tbody>
</table>

### Table 5.6. Units and conversion factors for pressure and stress.

<table>
<thead>
<tr>
<th>Viscosity</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 cp</td>
</tr>
<tr>
<td>1 cp</td>
</tr>
<tr>
<td>1 cp</td>
</tr>
<tr>
<td>1 cp</td>
</tr>
<tr>
<td>1 cp</td>
</tr>
<tr>
<td>1 Pa·s</td>
</tr>
</tbody>
</table>

### Table 5.7. Units and conversion factors for viscosity.
Table 5.8. Units and conversion factors for density.

| 1 g/cm³ | 1000 kg/m³ | 62.43 lb/ft³ |
| g/cm³ | 8.345 lb/U.S. gal |
| lb/ft³ | 16.0185 kg/m³ |

Table 5.9. Units and conversion factors for thermal conductivity.

| 1 btu/h·ft·F | 4.1365×10⁻³ cal/s·cm·°C |
| 1 btu/h·ft·F | 1.73073 W/m·K |

1 btu/h·ft²·F = 1.3571×10⁻⁴ cal/s·cm²·°C
1 btu/h·ft²·F = 5.6783×10⁻⁴ W/cm²·°C
1 btu/h·ft²·F = 5.6783 W/m²·°C
1 kcal/h·m²·°F = 0.2048 btu/h·ft²·°F

Table 5.10. Units and conversion factors for heat-transfer coefficients.
Chapter 6

Equation Specification

This chapter documents the EQ line commands within the current version of Aria. EQ line commands add equations and independent variables to Aria’s specification of the equation set to be solved for within each region. The equation is also associated with a field variable here that becomes part of the solution vector for Aria. EQ commands occur in Aria’s input file within Region blocks.

The EQ command line adds an equation to be solved for on a particular MESH_PART. The format is as follows

```
EQ equation FOR DOF on MESH_PART using INTERP with TERM0...TERMn
```

Equation is the string identifier for the individual equations listed in the previous chapter. The format for the equation string identifiers is listed in a subsection of Chapter 2. The DOF keyword specifies the independent unknown that is solved for in order to satisfy the equation. Normally, it is a strict function of the equation keyword. In other words, the temperature is the only valid DOF entry if the energy equation is being solved. MESH_PART is usually the name of an active element block in the finite element model. Unfortunately, if an equation is to be solved on the entire finite element model, this means that there must be multiple EQ keywords for each element block defined in the mesh.

INTERP defines the finite element interpolation to be used. Currently the valid entries for this keyword are P0, P1, Q1, Q2 and Q2S with standard volumes, e.g. hexahedron, tetrahedron, quadrilateral and triangle. These keywords imply the interpolation and polynomial order for the solution field where P denotes (polynomial) element interpolation and Q denotes the nodal interpolation (Q1 - linear, Q2 - quadratic and Q2S for near-quadratic, serendipity). We note that no syntactical distinction is made between element topology (i.e. triangle and quadrilateral or tetrahedron and hexahedron), that is, the syntax indicates only the level of interpolation. Here the association of element topology is handled internal to the code. When defining equations on shell elements the keywords above are paired (e.g. Q1P0) according to interpolation in the shell plane and interpolation through the shell thickness. For example Q1P0 represents linear nodal interpolation in the plane and constant through the thickness. In sets of coupled equations, arbitrary combinations of interpolations are sometimes not permitted for solution variables appearing in coupling terms of an equation as interpolation must be consistent in any given equation.

TERM refer to the broad categories for the terms in a general advection-diffusion continuity equation. Each term in the equation must be explicitly turned on for it to appear in the conservation equation. Admissible values of TERM are MASS, LUMPED_MASS, ADV, DIFF, SRC, and XFER. While MASS, LUMPED_MASS, ADV, DIFF, SRC terms imply the activation of mathematical operators appearing in an equation, XFER implies that a solution field values for this equation are being provided externally so the equation can still be thought of as belonging to the overall equation set.

Provisions are made for supplying multiple equation contributions for DIFF. These contributions are usually activated via settings in the Aria Materials command block 4.1. Multiple contributions can also be defined for SRC by adding additional source command lines 11. In order to accommodate different definitions of velocity for ADV the user can request the velocity to be provided from another equation, from transfer or from other available velocity models 6.27.
All equations are assumed to be formulated in Cartesian coordinates. For two-dimensional problems, axisymmetric formulations are available as well as the Cartesian forms. Invocation of the axisymmetric option is described in the Model Definition chapter of this manual 3.1 in the FINITE ELEMENT MODEL section.

6.1 EQ CONTINUITY

EQ CONTINUITY[<Subindex>][<Phase>] for DOF ON MESH_PART USING INTERP with TERM0...TERMn

Description
Activates the continuity equation 5.4.

Summary
The only admissible value for DOF is PRESSURE.
Admissible values of INTERP are Q1, Q2, Q2S, P1, P0, for standard volume elements and Q1P0, Q2P0, Q2SP0, Q1Q1 and Q1Q2 for shell elements.
Admissible values of TERMn are MASS, LUMPED_MASS, DIV, ADV, SRC, and XFER.
With the exception of XFER these terms are described in the Equations Aria Solves section 5 of the manual.
The MESH_PART must be an active element block.

Parent Block(s) ARIA_REGION

6.2 EQ CURRENT

EQ CURRENT[<Subindex>][<Phase>] for DOF ON MESH_PART USING INTERP with TERM0...TERMn

Description
Activates the electrical current equation 5.33.

Summary
The only admissible value for DOF is VOLTAGE.
Admissible values of INTERP are Q1, Q2, Q2S.
Admissible values of TERMn are DIFF, SRC, and XFER.
With the exception of XFER these terms are described in the Equations Aria Solves section 5 of the manual.
The MESH_PART must be an active element block.

Parent Block(s) ARIA_REGION

6.3 EQ CHARGE_DENSITY

EQ CHARGE_DENSITY[<Subindex>][<Phase>] for DOF ON MESH_PART USING INTERP with TERM0...TERMn
Description  Activates the charge density equation 5.36.

Summary  The only admissible value for DOF is CHARGE_DENSITY. 
Admissible values of INTERP are $Q_1, Q_2, Q_{2S}$. 
Admissible values of TERM are MASS, LUMPED_MASS, ADV, DIFF, and XFER. 
With the exception of XFER these terms are described in the Equations Aria Solves section 5 of the manual. 
The MESH_PART must be an active element block.

Parent Block(s)  ARIA_REGION

6.4 EQ ENERGY

EQ_ENERGY[<Subindex>][<Phase>] for DOF ON MESH_PART USING INTERP with TERM$_0$...TERM$_n$

Description  Activates the energy conservation transport equation 5.9.

Summary  Admissible values for DOF are TEMPERATURE and ENTHALPY. 
Admissible values of INTERP are $Q_1, Q_2, Q_{2S}$, for standard volume elements and $Q_1 P_0, Q_2 P_0, Q_1 Q_1$ and $Q_1 Q_2$ for shell elements. 
Admissible values of TERM$_n$ are MASS, LUMPED_MASS, ADV, DIFF, SRC, and XFER. 
With the exception of XFER these terms are described in the Equations Aria Solves section[5.3] of the manual. 
The MESH_PART must be an active element block.

Parent Block(s)  ARIA_REGION

6.5 EQ LEVEL_SET

EQ_LEVEL_SET[<Subindex>] for DOF ON MESH_PART USING INTERP with TERM$_0$...TERM$_n$

Description  Activates the level set (distance function) equation. This equation is often accompanied by additional commands described in 29.

Summary  The only admissible value for DOF is LEVEL_SET. 
Admissible values of INTERP are $Q_1, Q_2, Q_{2S}$. 
Admissible values of TERM$_n$ are MASS, LUMPED_MASS, ADV, DIFF, SRC, and XFER. 
With the exception of XFER these terms are described in the Equations Aria Solves section 5 of the manual. 
The MESH_PART must be an active element block.

Parent Block(s)  ARIA_REGION
6.6  EQ EXTENSION_SPEED

EQ EXTENSION_SPEED[<_Subindex>] for DOF ON MESH_PART USING INTERP with TERM<sub>0</sub>...<sub>n</sub>

Description  Activates an equation for extension speed, which is used by the level set equation (29) to define the interface speed.

Summary  The only admissible value for DOF is EXTENSION_SPEED. Admissible values of INTERP are \( Q_1, Q_2, Q_{2S} \). Admissible values of TERM<sub>n</sub> are DEF, SRC, and XFER. With the exception of XFER these terms are described in the Equations Aria Solves section 5 of the manual. The MESH_PART must be an active element block.

Parent Block(s)  ARIA_REGION

6.7  EQ MASS_BALANCE

EQ MASS_BALANCE[<_Subindex>] for DOF ON MESH_PART USING INTERP with TERM<sub>0</sub>...<sub>n</sub>

Description  The mass balance equation for porous media. See general porous flow information in section 5.11

Summary  Admissible value for DOF are PRESSURE, DENSITY, and MASS_FRACTION. Admissible values of INTERP are \( Q_1, Q_2, Q_{2S} \). Admissible values of TERM<sub>n</sub> are MASS, LUMPED_MASS, ADV, UPWIND_ADV, DIFF, SRC, and XFER. With the exception of XFER these terms are described in the Equations Aria Solves section 5 of the manual. The MESH_PART must be an active element block.

Parent Block(s)  ARIA_REGION

6.8  EQ MESH

EQ MESH[<_Subindex>] for DOF ON MESH_PART USING INTERP with TERM<sub>0</sub>...<sub>n</sub>

Description  Activates the pseudo-solid mesh equation 5.22.
Summary
The only admissible value for DOF is MESH_DISPLACEMENTS.
Admissible values of INTERP are $Q_1$, $Q_2$, $Q_{2S}$.
Admissible values of TERM$_n$ are DIFF, REF_DIFF, TTRAIN, SRC, and XFER.
With the exception of XFER these terms are described in the Equations Aria Solves section 5 of the manual.
If both the MESH and SOLID equations are defined in the same region, then the TALE mesh motion algorithm is activated automatically. In this case, the MESH equation must be defined on all blocks in the region. More details of the TALE method can be found in section 5.6.
The MESH_PART must be an active element block.

Parent Block(s) ARIA_REGION

6.9 EQ MOMENTUM

EQ MOMENTUM[_<Subindex>][_<Phase>] for DOF ON MESH_PART USING INTERP with TERM$_0$...TERM$_n$

Description Activates the fluid momentum equation 5.19.

Summary The only admissible value for DOF is VELOCITY.
Admissible values of INTERP are $Q_1$, $Q_2$, $Q_{2S}$.
Admissible values of TERM$_n$ are MASS, LUMPED_MASS, ADV, DIFF, SRC, and XFER.
With the exception of XFER these terms are described in the Equations Aria Solves section 5 of the manual.
The MESH_PART must be an active element block.

Parent Block(s) ARIA_REGION

6.10 EQ POROUS_SPECIES

EQ POROUS_SPECIES[_<Subindex>] for DOF ON MESH_PART USING INTERP with TERM$_0$...TERM$_n$

Description The species transport equation porous media. See general porous flow information in section 5.11

Summary Admissible value for DOF are PRESSURE, SPECIES, and SPECIES_FRACTION.
Admissible values of INTERP are $Q_1$, $Q_2$, $Q_{2S}$.
Admissible values of TERM$_n$ are MASS, LUMPED_MASS, ADV, UPWIND_ADV, DIFF, SRC, and XFER.
With the exception of XFER these terms are described in the Equations Aria Solves section 5 of the manual.

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The *MESH_PART* must be an active element block.
Additionally, one can choose whether to integrate the advection term by parts by setting the flag *INTEGRATE ADVECTION BY PARTS = false* in the *POROUS_FLOW_OPTIONS* block of *SOLUTION_OPTIONS*. The default is true.

**Parent Block(s)** ARIA_REGION

### 6.11 EQ POROUS_ENTHALPY

**EQ POROUS_ENTHALPY[_,<Subindex>] for DOF ON MESH_PART USING INTERP with TERM₀...TERMₙ**

**Description** The enthalpy transport equation porous media. See general porous flow information in section 5.11

**Summary** The only admissible value for DOF is TEMPERATURE.
Admissible values of *INTERP* are *Q₁, Q₂, Q₂S*.
Admissible values of *TERMₙ* are *MASS, LUMPED_MASS, ADV, UPWIND_ADV, DIFF, SRC, and XFER*.
With the exception of XFER these terms are described in the Equations Aria Solves section 5 of the manual.
The *MESH_PART* must be an active element block.
Additionally, one can choose whether to integrate the advection term by parts by setting the flag *INTEGRATE ADVECTION BY PARTS = false* in the *POROUS_FLOW_OPTIONS* block of *SOLUTION_OPTIONS*. The default is true.

**Parent Block(s)** ARIA_REGION

### 6.12 EQ POTENTIAL

**EQ POTENTIAL[_,<Subindex>][_,<Phase>] for DOF ON MESH_PART USING INTERP with TERM₀...TERMₙ**

**Description** Activates the potential projection equation for the potential, which is defined in equation 5.115.

**Summary** The only admissible value for DOF is POTENTIAL.
Admissible values of *INTERP* are *Q₁, Q₂, Q₂S*.
Admissible values of *TERMₙ* are *DEF and SRC, and both terms must be active for this equation to behave properly. The value used in SRC must be a separately defined source equation.*
The *MESH_PART* must be an active element block.

**Parent Block(s)** ARIA_REGION
6.13 EQ SHEAR

EQ SHEAR[<Subindex>][<Phase>] for DOF ON MESH_PART USING INTERP with TERM₀...TERMₙ

Description  Activates the shear-rate definition/intermediate equation.

Summary  The only admissible value for DOF is GAMMA_DOT.
Admissible values of INTERP are Q₁, Q₂, Q₂S.
Admissible values of TERMₙ are DEF, SRC, and XFER.
With the exception of XFER these terms are described in the Equations Aria Solves section 5 of the manual.
The MESH_PART must be an active element block.

Parent Block(s)  ARIA_REGION

6.14 EQ SOLID

EQ SOLID[<Subindex>][<Phase>] for DOF ON MESH_PART USING INTERP with TERM₀...TERMₙ

Description  Activates the solid momentum equation 5.22.

Summary  The only admissible value for DOF is SOLID_DISPLACEMENTS.
Admissible values of INTERP are Q₁, Q₂, Q₂S.
Admissible values of TERMₙ are DIFF, TSTRAIN, SRC, and XFER.
With the exception of XFER these terms are described in the Equations Aria Solves section 5 of the manual.
If both the MESH and SOLID equations are defined in the same region, then the TALE mesh motion algorithm is activated automatically. In this case, the reference configuration for the SOLID equation is converted to the TALE reference frame. More details of the TALE method can be found in section 5.6.
The MESH_PART must be an active element block.

Parent Block(s)  ARIA_REGION

6.15 EQ SPECIES

EQ SPECIES[<Subindex>][<Phase>] for DOF ON MESH_PART USING INTERP with TERM₀...TERMₙ

Description  Activates the species transport equation 5.13.
Summary

The only admissible value for **DOF** is **SPECIES**.

Admissible values of **INTERP** are **$Q_1$**, **$Q_2$**, **$Q_{2S}$**.

Admissible values of **TERM** are **MASS**, **LUMPED_MASS**, **ADV**, **DIFF**, **SRC**, **FRACBAL** and **XFER**. With the exception of **XFER** these terms are described in the Equations Aria Solves section 5 of the manual. The FRACBAL term may not be included with other terms.

Some other things to note about species equations in Aria.

- The FRACBAL term may be assigned to any species number.
- Species numbers in Aria are arbitrary; they may start at any value and need not be continuous.
- For the species fraction balances, Aria will automatically detect all species that are present in the problem and include them in the balance.

The **MESH_PART** must be an active element block.

Parent Block(s) **ARIA_REGION**

### 6.16 EQ SP

**EQ SP ORDER** for **DOF** ON **MESH_PART** USING **INTERP** with **TERM**

**Description**

Specifies the Spherical Harmonics equation set to solve for Participating Media Radiation (PMR). User provides the order, degree of freedom associated with the equation, the element type and the mesh part (element block) on which to apply the equation 23.2.

**Summary**

Admissible values for **ORDER** are odd values (1, 3, 5, ...).

The only admissible value for **DOF** is **angular_intensity**.

The **MESH_PART** must be an active element block.

Admissible values of **INTERP** are **$Q_1$**, **$Q_2$**, **$Q_{2S}$**

Admissible value of **TERM** is **SRC**.

Parent Block(s) **ARIA_REGION**

### 6.17 EQ SUSPENSION

**EQ SUSPENSION**[[_<Subindex>][<_Phase>_]] for **DOF** ON **MESH_PART** USING **INTERP** with **TERM**

**Description**

Activates the suspension transport equation 5.39.

**Summary**

The only admissible value for **DOF** is **PHI**.

Admissible values of **INTERP** are **$Q_1$**, **$Q_2$**, **$Q_{2S}$**.
Admissible values of $\text{TERM}_n$ are ADV, DIFF, SRC, and XFER.

With the exception of XFER these terms are described in the Equations Aria Solves section 5 of the manual.

The $\text{MESH\_PART}$ must be an active element block.

Parent Block(s) ARIA\_REGION

### 6.18 EQ VOLTAGE

$\text{EQ \ VOLTAGE[<_\text{Subindex}>][<_\text{Phase}>]\ for DOF ON MESH\_PART USING \text{INTERP with \text{TERM}_0...\text{TERM}_n}}$

<table>
<thead>
<tr>
<th>Description</th>
<th>Activates the voltage equation (electric-displacement formulation) 5.30. See also the CURRENT equation.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Summary</td>
<td>The only admissible value for DOF is VOLTAGE. Admissible values of $\text{INTERP}$ are $Q_1$, $Q_2$, $Q_{2S}$. Admissible values of $\text{TERM}_n$ are DIFF, SRC, and XFER. With the exception of XFER these terms are described in the Equations Aria Solves section 5 of the manual. The $\text{MESH_PART}$ must be an active element block.</td>
</tr>
</tbody>
</table>

Parent Block(s) ARIA\_REGION

### 6.19 EQ BRINKMAN\_MOMENTUM

$\text{EQ \ BRINKMAN\_MOMENTUM[<_\text{Subindex}>][<_\text{Phase}>]\ for DOF ON MESH\_PART USING \text{INTERP with \text{TERM}_0...\text{TERM}_n}}$

<table>
<thead>
<tr>
<th>Description</th>
<th>Activates the Brinkman momentum equation 5.108.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Summary</td>
<td>The only admissible value for DOF is VELOCITY. Admissible values of $\text{INTERP}$ are $Q_1$, $Q_2$, $Q_{2S}$. Admissible values of $\text{TERM}_n$ are MASS, LUMPED_MASS, ADV, DIFF, SRC, and XFER. With the exception of XFER these terms are described in the Equations Aria Solves section 5 of the manual. The $\text{MESH_PART}$ must be an active element block.</td>
</tr>
</tbody>
</table>

Parent Block(s) ARIA\_REGION
6.20 EQ LUBRICATION

EQ LUBRICATION[<Subindex>][<Phase>] for DOF ON MESH_PART USING INTERP with TERM₀...TERMₙ

Description  Activates the Reynolds’ lubrication equation 5.111.

Summary  The only admissible value for DOF is LUBRICATIONPRESSION.
Admissible values of INTERP is Q₁P₀.
Admissible values of TERMₙ are MASS, DIFF, and BOUND.
For problems with fluid-structural interactions, the lubrication equation needs to be defined on all blocks, including 3-D blocks adjoining the shell lubrication region. In order to accommodate this, there is an additional term, DEF, which applies the equation p = 0 on adjoining 3-D blocks where this is used. The other terms should not be used on 3-D blocks. It is weighted so that it does not affect the residual on the shell blocks.
The MESH_PART must be an active shell element block.

Parent Block(s)  ARIA_REGION

6.21 EQ Stress_Tensor_Projection

EQ Stress_Tensor_Projection[<Subindex>][<Phase>] for DOF ON MESH_PART USING INTERP with TERM₀...TERMₙ

Description  Activates the Stress Tensor Projection equation.

Summary  The only admissible value for DOF is Stress_Tensor.
Admissible values of INTERP are Q₁, Q₂, Q₂S.
The only admissible values of TERMₙ are DEF.
With the exception of XFER these terms are described in the Equations Aria Solves section 5 of the manual.
The MESH_PART must be an active element block.

Parent Block(s)  ARIA_REGION

6.22 ELASTICITY FORMULATION

ELASTICITY FORMULATION = PLANE TYPE

Description  Assigns the elasticity formulation TYPE for two-dimensional problems involving the MESH and SOLID equation 5.22.
Summary
Allowable formulation PLANE TYPE is PLANE STRESS or PLANE STRAIN.

Parent Block(s) ARIA_REGION

6.23 PRESSURE STABILIZATION

PRESSURE STABILIZATION IS TYPE WITH SCALING = C

Description
Prescribe a stabilization technique for solving the MOMENTUM 5.19 and CONTINUITY 5.4 equations with equal order interpolation.

Summary
Aria supports both PSPG (Pressure Stabilized Petrov-Galerkin) and PSPP (Pressure Stabilized Pressure Projection) stabilization techniques for solving the MOMENTUM and CONTINUITY equations with equal order interpolation.

Valid options for the TYPE specification are NO_STABILIZATION, PSPG_CONSTANT, PSPG_LOCAL, PSPG_GLOBAL and PSPP_CONSTANT.

NO_STABILIZATION disables any stabilization.

PSPP_CONSTANT results in the recently developed stabilization technique of [16] and [17].

In the PSPG forms of stabilization, introduced by [18], terms from the momentum equation are added to the continuity equation scaled by a multiplier, $\alpha$. The exact form of the multiplier depend on a global Reynolds number that is defined as

$$Re \equiv \frac{\rho |v| \langle h \rangle}{2\mu}$$

(6.1)

Here, $\rho$ is the density, $\mu$ is the viscosity, $|v|$ is a velocity scale and $\langle h \rangle$ is an element length scale. Armed with $Re$, the stabilization multiplier $\alpha$ is defined in one of two ways.

$$Re \leq 3 : \quad \alpha = \frac{\tau \langle h \rangle^2}{12\mu}$$

(6.2)

$$Re > 3 : \quad \alpha = \frac{\tau \langle h \rangle}{2\rho |v|}$$

(6.3)

NB: Currently, Aria always uses the low-Reynolds number for of $\alpha$, as defined in (6.2). The PSPG_LOCAL method computes $|v|$ and $\langle h \rangle$ within each element. The PSPG_GLOBAL method computes $|v|$ and $\langle h \rangle$ as averages over all of the elements with the MOMENTUM equation defined. The PSPG_CONSTANT gives $|v|$ and $\langle h \rangle$ a value of 1 (one) and just uses the scale factor.

Parent Block(s) ARIA_REGION

6.24 SAVE RESIDUALS

SAVE RESIDUALS = NODE

Description
Causes Aria to save the residuals to a field with the prefix residual->, e.g., residual->Temperature. This will be done for all fields (though we could make it a per-field option). Valid choices are OFF (default), BEFORE_BCS and AFTER_BCS.
Summary

Causes Aria to save the residuals to a field with the prefix residual->, e.g., residual->Temperature. This will be done for all fields (though we could make it a per-field option). Valid choices are OFF (default), BEFORE_BCS and AFTER_BCS.

For the choice BEFORE_BCS the residuals will be saved at the point in the assembly process where the primary equations have been assembled but prior to the assembly of any boundary conditions or distinguishing conditions. For the choice of AFTER_BCS the residuals will be saved after all BCs and distinguishing conditions have been applied. The default, OFF, is to not save the residuals.

This feature is only applicable when using the NEWTON nonlinear solution strategy.

Parent Block(s) ARIA_REGION

6.25 INTEGRATION RULE

INTEGRATION RULE for block_name = INT

Description

Overrides the default integration rule for the equations defined on block_name.

Summary

Overrides the default integration rule for the equations defined on block_name.

Parent Block(s) ARIA_REGION

6.26 MESH GROUP

MESH_GROUP alias_name = mesh_entity1 . . mesh_entityN
MESH_GROUP alias_name = group1 + mesh_entity
MESH_GROUP alias_name = group1 - group2
MESH_GROUP alias_name = block_1 to block_N

Description

Assigns aliased grouping of selected mesh entities (block, surface or nodelist). These aliases can then be used in defining equations, source terms, initial conditions, Dirichlet boundary conditions, surface flux conditions, postprocessing operations and the INTEGRATION RULE command.

In order to simplify the specification of mesh groups for large models, sum, difference and range operators are provided.

Summary

Only mesh entities can be aliased within a group and a mesh entity can appear in several groups. Mesh entities aliased within the FINITE_ELEMENT_MODEL command block can appear within MESH_GROUP and are treated in the same way as the mesh entity itself. Additionally mesh entities with names supplied from the input file can also be referenced.

By default the aliases all_blocks and all_volumes reference all active element blocks in the mesh and the all_surfaces alias references all active surfaces in the mesh. Default aliases are also defined for element blocks by topology and number of nodes, e.g. Hexahedron_8_blocks.

The first mesh_entity can be an alias to an existing MESH_GROUP provided subsequent entries define addition or subtraction to the group using (+, plus, and, -, minus, less) to define the
operation used to compose an alias. For example:

\[ \text{MESH GROUP alias\_name = all\_blocks - block\_0} \]

An additional form of the command defines a range of element blocks. In this case the group range is defined using (to, thru or through) to specify the first and last element blocks in the range based upon block number. Note that Mesh Group commands are parsed sequentially, thus command line ordering must be honored when using one Mesh Group to define another Mesh Group.

Parent Block(s)  ARIA\_REGION

### 6.27 ADVECTION VELOCITY

\[ \text{ADVECTION VELOCITY for block\_name = STRING} \]

<table>
<thead>
<tr>
<th>Description</th>
<th>Defines the velocity model to be used for the equations defined on block_name.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Summary</td>
<td>Velocity model can be XFER or any other vector velocity model defined in the Region.</td>
</tr>
</tbody>
</table>

Parent Block(s)  ARIA\_REGION
Chapter 7

Data for Material, BC, IC, SRC

In many numerical simulations the need often arises to define quantities that involve the same numerical values. Aria provides a means by which a user can collectively define problem dependent real and integer data as named members of a Data Block. These named data members can be defined either as independent values or as arrays. The utility of Data Block information in setting up problem descriptions is that rather than referencing individual data values, the data collection can be referenced as a whole using the name of the parent data block. This provides a convenient means of re-using data when defining material behavior, boundary conditions and initial conditions for a particular class of problem.

Because Data Block values are named then within the application code individual members of the Data Block can be easily referenced by name. This named feature is particularly useful in both Calore style user subroutines and in user plugin utilities. The usage of Data Blocks with Calore style user subroutines is detailed in the Chapter on Thermal Analysis 33.18.

The scope of Data Block information lies within an Aria Region. Each Data Block must have a unique name and will be referenced by that name within the problem setup. Reference to individual Data Block members will occur only within the application code.

7.1 Data Block

**Scope:** Aria Region

```
Begin Data DataBlockName
    Real {=|parameter_name|=} value...
    Integer {=|parameter_name|=} value...
End
```

**Summary**

Parameters for Data Blocks within Aria. The DataBlockName must be referenced within the definition of an initial condition, volumetric source, boundary condition or material property Calore_User_Sub in order to be available for use within a user subroutine.

The DataBlockName may also be used to define parameters for all equation line commands by specifying USE DATA BLOCK DataBlockName as the first parameter argument on the equation. When used in this manner, the typed name-value pairs of the Data Block are assigned as the parameters for the equation.

7.1.1 Real

**Scope:** Data Block
<table>
<thead>
<tr>
<th>Parameter values</th>
<th>Value real...</th>
<th>Default</th>
</tr>
</thead>
</table>

Summary Specifies the Real data values for this data block.

### 7.1.2 Integer

**Scope:** Data Block

<table>
<thead>
<tr>
<th>Parameter values</th>
<th>Value integer...</th>
<th>Default</th>
</tr>
</thead>
</table>

Summary Specifies the Integer data values for this data block.
This chapter documents the initial condition, IC, line commands within the current version of Aria. The IC simply represents an initialization the Field variable. Technically speaking initial conditions have meaning only in time-dependent partial differential equations. However, within the Aria code the IC command line results in an initialization of Field values used in the assembly of a Jacobian matrix. Thus the IC is equally applicable for assigning starting values in a steady solution. Note that in the case of steady solutions the Aria output will contain two "time" steps, an initialization output plane, and the steady solution output. A distinction between the meaning of these two steps is particularly important when using a steady solution result to provide the initial condition from file for a transient problem.

Within the input file initial conditions command lines will appear at the Region scope as illustrated below. Note that input specification of initial conditions on portions of the FEM mesh can be simplified by aggregation of element blocks into Mesh Groups 6.26.

```
Begin Procedure My_Aria_Procedure
  Begin Aria Region My_Region
    IC related commands
  End
End
```

8.1 IC CIRC_X

```
IC CIRC_X AT MESH_PART DOF AMP = Ω
```

**Description**
Initial boundary condition for the x component of a vector variable with constant tangential magnitude along circles of radius $r(x, y)$ defined on the mesh entity.

**Summary**
Sets $DOF$ to the provided value on nodeset associated with the name $MESH\_PART$ according to
the relation

\[ DOF = -\Omega r(x, y) \sin \theta. \] (8.1)

The rotation \( \Omega \) is assumed to be about the vector \((0, 0, 1)\) passing through point \((0, 0, 0)\).

Parent Block(s) ARIA_REGION

### 8.2 IC CIRC\_Y

**IC CIRC\_Y AT MESH\_PART DOF AMP = \( \Omega \)**

**Description** Initial boundary condition for the \( y \) component of a vector variable with constant tangential magnitude along circles of radius \( r(x, y) \) defined on the mesh entity.

**Summary** Sets \( DOF \) to the provided value on nodeset associated with the name \( MESH\_PART \) according to the relation

\[ DOF = \Omega r(x, y) \cos \theta. \] (8.2)

The rotation \( \Omega \) is assumed to be about the vector \((0, 0, 1)\) passing through point \((0, 0, 0)\).

Parent Block(s) ARIA_REGION

### 8.3 IC CONSTANT

**IC CONST AT MESH\_PART DOF = REAL**

**Description** Constant initial condition.

**Summary** Sets \( DOF \) to the provided constant value on mesh entity associated with the name \( MESH\_PART \).

Parent Block(s) ARIA_REGION

### 8.4 IC COUETTE\_X

**IC COUETTE\_X AT MESH\_PART DOF PARAMS = \( r_i \) \( r_o \) \( \Omega \)**

**Description** Initial condition for \( x \) component of a vector variable in a Couette device with inner radius \( r_i \), outer radius \( r_o \) and driven at angular velocity \( \Omega \), that varies spatially over a mesh entity.

**Summary** Sets \( DOF \) to vary spatially over the nodeset associated with the name \( MESH\_PART \) according to the relation

\[ DOF = -\Omega r(x, y) \frac{r_i r_o}{r_o^2 - r_i^2} \left( \frac{r_o^2}{r} - r \right) \sin \theta \] (8.3)

The rotation \( \Omega \) is assumed to be about the vector \((0, 0, 1)\) passing through point \((0, 0, 0)\).
8.5 IC COUETTE_Y

IC COUETTE_Y AT MESH_PART DOF PARAMS = $r_i$ $r_o$ $\Omega$

Description
Initial condition for x component of a vector variable in a Couette device with inner radius $r_i$, outer radius $r_o$ and driven at angular velocity $\Omega$, that varies spatially over a mesh entity.

Summary
Sets DOF to vary spatially over the nodeset associated with the name MESH_PART according to the relation

$$DOF = \Omega r(x, y) \frac{r_i r_o}{r_o^2 - r_i^2} \left( \frac{r_o^2}{r} - \frac{1}{r(x, y)} \right) \cos \theta$$

The rotation $\Omega$ is assumed to be about the vector (0, 0, 1) passing through point (0, 0, 0).

8.6 IC COUETTE_SH

IC COUETTE_SH AT MESH_PART DOF PARAMS = $r_i$ $r_o$ $\Omega$

Description
Initial condition for generalized shear rate in a Couette device with inner radius $r_i$, outer radius $r_o$ and driven at angular velocity $\Omega$, that varies spatially over a mesh entity.

Summary
Sets DOF to the provided value on nodeset associated with the name MESH_PART according to the relation

$$DOF = \frac{\Omega}{r^2(x, y)} \frac{r_i r_o}{r_o^2 - r_i^2} \left( \frac{1}{r(x, y)} - \frac{r(x, y)}{r_o^2} \right)$$

The rotation $\Omega$ is assumed to be about the vector (0, 0, 1) passing through point (0, 0, 0).

8.7 IC READ_FILE

IC READ_FILE DOF = STRING AT TIME REAL

Description
This IC command will initialize the field DOF with values from the field with the name given by the STRING argument in the input mesh database.
This IC command will initialize the field \texttt{DOF} in the current analysis with values from the input mesh database containing variable \texttt{STRING}. For example, if the mesh database as specified in the \texttt{FINITE ELEMENT MODEL} command block (3.1) contains a vector field named \texttt{U} then one could use the file to initialize a velocity field by setting \texttt{DOF} to \texttt{VELOCITY} and the \texttt{STRING} argument to \texttt{U}.

In cases where the field \texttt{DOF} has a several components (i.e. \texttt{DOF} is a vector field) and the input mesh database contains the vector components defined as scalar fields, the command must be called once for each component. Here the \texttt{DOF} is specified as \texttt{DOF\_component}.

A typical usage example would be initialize a transient calculation from a previously performed steady-state calculation in the following way. The steady-state calculation would be performed and its output variables saved to some output file. In most cases one might employ a copy of the output file to be used as a mesh file for the transient calculation. Finally, one would use the file IC command line to tie the variable named "FileVariableName" in the mesh file to the variable named \texttt{DOF} in the current simulation.

When the mesh database file contains multiple timesteps and the \texttt{TIME} option is not supplied, the default behavior will be to extract the requested field from the last timeplane in the file. The \texttt{TIME} option must be used if the initial condition is to be extracted from any specific database timeplane other than the last timeplane. In using the \texttt{TIME} option the database time step nearest to the specified time value will be used for the assignment. For an initial condition taken from a steady-state results file, the optional \texttt{TIME} parameter should be omitted.

Steady solution results are sometimes written to file in two steps. When this command is used to initialize values from a steady solution one must select the solution file step according to the content of the steady solution file results. If the file contains two solution steps then one should employ the time option and select the time corresponding to the second step.

CAVEAT: Current framework services do not allow restricting this initialization by mesh subsets. e.g. If one wishes to initialize the temperature, then all nodes in the mesh will be initialized.

NOTE: Those wishing to perform initializations as part of a formal restart are referred to \texttt{RESTART DATA} services (25.1).

\textbf{Parent Block(s)} ARIA\_REGION

\section*{8.8 IC LINEAR}

\begin{verbatim}
IC LINEAR AT MESH_PART DOF COEF = C_0 C_1 C_2 C_3
\end{verbatim}

\textbf{Description} Initial condition that varies spatially over a mesh entity in a linear fashion.

\textbf{Summary} Sets \texttt{DOF} to the provided value on nodeset associated with the name \texttt{MESH\_PART} in the following manner

\[ \texttt{DOF} = C_0 + C_1 x + C_2 y + C_3 z. \]  

\textbf{Parent Block(s)} ARIA\_REGION
8.9 IC PARAB

\texttt{IC\_PARAB\_AT \textit{MESH\_PART} DOF COEF = C_0 C_1 C_2 C_3 C_4 C_5 C_6 C_7 C_8 C_9}

\begin{description}
\item[Description] Initial condition that varies spatially over a mesh entity in a parabolic fashion.
\item[Summary] Sets \textit{DOF} to the provided value on nodeset associated with the name \textit{MESH\_PART} in the following manner
\[ \textit{DOF} = C_0 + C_1 x + C_2 y + C_3 z + C_4 x y + C_5 x z + C_6 y z + C_7 x^2 + C_8 y^2 + C_9 z^2. \] (8.7)
\end{description}

Parent Block(s) ARIA\_REGION

8.10 ANNEAL MESH ON STARTUP

\texttt{ANNEAL\_MESH\_ON\_STARTUP}

\begin{description}
\item[Description] With the command in the Region block Aria will search for a displacements field and, if found, will apply the displacements to the model and physical coordinates.
\item[Summary] With the command in the Region block Aria will search for a displacements field and, if found, will apply the displacements to the model and physical coordinates.
\end{description}

Parent Block(s) ARIA\_REGION
Chapter 9

Boundary Conditions

Boundary conditions are needed in order to define a well-posed boundary value problem as described by the FEM equations. These boundary conditions are often denoted as Dirichlet, Neumann or Robin conditions. Dirichlet or essential boundary conditions for a solution field stem directly from the governing equations for a particular physics. In Aria, Neumann and Robin conditions are generally categorized as flux boundary conditions.

This section documents primarily the native Aria boundary condition, BC, line commands within the current version of the code. Recall from a previous chapter outlining the general FEM that the method gives rise to boundary terms 5.3, i.e. various flux boundary conditions. In what follows, Dirichlet boundary conditions are first described 9.1 followed by flux boundary condition descriptions 9.2.

Boundary conditions are defined at the Region scope of the input file as illustrated below.

```
Begin Sierra myJob
  ...
  Begin Procedure My_Aria_Procedure
    ...
    Begin Aria Region My_Region
      ...
      BC related commands
      ...
    End
  End
End Sierra myJob
```

9.1 BC Dirichlet

Dirichlet or essential boundary conditions for a solution field stem directly from the governing equations for a particular physics. Here the boundary conditions are strongly enforced by substituting the user prescribed evaluation of solution unknown into the system residual and the corresponding solution increment equal to zero. In Aria default enforcement of the Dirichlet boundary condition involves a matrix row modification. Optional methods of Dirichlet BC enforcement can be specified through the linear solver interface tt BC Enforcement command line.
The boundary condition specification corresponds to the solution variable or degree-of-freedom (DOF) for a given equation as it appears in the EQ command line. As an example in the EQ command line:

\[ \text{EQ Energy for Temperature on block_2 using Q1 with Diff mass} \]

Temperature is the DOF that used in specification of the Dirichlet boundary condition whereas Energy is the equation to which the condition is applied.

For thermal analysis another interface for Dirichlet BC specification is described in the Thermal Analysis chapter 33.1. The advantage of this approach is that the syntax is fully supported within the Sandia Analysis Workbench (SAW).

Dirichlet boundary conditions are applied at a surface of the physical problem domain. Within the context of Sierra Mechanics that surface corresponds to a surface (e.g. surface_3) or a nodelist (e.g. nodelist_2). In what follows the Constant, Encore Function, Global, User Function, Polynomial, User Field and Exponential conditions must be applied to a surface. The remainder of the Dirichlet BCs listed can be applied to either a surface or a nodelist.

Note that stylistically the syntax for Constant, Encore Function, Global, User Function, Polynomial, User Field and Exponential conditions differ from the remaining Dirichlet BC command lines. As an example:

\[ \text{BC Dirichlet BC for Energy on surface_3 = constant value = 273.0} \]

implicitly refers to a Temperature boundary condition. This is in contrast to the remaining Dirichlet BC command lines:

\[ \text{BC Const Dirichlet at surface_3 temperature = 273.0} \]

where the DOF is explicitly stated.

### 9.1.1 Generic: Constant

**Scope:** Equation System

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dof Name</td>
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</tr>
<tr>
<td>Species</td>
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<td>value</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** Constant value

### 9.1.2 Generic: Encore Function

**Scope:** Equation System

<table>
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<th>Parameter</th>
<th>Value</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Dof Name</td>
<td>{of</td>
<td>species</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
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<tr>
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<td>Eval_Type</td>
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</tr>
</tbody>
</table>
### 9.1.3 Generic: Global

**Scope:** Equation System

**Generic: Global**

For Dof Name [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Global [ Using Data Specification Data Spec Name ][ Global_Name = global_name ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dof Name</td>
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</table>

**Summary**

Value from an Encore function

### 9.1.4 Generic: User Function

**Scope:** Equation System

**Generic: User Function**

For Dof Name [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = User_Function [ Using Data Specification Data Spec Name ][ Name = name |X = x |X.Multiplier = x_multiplier |Multiplier = multiplier |Toggle = toggle ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dof Name</td>
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</tr>
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</table>

**Summary**

Value from a global variable
### 9.1.5 Generic: Polynomial

**Scope:** Equation System

<table>
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<th>Value</th>
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</thead>
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### 9.1.6 Generic: User Field

**Scope:** Equation System

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</tbody>
</table>
9.1.7 Generic: Exponential

Scope: Equation System

Generic: Exponential
For Dof Name [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Exponential [ Using Data Specification Data Spec Name ][ Variable = variable | Constant = constant | Multiplier = multiplier | Exponent = exponent ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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Summary Value from an exponential function

9.1.8 BC Dirichlet = BC CIRC_X

BC CIRC_X DIRICHLET AT MESH_PART DOF AMP = Ω

Description Dirichlet boundary condition for the x component of a vector variable with constant tangential magnitude along circles of radius \( r(x, y) \) defined on the mesh entity.

Summary Sets DOF to the provided value on nodeset associated with the name MESH_PART according to the relation

\[
DOF = -\Omega r(x, y) \sin \theta.
\]  

The rotation \( \Omega \) is assumed to be about the vector \((0, 0, 1)\) passing through point \((0, 0, 0)\).

Parent Block(s) ARIA_REGION

9.1.9 BC Dirichlet = BC CIRC_Y

BC CIRC_Y DIRICHLET AT MESH_PART DOF AMP = Ω

Description Dirichlet boundary condition for the y component of a vector variable with constant tangential magnitude along circles of radius \( r(x, y) \) defined on the mesh entity.
Summary

Sets $DOF$ to the provided value on nodeset associated with the name $MESH\_PART$ according to the relation

$$DOF = \Omega r(x, y) \cos \theta.$$  \hspace{1cm} (9.2)

The rotation $\Omega$ is assumed to be about the vector $(0, 0, 1)$ passing through point $(0, 0, 0)$.

Parent Block(s) ARIA\_REGION

### 9.1.10 BC Dirichlet = BC CONST

**BC CONST DIRICHLET AT MESH\_PART DOF = REAL**

Description

Constant Dirichlet condition.

Summary

Sets $DOF$ to the provided constant value on mesh entity associated with the name $MESH\_PART$.

Parent Block(s) ARIA\_REGION

### 9.1.11 BC Dirichlet = BC DISTRIBUTION FACTOR

**BC DISTRIBUTION FACTOR DIRICHLET AT MESH\_PART DOF SCALING = REAL**

Description

Dirichlet condition from node distribution factors.

Summary

Sets $DOF$ to the a scaled value of the distribution factors present on mesh entity associated with nodeset of name $MESH\_PART$.

Parent Block(s) ARIA\_REGION

### 9.1.12 BC Dirichlet = BC LINEAR

**BC LINEAR DIRICHLET AT MESH\_PART DOF COEF = $C_0 \ C_1 \ C_2 \ C_3$**

Description

Dirichlet boundary condition that varies spatially over a mesh entity in a linear fashion.

Summary

Sets $DOF$ to the provided value on nodeset associated with the name $MESH\_PART$ in the following manner

$$DOF = C_0 + C_1x + C_2y + C_3z.$$  \hspace{1cm} (9.3)

Parent Block(s) ARIA\_REGION
9.1.13 BC Dirichlet = BC LINEAR_IN_TIME

**BC LINEAR_IN_TIME DIRICHLET AT MESH_PART DOF COEF = C₀ C₁**

**Description**  
Dirichlet boundary condition whose value is a linear function in time.

**Summary**  
Sets DOF to the provided value on nodeset associated with the name MESH_PART according to the relation

$$DOF = C₀ + C₁t \quad (9.4)$$

**Parent Block(s)** ARIA_REGION

9.1.14 BC Dirichlet = BC PARAB

**BC PARAB DIRICHLET AT MESH_PART DOF COEF = C₀ C₁ C₂ C₃ C₄ C₅ C₆ C₇ C₈ C₉**

**Description**  
Dirichlet boundary condition for DOF that varies spatially over a mesh entity in a parabolic fashion.

**Summary**  
Sets DOF to the provided value on nodeset associated with the name MESH_PART in the following manner

$$DOF = C₀ + C₁x + C₂y + C₃z + C₄xy + C₅xz + C₆yz + C₇x² + C₈y² + C₉z². \quad (9.5)$$

**Parent Block(s)** ARIA_REGION

9.1.15 BC Dirichlet = BC PERIODIC_LINEAR_IN_TIME

**BC PERIODIC_LINEAR_IN_TIME DIRICHLET AT MESH_PART DOF COEF = C₀ C₁ C₂ C₃**

**Description**  
Dirichlet boundary condition that provides a periodic linear (ramp) function in time over a portion of the local time period \(τ = C₃\).

**Summary**  
The value of DOF is given by a periodic linear (ramp) function in time, within a local time interval \(τ\) as illustrated in figure below.

$$DOF = \begin{cases} 
C₀ + C₁t_p & t_p \leq t_d \\
C₀ & \text{otherwise}
\end{cases} \quad (9.6)$$

where \(t_p = t - τ\text{Int}(t/τ)\) is a local time period and \(t_d = C₂\) is the dwell time within this time period. Note that \(t_d < τ\) in order for the boundary condition to be uniquely defined.
9.1.16 BC Dirichlet = BC PERIODIC_STEP_IN_TIME

BC PERIODIC_STEP_IN_TIME DIRICHLET AT MESH_PART DOF COEF = \( C_0 \ C_1 \ C_2 \ C_3 \)

**Description**  
Dirichlet boundary condition that provides a periodic step function in time over a portion of the local time period \( \tau = C_3 \).

**Summary**  
The value of \( DOF \) is given by a periodic step function in time, as illustrated in figure below.

\[
DOF = \begin{cases} 
C_0 & t_p \leq t_d \\
C_1 & \text{otherwise}
\end{cases} 
\]  
(9.7)

where \( t_p = t - \tau \text{Int}(t/\tau) \) is a local period time and \( t_d = C_2 \) is the dwell time within this time period. Note that \( t_d < \tau \) in order for the boundary condition to be uniquely defined.

9.1.17 BC Dirichlet = BC RAMP_LINEAR_IN_TIME

BC RAMP_LINEAR_IN_TIME DIRICHLET AT MESH_PART DOF COEF = \( C_0 \ C_1 \ C_2 \)

**Description**  
Dirichlet boundary condition that provides a linear function in time over a range \( t = 0 \) to \( t = C_2 \) and constant value thereafter.
The value of \( \text{DOF} \) is given by a linear function of time, as illustrated in figure below.

\[
\text{DOF} = C_0 + (C_1 - C_0) \min(t/C_2, 1.0)
\]  

(9.8)

where \( C_0 \) is the initial value, \( C_1 \) is the final value and \( C_2 \) is the time at which the final value \( C_1 \) is achieved.

\[C0\]
\[\uparrow\]
\[C1\]
\[\downarrow\]
\[C2\]
\[t\]

9.1.18 BC Dirichlet = BC ROTATING _X

**Description**  
Dirichlet boundary condition that applies the \( x \)-component of a rotation in time about the \( z \)-axis.

**Summary**  
Sets \( \text{DOF} \) to the provided value on nodeset associated with the name \( \text{MESH_PART} \) according to the relation

\[
\text{DOF}_X = X_0 (\cos \omega t - 1) - Y_0 \sin \omega t
\]  

(9.9)

The rotation \( \Omega \) is assumed to be about the vector \((0, 0, 1)\) passing through point \((0, 0, 0)\).

9.1.19 BC Dirichlet = BC ROTATING _Y

**Description**  
Dirichlet boundary condition that applies the \( y \)-component of a rotation in time about the \( z \)-axis.

**Summary**  
Sets \( \text{DOF} \) to the provided value on nodeset associated with the name \( \text{MESH_PART} \) according to the relation

\[
\text{DOF}_Y = X_0 \sin \omega t + Y_0 (\cos \omega t - 1)
\]  

(9.10)

The rotation \( \Omega \) is assumed to be about the vector \((0, 0, 1)\) passing through point \((0, 0, 0)\).
9.1.20 BC Dirichlet = BC TRANSLATE

BC TRANSLATE DIRICHLET AT MESH_PART DOF SCALE = $C_0$

Description  
Dirichlet boundary condition for translating a value in time.

Summary  
Sets $DOF$ to the provided value on nodeset associated with the name $MESH\_PART$ according to the relation

$$DOF = C_0 t$$  \hfill (9.11)

Parent Block(s)  
ARIA\_REGION

9.1.21 BC Dirichlet = BC UNIDIRECTIONAL\_FLOW\_X

BC UNIDIRECTIONAL\_FLOW\_X DIRICHLET AT MESH_PART DOF coef = $c_0$ $c_1$ $c_2$

Description  
A specialized boundary condition for conveniently specifying inflow boundary conditions on simple geometries.

Summary  
This boundary condition is really meant for specifying inflow velocity profiles in a convenient way. The three coefficients provide the magnitudes of plug, shear and parabolic flow as a function of the $y$-coordinate. Specifically,

$$DOF = c_0 + c_1 \frac{y - y_o}{H} + c_2 \left[ 1 - \left( \frac{y - y_o}{H} \right)^2 \right]$$  \hfill (9.12)

Here $y_o$ is the $y$-coordinate of the middle of the surface (sideset) or nodelist (nodeset) and $H$ is the half-width of the surface or nodelist. Both $y_o$ and $H$ are automatically calculated by Aria. This combination of flows is represented graphically as

Parent Block(s)  
ARIA\_REGION

9.1.22 BC Dirichlet = BC UNIDIRECTIONAL\_FLOW\_Y

BC UNIDIRECTIONAL\_FLOW\_Y DIRICHLET AT MESH_PART DOF coef = $c_0$ $c_1$ $c_2$

Description  
A specialized boundary condition for conveniently specifying inflow boundary conditions on simple geometries.
Summary

This boundary condition is really meant for specifying inflow velocity profiles in a convenient way. The three coefficients provide the magnitudes of plug, shear and parabolic flow as a function of the $x$-coordinate. Specifically,

$$
DOF = c_0 + c_1 \frac{x - x_0}{H} + c_2 \left[ 1 - \left( \frac{x - x_0}{H} \right)^2 \right]
$$

(9.13)

Here $x_0$ is the $x$-coordinate of the middle of the surface (sideset) or nodelist (nodeset) and $H$ is the half-width of the surface or nodelist. Both $x_0$ and $H$ are automatically calculated by Aria. This combination of flows is represented graphically as

\[ c_0 + c_1 + c_2 \]

Parent Block(s) ARIA_REGION

9.1.23 BC Dirichlet = BC XFER

BC XFER DIRICHLET AT MESH_PART DOF

Description

Constant Dirichlet condition where the value is set via a transfer.

Summary

Allows $DOF$ to set to a value that comes from a transfer operation on mesh entity associated with the name $MESH_PART$. Within the transfer operation the target (destination) Field should be defined as solution->$DOF$ state NEW.

Example

Assuming that a scalar Field for temperature is being provided from Transfer then use of the XFER BC could be defined as follows:

BEGIN PROCEDURE myProcedure
  BEGIN ARIA REGION myRegion
    BC Xfer Dirichlet at surface_2 Temperature
  END ARIA REGION myRegion
  BEGIN transfer my_transfer
    send field afield state none to solution->temperature state new
  END transfer my_transfer
END PROCEDURE myProcedure

Parent Block(s) ARIA_REGION
9.1.24  BC Dirichlet = BC USER FIELD

BC USER FIELD DIRICHLET AT MESH_PART DOF = UserFieldName

Description  Dirichlet boundary condition to provide values as a function of time via a user defined Field.

Summary  This Dirichlet boundary condition provides DOF values as a function of time from a user defined Field. Typically these values would be supplied by an Encore utility or from an external source via the Transfer utility.

Example  Assuming that a scalar Field tf_BC is being provided from Transfer then use of the BC could be defined as follows:

BEGIN PROCEDURE myProcedure
  .
  BEGIN ARIA REGION myRegion
    .
    USER FIELD REAL NODAL SCALAR tf_BC on surface_2
    .
    BC User Field Dirichlet at surface_2 Temperature = tf_BC
    .
    END ARIA REGION myRegion
    BEGIN transfer my_transfer
      .
      send field afield state none to tf_BC state none
      .
      END transfer my_transfer
    .
  END ARIA REGION myRegion
  .
END PROCEDURE myProcedure

Parent Block(s)  ARIA_REGION

9.1.25  BC Dirichlet = BC USER_FUNCTION_IN_TIME

BC USER_FUNCTION_IN_TIME DIRICHLET AT MESH_PART DOF = UserFunctionName

Description  Dirichlet boundary condition that provides values as a function of time via a user input function.

Summary  Dirichlet boundary condition that provides values as a function of time via a user input function.
Example

Begin Sierra myJob

.  
Begin Definition for Function T_BC_func
    Type is Piecewise Linear
    Column Titles Time Temperature
    Begin Values
        0  0
        10 50
    End
End
BEGIN PROCEDURE myProcedure
    .  
    BEGIN ARIA REGION myRegion
        .
        BC User_Function_In_Time Dirichlet at surface_3 Temperature = T_BC_func
        .
    END ARIA REGION myRegion
    .
    END PROCEDURE myProcedure
End Sierra myJob

Parent Block(s) ARIA_REGION

9.2 BC FLUX

In Aria, Neumann and Robin conditions are categorized as flux boundary conditions applied to a surface of the problem domain. Here the flux boundary conditions are applied by performing a surface integration of the user prescribed evaluation to generate the appropriate system residual contributions.

The flux boundary condition specification corresponds to a given equation as it appears in the EQ command line. As an example in the EQ command line:

EQ Energy for Temperature on block_2 using Q1 with Diff mass

Energy is the equation to which the residual contributions will be applied.

For thermal analysis another form of Flux BC specification (Calore-style syntax) is described in the Thermal Analysis chapter 33.1. The advantage of this approach is that the syntax is fully supported within the Sandia Analysis Workbench (SAW).

Native Aria boundary conditions differ from those in the Calore-style in that they are succinctly contained within a single line rather than in a command block. In what follows the Constant, Distribution_Factor, Encore_Function and User_Field Flux conditions can be applied to any equation. The remainder of the Flux conditions listed apply to specific equations.

One additional difference between the native Aria and Calore-style flux boundary condition syntax is that the native Aria constant flux condition is defined based upon a mathematical convention rather than an engineering convention. By contrast, a constant flux condition in the Calore-style flux condition respects the engineering sign convention, i.e. energy input is positive while energy output is negative. Thus the native Aria constant flux sign convention is opposite that of the Calore-style convention. For other flux boundary conditions these differences are resolved internal code so that problems are always solved in a consistent manner. Nevertheless, users must be aware of the difference in these sign conventions when setting up thermal problems.
For both the native Aria and Calore-style syntax, specification of boundary conditions on portions of the FEM mesh can be simplified by aggregation of surface parts into Mesh Groups 6.26. Flux boundary conditions available in Aria are listed following a definition of the general syntax and the associated operation.

\[
\text{BC FLUX FOR } EQNAME \text{ ON MESH}_\text{PART} = \text{MODEL}[\text{param}_1 = \text{val}_1, \text{param}_2 = \text{val}_2 \ldots] 
\]

**Description**
Neumann boundary condition that sets the surface normal flux of the degree of freedom associated with equation \( EQNAME \) to that provided by the specified \( \text{MODEL} \).

**Summary**
\( \text{MODEL} \) the supplies a diffusive flux \( f_n \equiv \mathbf{n} \cdot \mathbf{f} \) in accordance with equation 5.3. I.e., it adds the surface integral

\[
\int_{MESH}_\text{PART} \mathbf{n} \cdot \mathbf{f} \phi^i \, dS \quad (9.14)
\]

to the residual for equation \( EQNAME \). See, e.g., \( q_n \) in equation 5.10.

**Parent Block(s)**
ARIA_REGION

### 9.2.1 BC FLUX = CONSTANT

**Parameters**
\( \text{FLUX} = \text{REAL} \)

**Example**
BC Flux for Energy on surface_10 = Constant Flux=3.14159

**Description**
\( \text{FLUX} \) is the value of the constant flux where positive values indicate a loss, i.e., positive flux leaves the volume.

### 9.2.2 BC FLUX = CALORE_USER_SUB

**Parameters**
\( \text{NAME} = \text{STRING} \)
\( \text{TYPE} = \text{STRING} \)
\( \text{MULTIPLIER} = \text{REAL} \)
\( \text{NR} = \text{INT} \)
\( \text{RO} = \text{INT} \)
\( \text{R1} = \text{INT} \) (etc.)
\( \text{NI} = \text{INT} \)
\( \text{IO} = \text{INT} \)
\( \text{I1} = \text{INT} \) (etc.)

**Example**
BC FLUX for Energy on surface_10 = Calore_User_Sub name=w80afftuser
type=element NR=2 R0=1000 R1=0.7

**Description**
\( \text{NAME} \) is the name of the user subroutine was registered with. \( \text{NR} \) and \( \text{NI} \) are the numbers of real and integer parameters, respectively. Each real (int) parameter is supplied by the \( \text{Rn} \) \( \text{In} \).
parameter. **Note:** the parameters use zero based counting. The `TYPE` parameter denotes the API the user subroutine uses; currently Aria only supports types `ELEMENT` and `NODE`.

The optional `MULTIPLIER` parameter (defaults to 1.0) can be supplied to scale the output of the user subroutine, i.e., the flux $f$ will be $f = mf_{user}$ where $m$ is the multiplier and $f_{user}$ is the output of the user subroutine.

The following is an example user subroutine that provides a free-space radiation energy flux. Note that the flux sign convention is that positive values indicate a heat loss, i.e., positive flux leaves the volume.
Example

```c
#include "math.h"
#include "Aria_Calore_User_Sub_Support.h"

Int
w80afftuser(UserQuery & user_query,
Int element_id,
Int num_elements,
Int num_points,
Int spatial_dimension,
CoordinateElementIntegrationPoints coordinate_array,
TemperatureElementIntegrationPoints temperature_array,
FluxElementIntegrationPoints flux_array)
{

RealArray1d rdat(2);

user_query.getUserRealInstanceData(2, rdat.ptr());

const Real Tref = rdat(0);
const Real emissivity = rdat(1);

const Real sb_emissivity = 5.67E-8 * emissivity;
const Real Tref4 = Tref * Tref * Tref * Tref;

for(Int element=0; element < num_elements; ++element)
{
    for(Int point=0; point < num_points; ++point)
    {
        const Real & T = temperature_array(element,point);
        const Real T2 = T*T;
        flux_array(element,point) = sb_emissivity * (T2*T2 - Tref4);
    }
}
return 0;
}

extern "C"
void
#if defined(DYNAMIC_USER_PLUGIN)
dl_register()
#else
SIERRA_FORTRAN(fmwk_reg_user_subs)()
#endif
{
    sierra::Plugin::
        UserSubroutine<aucal_elem_sub_c>::registerFunction("w80afftuser",
                &w80afftuser);
}
```

Description
To use this subroutine, you must add a line command at the domain level of your Aria input file that gives the name of the user subroutine source file, e.g.,
`User Subroutine file = w80afftuser.C`

Then, to build the executable with the user subroutine, you can use a command line like this,
sierra --make aria -i w80afftuser_aria.i
And, finally, the problem can be run like this,

sierra ./UserSubsProject/bin/aria -i w80afftuser_aria.i

Note that the UserSubsProject is created as a part of the build process.

9.2.3 BC FLUX = DISTRIBUTION_FACTOR

Parameters  MULTIPLIER = REAL

Example  BC Flux for Energy on MESH_PART= distribution_factor multiplier=3.2

Description  Sets the surface flux to the value of the distribution factor scaled by the MULTIPLIER parameter. Here the distribution factor is present on a mesh entity associated with a sideset (surface) of named MESH_PART.

9.2.4 BC FLUX = ENCLOSURE_RADIATION

Parameters  [MULTIPLIER = REAL]

Example  BC Flux for Energy on surface_10 = Enclosure_Radiation

Description  This boundary condition incorporates the heat flux that’s computed using Chaparral for enclosure radiation.
See chapter 22 for more information on enclosure radiation.

9.2.5 BC FLUX = ENCORE_FUNCTION

Parameters  NAME = STRING
[EVAL_TYPE = STRING]

Example  BC Flux for Energy on surface_10 = Encore_Function Name=My_Function

Description  Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```
Begin String Function My_Function
    Value is "200 + 1.0*t"
End
```
Then, this function can be used as shown in the example above.

The optional argument `EVAL_TYPE` can be used to select the particular function on the Encore function object. Valid options include `VALUE`, `DOT`, `GRADIENT`, `FLUX` and `STRESS`. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – `VALUE` for `NO_OP` expressions, `DOT` for `DT_OP` expressions and `GRADIENT` for `GRAD_OP` expressions.

### 9.2.6 BC FLUX = LASER

**Parameters**

- `T_ON = REAL`
- `T_OFF = REAL`
- `X_0 = REAL`
- `Y_0 = REAL`
- `R0 = REAL`
- `W = REAL`
- `FLUX = REAL`
- `R = REAL`

**Example**

```
BC Flux for Energy on surface_10 = Laser t_on=0 t_off=10 X_0=0 Y_0=0 w=1.0 r=0.025 flux=3.14159 r0=0.01
```

**Description**

This boundary condition imposes an energy flux due to an incident laser. The laser is directed in the \( z \) direction and travels in a circular path of radius \( r_0 \).

- `T_ON` is the time when the laser is turned on.
- `T_OFF` is the time when the laser is turned off.
- `X_0` is the \( x \) coordinate of the center of the laser path.
- `Y_0` is the \( y \) coordinate of the center of the laser path.
- `R0` is the radius of the circular laser path.
- `W` is the angular velocity of the laser.
- `FLUX` is the value of the energy flux into the surface. (The usual convention is that positive fluxes indicate loss.)
- `R` is the radius of the laser beam.

### 9.2.7 BC FLUX = LASER_WELD

**Parameters**

- `PATH_FUNCTION = STRING`
- `FLUX = REAL`
- `R = REAL`
- `[NORMAL_TOLERANCE = REAL]`

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Example
# This function lives at the top level, inside the 'Begin Sierra' block

Begin Definition for Function PATH
Type is Multicolumn Piecewise Linear
Column Titles Time X Y Z
Begin Values
0.0000000E+00 0.0000000E+00 0.0000000E+00 0.0000000E+00
0.2250000E+00 0.1500000E+00 0.0000000E+00 0.0000000E+00
0.2328540E+00 0.1552336E+00 -0.1370490E-03 0.0000000E+00
...
End
End

# This goes in the Aria Region with all of the other BCs
BC Flux for Energy on surface_10 = Laser Path Function=PATH R=0.025
Flux=3.14159

Description
This boundary condition imposes an energy flux due to an incident laser. The position of the laser is dictated by a user supplied function which contains \( x \), \( y \) and \( z \) coordinates as a function of time; Aria uses linear piecewise interpolation to obtain the location of the laser at a given time.

PATH is the name of the user supplied function which has time as the first column and contains columns titled \( X \), \( Y \) and \( Z \) containing the \( x \), \( y \) and \( z \) coordinates.

FLUX is the value of the energy flux into the surface. (The usual convention is that positive fluxes indicate loss.)

\( R \) is the radius of the laser beam.

The determination of whether a point in space has an incident laser flux is done as follows. At time \( t \) the laser is at a point \( P \) and the Laser_Weld boundary condition is being evaluated at a point \( x \) (typically an integration point). The distance of \( x \) from \( P \) is defined as \( r \equiv P - x \). If a tolerance NORMAL_TOLERANCE is supplied and the out-of-surface portion of \( r \) is greater than the tolerance then the flux is taken to be zero. Otherwise, the out-of-surface portion of \( r \) is subtracted to neglect minor differences between the PATH_FUNCTION and the discretized mesh, \( r_s \equiv r - (n \cdot r)n \) where \( n \) is the outward unit normal at \( x \). If \( |r_s| < R \) then the point \( x \) is considered to be in the beam and the flux is applied; otherwise, no flux is applied.
9.2.8 BC FLUX = GAUSSIAN_LINE_WELD

Parameters

SR_C_X = REAL
SR_C_Y = REAL
[SR_C_Z = REAL]
DIR_X = REAL
DIR_Y = REAL
[DIR_Z = REAL]
VEL_X = REAL
VEL_Y = REAL
[VEL_Z = REAL]
FLUX = REAL
R = REAL
R_EFF = REAL
[T_ON = REAL]
[T_OFF = REAL]
[ALPHA = REAL]
[COMPUTE_VISIBILITY_FIELD = STRING]
[DEPTH_ENHANCED_LASER_ABSORPTION_COEFFICIENT = STRING]

Example
BC Flux for Energy on surface_10 = Gaussian_Line_Weld Src_X=0 Src_Y=3
Dir_X=0 Dir_Y=-1 Flux=1000 R=0.01 VEL_X=0.1 VEL_Y=0

Description
This boundary condition imposes an energy flux due to an incident laser. The laser source is at the coordinates
\[ s = x_s + v_s(t-t_{on}) \]  
where \( x_s \) is provided by the SRC vector coordinates and \( v_s \) is provided by the VEL vector coordinates. The laser is directed from \( s \) by the DIR vector (\( d \) in the figure below).

The point-wise energy flux, \( f_{eff} \), is computed using a Gaussian distribution based on the effective radius \( R_{eff} \) (\( R_EFF \)),
\[ f_{eff} = 2R_{eff}e^{-\frac{R_{eff}^2}{r_p^2}} \]  
where \( r_p \) is the radial coordinate from the center of the incident laser.
The radius of the laser is provided by $R$ ($R$ in the figure). A positive FLUX is defined as energy input to the surface. That is, the net normal flux is

$$q_n = \frac{(-n) \cdot d}{|d|} (-f_{eff}) = \frac{n \cdot d}{|d|} f_{eff}$$

(9.17)

Here, the term $-n \cdot d$ accounts for the fact that the surface may not be orthogonal to the laser.

The optional parameter **ALPHA** is the absorption coefficient for the incident laser flux (defaults to 1.0).

The optional parameter **COMPUTE_VISIBILITY_FIELD** is used to enable the calculation of the visibility of the laser to the given surface. When this parameter is given, the provided field will be populated with 1 for each node of the surface that is visible to the laser, and 0 for each node that is obstructed by other portions of the surface. This field is then used to multiply the incident flux so that the obstructed portions of the surface do not get heated by the laser.

The optional parameter **DEPTH_ENHANCED_LASER_ABSORPTION_COEFFICIENT** is used to specify a filename containing tabular data that will be used to compute an absorption coefficient that depends on the relative depth of the point and the aspect ratio of the given surface, defined as the surface depth/$R$. The text file is formatted as follows. The first line contains a list of the relative depth values. The second line contains of list of aspect ratio values. The table rows are for increasing depth values for a fixed aspect ratio. The rows are for increasing aspect ratios. Bilinear interpolation is used to evaluated the enhanced absorption coefficient for each point on the surface. For any point that lies outside of the specified range of relative depth or aspect ratio values, the value will be extrapolated using the nearest bilinear cell.

The net absorption coefficient is the product of **ALPHA** (defaults to 1.0), the results of the optional **COMPUTE_VISIBILITY_FIELD** calculation, and the optional **DEPTH_ENHANCED_LASER_ABSORPTION_COEFFICIENT** calculation.

### 9.2.9 BC FLUX = GAUSSIAN_SPOT_WELD

**Parameters**

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>SRC_X</td>
<td>REAL</td>
</tr>
<tr>
<td>SRC_Y</td>
<td>REAL</td>
</tr>
<tr>
<td>[SRC_Z = REAL]</td>
<td></td>
</tr>
<tr>
<td>DIR_X</td>
<td>REAL</td>
</tr>
<tr>
<td>DIR_Y</td>
<td>REAL</td>
</tr>
<tr>
<td>[DIR_Z = REAL]</td>
<td></td>
</tr>
<tr>
<td>FLUX</td>
<td>REAL</td>
</tr>
<tr>
<td>R = REAL</td>
<td></td>
</tr>
<tr>
<td>R_EFF = REAL</td>
<td></td>
</tr>
<tr>
<td>[T_ON = REAL]</td>
<td></td>
</tr>
<tr>
<td>[T_OFF = REAL]</td>
<td></td>
</tr>
<tr>
<td>[ALPHA = REAL]</td>
<td></td>
</tr>
<tr>
<td>[COMPUTE_VISIBILITY_FIELD = STRING]</td>
<td></td>
</tr>
<tr>
<td>[DEPTH_ENHANCED_LASER_ABSORPTION_COEFFICIENT = STRING]</td>
<td></td>
</tr>
</tbody>
</table>

**Example**

BC Flux for Energy on surface_10 = Gaussian_Spot_Weld Src_X=0 Src_Y=3
Dir_X=0 Dir_Y=-1 Flux=1000 R=0.01

**Description**

This boundary condition imposes an energy flux due to an incident laser. The laser source is at the coordinates provided by the SRC vector coordinates ($s$ in the figure below) and it is directed from there as specified by the DIR vector ($d$ in the figure below).
The point-wise energy flux, $f_{\text{eff}}$, is computed using a Gaussian distribution based on the effective radius $R_{\text{eff}}$ (R\_EFF),

$$f_{\text{eff}} = 2R_{\text{eff}}e^{-\frac{R_{\text{eff}}^2}{r^2}}$$  \hspace{1cm} \text{(9.18)}

where $r_p$ is the radial coordinate from the center of the incident laser.

The radius of the laser is provided by $R$ ($R$ in the figure). A positive FLUX is defined as energy input to the surface. That is, the net normal flux is

$$q_n = \left(\frac{-\boldsymbol{n} \cdot \boldsymbol{d}}{|\boldsymbol{d}|}\right)(-f_{\text{eff}}) = \frac{\boldsymbol{n} \cdot \boldsymbol{d}}{|\boldsymbol{d}|}f_{\text{eff}}$$  \hspace{1cm} \text{(9.19)}

Here, the term $-\boldsymbol{n} \cdot \boldsymbol{d}$ accounts for the fact that the surface may not be orthogonal to the laser.

The optional parameter $\text{ALPHA}$ is the absorption coefficient for the incident laser flux (defaults to 1.0).

The optional parameter $\text{COMPUTE_VISIBILITY_FIELD}$ is used to enable the calculation of the visibility of the laser to the given surface. When this parameter is given, the provided field will be populated with 1 for each node of the surface that is visible to the laser, and 0 for each node that is obstructed by other portions of the surface. This field is then used to multiply the incident flux so that the obstructed portions of the surface do not get heated by the laser.

The optional parameter $\text{DEPTH_ENHANCED_LASER_ABSORPTION_COEFFICIENT}$ is used to specify a filename containing tabular data that will be used to compute an absorption coefficient that depends on the relative depth of the point and the aspect ratio of the given surface, defined as the surface depth/$R$. The text file is formatted as follows. The first line contains a list of the relative depth values. The second line contains of list of aspect ratio values. The table rows are for increasing depth values for a fixed aspect ratio. The rows are for increasing aspect ratios. Bilinear interpolation is used to evaluated the enhanced absorption coefficient for each point on the surface. For any point that lies outside of the specified range of relative depth or aspect ratio values, the value will be extrapolated using the nearest bilinear cell.

The net absorption coefficient is the product of $\text{ALPHA}$ (defaults to 1.0), the results of the optional $\text{COMPUTE_VISIBILITY_FIELD}$ calculation, and the optional $\text{DEPTH_ENHANCED_LASER_ABSORPTION_COEFFICIENT}$ calculation.

9.2.10 \text{BC FLUX = GENERALIZED\_ NAT\_ CONV}

Parameters \hspace{1cm} [MULTIPLIER = \text{REAL}]
Example  BC Flux for Energy on surface_10 = Generalized_Nat_Conv

Description  This boundary condition adds a heat flux due to Newton’s law of cooling. This is similar to the NAT_CONV boundary condition except that it utilizes the HEAT_TRANSFER_COEFFICIENT and BC_REFERENCE_TEMPERATURE models.

\[ q = mh(T - T_{\text{ref}}) \]  

(9.20)

where \( m \) is the optional MULTIPLIER parameter (defaults to 1.0), \( h \) is the heat transfer coefficient, \( T \) is the temperature at the surface and \( T_{\text{ref}} \) is a reference temperature.

9.2.11 BC FLUX = GENERALIZED_RAD

Parameters  
[MULTIPLIER = REAL]

Example  BC Flux for Energy on surface_10 = Generalized_Rad

Description  This boundary condition accounts for the heat flux due to radiation in free space, similar to the RAD boundary condition but the model is more general in that it uses specific models for EMISSIVITY, RADIATION FORM FACTOR and BC RAD REFERENCE TEMPERATURE. The functional form of this boundary condition is

\[ q = mF\sigma(T^4 - T_{\text{ref}}^4) \]  

(9.21)

where \( m \) is the optional MULTIPLIER parameter (defaults to 1.0), \( F \) is the RADIATION FORM FACTOR, \( \sigma \) is the Stefan-Boltzmann constant, \( \epsilon \) is the emissivity of the surface, \( T \) is the temperature at the surface and \( T_{\text{ref}} \) is a reference temperature which must be supplied from a BC RAD REFERENCE TEMPERATURE material model.

The Stefan-Boltzmann constant must be defined in the input file using a Global Constants block, e.g.,

Begin Global Constants
  Stefan Boltzmann Constant = 5.67e-14
End

9.2.12 BC FLUX = SHARP_LINE_WELD

Parameters  
SRC_X = REAL
SRC_Y = REAL
[SRC.Z = REAL]
DIR_X = REAL
DIR_Y = REAL
[DIR.Z = REAL]
VEL_X = REAL
VEL_Y = REAL
[VEL.Z = REAL]
FLUX = REAL
R = REAL
[T.ON = REAL]
[T.OFF = REAL]
Example  BC Flux for Energy on surface_10 = Sharp_Line_Weld Src_X=0 Src_Y=3 Dir_X=0 Dir_Y=-1 Flux=1000 R=0.01 VEL_X=0.1 VEL_Y=0

Description  This boundary condition imposes an energy flux due to an incident laser. The laser source is at the coordinates

\[ s = x_s + v_s(t - t_{on}) \]  

(9.22)

where \( x_s \) is provided by the SRC vector coordinates and \( v_s \) is provided by the VEL vector coordinates. The laser is directed from \( s \) by the DIR vector (\( d \) in the figure below).

\[ p \]

\[ s \]

\[ h \]

\[ d \]

\[ g \]

\[ r_p \]

\[ R \]

The radius of the laser is provided by \( R \) (\( R \) in the figure) and the energy flux is provided by the FLUX argument (written as \( f \) below). A positive FLUX is defined as energy input to the surface. That is, the net normal flux is

\[ q_n = \frac{(-n) \cdot d}{|d|} (-f) = \frac{n \cdot d}{|d|} f \]  

(9.23)

Here, the term \(-n \cdot d\) accounts for the fact that the surface may not be orthogonal to the laser.

9.2.13  BC FLUX = SHARP_SPOT_WELD

Parameters  

SRC_X = REAL  
SRC_Y = REAL  
[SRC_Z = REAL]  
DIR_X = REAL  
DIR_Y = REAL  
[DIR_Z = REAL]  
FLUX = REAL  
R = REAL  
[T_ON = REAL]  
[T_OFF = REAL]

Example  BC Flux for Energy on surface_10 = Sharp_Spot_Weld Src_X=0 Src_Y=3 Dir_X=0 Dir_Y=-1 Flux=1000 R=0.01
This boundary condition imposes an energy flux due to an incident laser. The laser source is at the coordinates provided by the \( \text{SRC} \) vector coordinates (\( s \) in the figure below) and it is directed from there as specified by the \( \text{DIR} \) vector (\( d \) in the figure below).

The radius of the laser is provided by \( R \) (\( R \) in the figure) and the energy flux is provided by the \( \text{FLUX} \) argument (written as \( f \) below). A positive \( \text{FLUX} \) is defined as energy input to the surface. That is, the net normal flux is

\[
q_n = \frac{(-n) \cdot d}{|d|} (-f) = \frac{n \cdot d}{|d|} f
\]

(9.24)

Here, the term \(-n \cdot d\) accounts for the fact that the surface may not be orthogonal to the laser.

### 9.2.14 \( \text{BC FLUX} = \text{LATENT\_HEAT} \)

**Parameters**

\[ Y_0 = \text{REAL} \]
\[ \text{SPECIES} = \text{INT} \]

**Example**

BC Flux for Energy on surface\_10 = Latent_Heat \( Y_0=0.2 \) \( \text{SPECIES}=0 \)
BC Flux for Energy on surface\_10 = Latent_Heat \( Y_0=0.4 \) \( \text{SPECIES}=1 \)
BC Flux for Energy on surface\_10 = Latent_Heat \( Y_0=0.0 \) \( \text{SPECIES}=2 \)

**Description**

This boundary condition accounts for the heat flux due to the latent heat of vaporization (evaporation).

\[
q = H_v \rho (Y_i - Y_{\infty,i})
\]

(9.25)

where \( H_v,i \) is the heat of vaporization of species \( i \), \( \rho \) is the density of the material, \( Y_i \) is the mass fraction of species \( i \) and \( Y_{\infty,i} \) is the mass fraction of species \( i \) far from the surface.

\( \text{SPECIES} \) is the index of the species to use (in multicomponent systems), \( = i \) in equation 9.25.

\( Y_{\infty} \) is the mass fraction of species \( i \) far from the surface, \( = Y_{\infty,i} \) in equation 9.25.

### 9.2.15 \( \text{BC FLUX} = \text{NAT\_CONV} \)

**Parameters**

\[ T_{\text{REF}} = \text{REAL} \]
\[ H = \text{REAL} \]
Example

BC Flux for Energy on surface_10 = Nat_Conv T_REF=273.15 H=300

Description

This boundary condition accounts for the heat flux due to natural heat convection:

\[ q = h(T - T_{ref}) \]  

(9.26)

\( T_{REF} \) is the reference temperature of free space.
\( H \) is the heat transfer coefficient.

9.2.16 BC FLUX = RAD

Parameters

\( T_{REF} = \text{REAL} \)
\( CRAD = \text{REAL} \)

Example

BC Flux for Energy on surface_10 = Rad T_REF=273.15 CRAD=1.3e-8

Description

This boundary condition accounts for the heat flux due to radiation in free space:

\[ q = c_{rad}(T^4 - T_{ref}^4) \]  

(9.27)

\( T_{REF} \) is the temperature of free space.
\( CRAD \) is the coefficient that multiplies the radiation term, viz., the product of the emissivity and the Stefan-Boltzmann constant, which is \( 5.67 \times 10^{-08} \text{ W m}^{-2} \text{ K}^{-4} \) (SI Units).

9.2.17 BC FLUX = VAPOR_COOLING

Parameters

\( TBOIL = \text{REAL} \)

Example

BC Flux for Energy on surface_10 = Vapor_Cooling Tboil=3000

Description

This boundary condition accounts for the cooling of a material due to vaporization. See Allen Roach (raroach@sandia.gov) for more information.
\( TBOIL \) is the boiling point of the material.

9.2.18 BC FLUX = NAT_CONV

Parameters

\( Yinf = \text{REAL} \)
\( k = \text{REAL} \)

Example

BC Flux for Species_2 on surface_10 = Nat_Conv k=0.15 Yinf=0.8

Description

This boundary condition accounts for the mass flux due to natural convection:

\[ q = k(Y - Y_{\infty}) \]  

(9.28)

\( Yinf \) is the bulk species concentration, \( Y_{\infty} \).
\( k \) is the mass transfer coefficient.
9.2.19 BC FLUX = CAPILLARY

Parameters (none)

Example BC FluxBP for Momentum on surface_10 = Capillary

Description This boundary condition implements the capillary (surface tension) contributions to the traction boundary condition [19]. Specifically, this boundary condition adds to the \( k \)th component of the \( i \)th momentum residual

\[
\int_S \sigma (I - nn) : \nabla (\phi^i e^k) \, dS. 
\]  

(9.29)

where \( \sigma \) is the surface tension and \( n \) is the unit outward normal to the interface. This condition accounts for both the curvature and surface tension gradient contributions to the traction condition.

9.2.20 BC FLUX = CONSTANT_TRACTION

Parameters [\( X = REAL \)]
[\( Y = REAL \)]
[\( Z = REAL \)]

Example BC Flux for Momentum on surface_10 = Constant_Traction \( Y=0.5 \)
or BC Flux for Mesh on surface_10 = Constant_Traction \( X=0.1 \) \( Y=0.5 \)
or BC Flux for Solid on surface_10 = Constant_Traction \( Z=0.5 \)

Description This boundary condition integrates a constant and uniform traction over a surface for either fluid momentum, mesh elasticity or solid elasticity. Specifically, this boundary condition adds to the \( k \)th component of the \( i \)th momentum/mesh/solid residual

\[
\int_S f_i \phi^i \, dS. 
\]  

(9.30)

where \( f_i \) is the constant traction vector whose components are given by the parameters \( X, Y \) and \( Z \).

9.2.21 BC FLUX = ELECTRIC_TRACTION

Parameters [\( SIGN = REAL \)]

Example BC Flux for Momentum on surface_10 = Electric_Traction
BC Flux for Mesh on surface_10 = Electric_Traction
BC Flux for Solid on surface_10 = Electric_Traction
This boundary condition adds the stress contribution for due to the presence of an electric field. Here, the electric stress tensor is taken to be

\[ T_e = \varepsilon EE - \frac{1}{2} \varepsilon E \cdot EI \]  

(9.31)

where \( E = -\nabla V \) is the electric field and \( V \) is voltage and \( \varepsilon \) is the electric permittivity. The electric traction is then defined as

\[ t = \varepsilon n \cdot EE - \frac{1}{2} \varepsilon E \cdot En \]  

(9.32)

where \( n \) is the outward normal to the boundary.

### 9.2.22 BC FLUX = NAT_E

**Example**

BC Flux for Charge Density on surface 1 = Nat_E

**Description**

This boundary condition creates a no-flux boundary condition for the charge density equation when DIFF is activated in the equation. The diffusion operator for this equation is not of the standard no-flux type, and so this term needs to be added to get the correct surface term contribution.

### 9.2.23 BC FLUX = FLOW_HYDROSTATIC

**Parameters**

- \([GX = \text{REAL}]\)
- \([GY = \text{REAL}]\)
- \([GZ = \text{REAL}]\)
- \([P_{\text{REF}} = \text{REAL}]\)

**Example**

BC Flux for Momentum on surface 10 = Flow_Hydrostatic Gy=-9.8

**Description**

This boundary condition provides a hydrostatic pressure head along a boundary. The acceleration vector \( g \) is specified with the parameters \( GX, GY \) and \( GZ \) (all default to zero). The reference pressure \( P_{\text{ref}} \) (\( P_{\text{REF}}, \) defaults to zero) is taken to be the pressure datum at the origin \((0, 0, 0)\). Specifically, this BC adds

\[ n \cdot T = - \left( P_{\text{ref}} + \sum_{i} \rho g_i x_i \right) n \]  

(9.33)

to the momentum equation. Here \( N_d \) is the spatial dimension of the problem, \( g_i \) are the components of the acceleration, \( x_i \) are the coordinate components and \( \rho \) is the density. **NOTE:** This BC evaluates the density material model for \( \rho \) thus the parameters for \( g \) should not include the density.

### 9.2.24 BC FLUX = FREE_OPEN_FLOW

**Parameters**

- \( \text{PRESSURE} = \text{REAL} \)
Example: BC Flux for Momentum on surface_10 = Free_Open_Flow Pressure=1000

Description: This boundary condition adds back the stress contribution for inlet/outlet flows where the pressure is prescribed. This command line is used in open-flow applications to set the pressure datum but imposes no other constraints on the flow profile.

\[ q = -p_0 n + \mu n \cdot (\nabla v + \nabla v^T) \]  
\[ (9.34) \]

Here, \( p_0 \) is the pressure provided by the user, \( \mu \) is the viscosity, \( \nabla v \) is the velocity gradient, and \( n \) is the outward normal to the boundary.

9.2.25 BC FLUX = OPEN_FLOW

Parameters

PRESSURE = REAL

Example: BC Flux for Momentum on surface_10 = Open_Flow Pressure=1000

Description: This boundary condition adds back the stress contribution for inlet/outlet flows where the pressure is prescribed, and the flow is unidirectional and fully developed. Specifically, it removes the normal component of the viscous stresses (and any other non-pressure stresses).

\[ q = -p_0 n + n \cdot \tau \cdot (I - nn) \]  
\[ (9.35) \]

Here, \( p_0 \) is the pressure provided by the user \( n \) is the outward normal to the boundary and \( \tau \) is the sum of all other stresses except the the isotropic pressure (e.g., NEWTONIAN_PRESSURE_MOMENTUM_STRESS).

9.2.26 BC FLUX = PRESSURE

Parameters

\[ P = REAL \]
\[ [C_T = REAL] \]

Example: BC Flux for Momentum on surface_10 = Pressure P=101325
or
BC Flux for Mesh on surface_10 = Pressure P=101325
or
BC Flux for Solid on surface_10 = Pressure P=101325

Description: This boundary condition integrates a uniform pressure over a surface for either fluid momentum, mesh elasticity or solid elasticity. The optional parameter \( C_T \) allows the pressure to vary linearly in time. Specifically, this boundary condition adds to the \( k^{th} \) component of the \( i^{th} \) momentum/mesh/solid residual

\[ \int_S - (p + c_t t) n \phi^i \, dS. \]  
\[ (9.36) \]

where \( p \) is the pressure provided by the parameter \( P \), \( t \) is time and \( c_t \) is a constant provided by the \( C_T \) parameter.
### 9.2.27 BC FLUX = LUBRICATION_PRESSURE

**Parameters**

- (none)

**Example**

- BC Flux for Mesh on surface_10 = Lubrication_Pressure
- BC Flux for Solid on surface_10 = Lubrication_Pressure

**Description**

This boundary condition integrates the lubrication pressure (the DOF from the lubrication equation) over a surface. This is intended to apply the fluid pressure to a solid region, and can be used for either the MESH or SOLID equations.

### 9.2.28 BC FLUX = ORIENTED_SLIP

**Parameters**

- BETA_NORMAL = REAL
- BETA_TANGENT = REAL
- VS_X = REAL
- VS_Y = REAL
- VS_Z = REAL

**Example**

- BC Flux for Momentum on surface_7 = Oriented_Slip Beta_Tangent=1e-1 Beta_Normal=1e-5

**Description**

This is similar to the classic SLIP condition except that it permits separate slip coefficients for the normal and tangent directions.

\[
q = n \cdot T = \frac{1}{\beta_t} (v_s - v) + \left( \frac{1}{\beta_n} - \frac{1}{\beta_t} \right) (v_s - v) \cdot nn \tag{9.37}
\]

where \( \beta_t \) is the Navier slip coefficient for the tangent direction (BETA_TANGENT), \( \beta_n \) is that for the normal direction (BETA_NORMAL), \( v \) is the fluid velocity and \( v_s \) is the velocity of the surface. The surface velocity is zero by default but a nonzero velocity can be supplied in component form using one or more of the optional VS_X, VS_Y and VS_Z parameters.

### 9.2.29 BC FLUX = SLIP

**Parameters**

- BETA = REAL
- VS_X = REAL
- VS_Y = REAL
- VS_Z = REAL

**Example**

- BC Flux for Momentum on surface_10 = Slip Beta = 0.01

**Description**

This boundary condition implements the Navier slip boundary condition along a surface wherein the tangential velocity along the surface is proportional to the fluid stress,

\[
q = n \cdot T = \frac{1}{\beta} (v_s - v) \tag{9.38}
\]
where $\beta$ is the Navier slip coefficient, $v$ is the fluid velocity and $v_s$ is the velocity of the surface. The surface velocity is zero by default but a nonzero velocity can be supplied in component form using one or more of the optional VS_X, VS_Y and VS_Z parameters.

### 9.2.30 BC FLUX = TRANSIENT_TRACTION

**Parameters**
- $[A_X = \text{REAL}]$
- $[A_Y = \text{REAL}]$
- $[A_Z = \text{REAL}]$
- $[B_X = \text{REAL}]$
- $[B_Y = \text{REAL}]$
- $[B_Z = \text{REAL}]$

**Example**
- BC Flux for Momentum on surface_10 = Transient_Traction B_Y=0.5
- or
- BC Flux for Mesh on surface_10 = Transient_Traction A_X=0.1 A_Y=0.5
- or
- BC Flux for Solid on surface_10 = Transient_Traction B_Z=0.5

**Description**
This boundary condition integrates a uniform but time dependent traction over a surface for either fluid momentum, mesh elasticity or solid elasticity. Specifically, this boundary condition adds to the $k^{th}$ component of the $i^{th}$ momentum/mesh/solid residual

$$\int_S (f_a + tf_b) \phi^i \, dS.$$  \hspace{1cm} \text{(9.39)}

where $t$ is time and $f_a$ and $f_b$ are is constant traction vectors whose components are given by the parameters $A_X$, $A_Y$ and $A_Z$ and $B_X$, $B_Y$ and $B_Z$ respectively.

### 9.2.31 BC FLUX = WETTING_SPEED_BLAKE_LS

**Parameters**
- $V_w = \text{REAL}$
- $G = \text{REAL}$
- $\text{THETA} = \text{REAL}$
- $\text{WIDTH} = \text{REAL}$

**Example**
- BC Disting for Momentum_A on surface_3 = Wetting_Speed_Blake_LS V_w=1e-1 g=1 Width=1 Theta=60
- BC Disting for Momentum_B on surface_3 = Wetting_Speed_Blake_LS V_w=1e-1 g=1 Width=1 Theta=60

**Description**
This boundary condition is a distinguishing condition that enforces a slip velocity in accordance with the model provided by [20].

$$v - f(\phi)v_w \sinh (g (\cos \theta_s - \cos \theta_a)) \frac{n_s - \cos \theta_a n_w}{1 - \cos^2 \theta_a} = 0$$ \hspace{1cm} \text{(9.40)}

Here $\theta_s$ is the static or equilibrium contact angle and is provided by the THETA input parameter, $\theta_a$ is the actual (or “observed” or “current”) contact angle, $g$ is a constant parameter given by the G input parameter and $v_w$ is the characteristic slip velocity and is given by the input parameter $V_w$. In their paper, Blake and De Coninck develop $g$ and $v_w$ theoretically.
The function $f(\phi)$ where $\phi$ is the level set distance function is simply a triangle shape function which causes the velocity to vary from $v_w$ to zero over the distance given by the input parameter $\text{WIDTH}$. Taking $h_w$ to half of the input $\text{WIDTH}$ parameter, $f(\phi)$ is given as

$$f(\phi) = \begin{cases} 
0 & \text{if } \phi < -h_w \\
1 + \phi/h_w & \text{if } -h_w \leq \phi < 0 \\
1 - \phi/h_w & \text{if } 0 \leq \phi < h_w \\
0 & \text{if } h_w \leq \phi 
\end{cases}$$

This boundary condition is a distinguishing condition which means the momentum equations are discarded at the nodes where this is applied. Also, the velocity provided by this boundary condition is purely tangential. Thus, this boundary condition automatically enforces no-penetration in addition to slip/no-slip.

**Note:** The interface normal points out of the negative phase (negatively signed level set distance $\phi$) into the positive phase. Thus, the contact angle is measured from the wall, through the negative phase and to the level set interface $\phi = 0$.

### 9.2.32 BC FLUX = USER_FUNCTION

**Parameters**

- **NAME** = STRING
- **X** = STRING

**Example**

BC Flux for Energy on surface_6 = USER_FUNCTION NAME=testflux X=TIME

**Description**

The flux boundary condition value is defined by a tabular function of the given name which is evaluated as a function of argument $X$.

### 9.2.33 Shear Free

**Scope:** Equation System

Shear Free Flux For Solid [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Shear_Free [ Using Data Specification Data Spec Name ]

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**Summary**

Shear-free traction solid flux

### 9.2.34 Electric Traction

**Scope:** Equation System

Electric Traction Flux For Solid [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Electric_Traction [ Using Data Specification Data Spec Name ][ Sign = sign ]
## 9.2.35 Pressure User Function

**Scope:** Equation System

Pressure User Function Flux For Solid [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ][@|at|for|in|on|over] Mesh Extent Name = Pressure_User_Function [ Using Data Specification Data Spec Name ][ Name = name |X = x |X_Multiplier = x_multiplier |Multiplier = multiplier |Toggle = toggle ]

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**Summary** Electric traction solid flux

## 9.2.36 Pressure

**Scope:** Equation System

Pressure Flux For Solid [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ][@|at|for|in|on|over] Mesh Extent Name = Pressure [ Using Data Specification Data Spec Name ][ Use_Bulk_Node = use_bulk_node |P = p |C_T = c_t |C_X = c_x |C_Y = c_y |C_Z = c_z |C_Cos_W = c_cos_w |C_Sin_W = c_sin_w |C1_Cos = cl_cos |C1_Sin = cl_sin ]

**Summary** Pressure user function solid flux
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Summary: Pressure solid flux

9.2.37 Constant Traction

Scope: Equation System

Constant Traction Flux For Solid [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@at|for|in|on|over} Mesh Extent Name = Constant_Traction [ Using Data Specification Data Spec Name ][ X = x | Y = y | Z = z ]

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Summary: Constant traction solid flux

9.2.38 Transient Traction

Scope: Equation System

Transient Traction Flux For Solid [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@at|for|in|on|over} Mesh Extent Name = Transient_Traction [ Using Data Specification Data Spec Name ][ A_X = a_x | A_Y = a_y | A_Z = a_z | B_X = b_x | B_Y = b_y | B_Z = b_z ]
### 9.2.39 Capillary

**Scope:** Equation System

Capillary Fluxbp For Solid [{of|species|subindex} Species |{in|material_phase} MaterialPhase ][@|at|for|in|on|over} Mesh Extent Name = Capillary [ Using Data Specification Data Spec Name ]

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**Summary**

Transient traction solid flux

### 9.2.40 Wall Function

**Scope:** Equation System

Wall Function Disting For Cvfem_Turbulence_Dissipation_Rate [{of|species|subindex} Species |{in|material_phase} MaterialPhase ][@|at|for|in|on|over} Mesh Extent Name = Wall_Function [ Using Data Specification Data Spec Name ] [ Wall_Friction_Factor = Wall_Friction_Factor | Wall_Velocity_X = Wall_Velocity_X | Wall_Velocity_Y = Wall_Velocity_Y | Wall_Velocity_Z = Wall_Velocity_Z ]

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**Summary**

CVFEM wall function turbulence dissipation rate distinguishing condition
9.2.41 Simple Interp

Scope: Equation System

Simple Interp Flux For Cvfem_Tke_Edge_Gradient_Projection [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ][@at|for|in|on|over} Mesh Extent Name = Simple_Interp [ Using Data Specification Data Spec Name ][ ]

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Summary: CVFEM edge-based projection simple interpolation flux BC

9.2.42 Kinematic

Scope: Equation System

Kinematic Disting For Solid_Y [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ][@at|for|in|on|over} Mesh Extent Name = Kinematic [ Using Data Specification Data Spec Name ][ ] V0 = v0 |Shared_Normal = shared_normal |No_Mesh_Movement = no_mesh_movement ]

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Summary: Kinematic model for SOLID_y distinguishing condition

9.2.43 Wall Function

Scope: Equation System

Wall Function Flux For Hfem_Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ][@at|for|in|on|over} Mesh Extent Name = Wall_Function [ Using Data Specification Data Spec Name ][ Scale_Factor = scale_factor |Wall_Friction_Factor = wall_friction_factor |Wall_Velocity_X = wall_velocity_x |Wall_Velocity_Y = wall_velocity_y |Wall_Velocity_Z = wall_velocity_z ]
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Summary: HFEM momentum symmetry flow flux BC

### 9.2.44 Capillary

**Scope:** Equation System

\[
\text{Capillary Fluxbp For Cvfem_Momentum } \{ \text{of|species|subindex} \text{Species} | \{\text{in|material_phase} \text{MaterialPhase} \} \{\text{at|for|in|on|over} \text{Mesh Extent Name} = \text{Capillary} \} [ \text{Using Data Specification Data Spec Name } ] [ ]
\]

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Summary: Momentum capillary flux BP BC

### 9.2.45 Ls Capillary

**Scope:** Equation System

\[
\text{Ls Capillary Fluxbp For Cvfem_Momentum } \{ \text{of|species|subindex} \text{Species} | \{\text{in|material_phase} \text{MaterialPhase} \} \{\text{at|for|in|on|over} \text{Mesh Extent Name} = \text{Ls_Capillary} \} [ \text{Using Data Specification Data Spec Name } ] [ ]
\]

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Summary: Momentum capillary flux BP BC using level set normal

### 9.2.46 Porous Robin Interface

**Scope:** Equation System

\[
\text{Porous Robin Interface Disting For Cvfem_Momentum } \{ \text{of|species|subindex} \text{Species} | \{\text{in|material_phase} \text{MaterialPhase} \} \{\text{at|for|in|on|over} \text{Mesh Extent Name} = \text{Porous_Robin_Interface} \}
\]
Summary: No slip model for momentum distinguishing condition

### 9.2.47 Porous Robin One Region

**Scope:** Equation System

Porous Robin One Region Disting For Cvfem_Momentum [ {of|species|subindex} Species | {in |material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Porous_Robin_One_Region [ Using Data Specification Data Spec Name ][ Alpha = alpha ]

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Summary: No slip model for momentum distinguishing condition

### 9.2.48 No Slip

**Scope:** Equation System

No Slip Disting For Cvfem_Momentum [ {of|species|subindex} Species | {in |material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = No_Slip [ Using Data Specification Data Spec Name ][ ]

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Summary: No slip model for momentum distinguishing condition

### 9.2.49 Mass Flux

**Scope:** Equation System
Mass Flux Flux For Cvfem_Mixture_Fraction [ \{of\species{species}{subindex}\} \Species \{in\material_phase{MaterialPhase}\}@\at\for\in\on\over\ Mesh Extent Name = Mass_Flux [ Using Data Specification Data Spec Name ][ Total_Mdot = total_mdot | Inflow_Phi = inflow_phi ]

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9.2.50 Open Adv Flow
Scope: Equation System

Open Adv Flow Flux For Cvfem_Mixture_Fraction [ \{of\species{species}{subindex}\} \Species \{in\material_phase{MaterialPhase}\}@\at\for\in\on\over\ Mesh Extent Name = Open_Adv_Flow [ Using Data Specification Data Spec Name ][ Pressure = pressure | Total_Pressure = total_pressure | Far_Field_Entrainment_Value = far_field_entrainment_value ]

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9.2.51 Simple Dp Interp
Scope: Equation System

Simple Dp Interp Flux For Hfem_Velocity_Pressure_Projection [ \{of\species{species}{subindex}\} \Species \{in\material_phase{MaterialPhase}\}@\at\for\in\on\over\ Mesh Extent Name = Simple_Dp_Interp [ Using Data Specification Data Spec Name ][ ]

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Summary
Simple DP interpolation model for HFEM velocity pressure projection flux

9.2.52 Open Adv Flow
Scope: Equation System

Open Adv Flow Flux For Hfem_Turbulent_Kinetic_Energy [ \{of\species{species}{subindex}\} \Species \{in\material_phase{MaterialPhase}\}@\at\for\in\on\over\ Mesh Extent Name = Open_Adv_Flow
[ Using Data Specification Data Spec Name ][ Pressure = pressure | Total_Pressure = total_pressure | Far_Field_Entrainment_Value = far_field_entrainment_value ]

<table>
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Summary: HFEM turbulent kinetic energy open advective flow flux BC

### 9.2.53 Calore User Sub

**Scope:** Equation System

Calore User Sub Flux For Spherical_Harmonic [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Calore_User_Sub [ Using Data Specification Data Spec Name ][ Name = name | Type = type | Multiplier = multiplier | Material_Data_Block = material_data_block | Data = data | Scaling_Field = scaling_field ]

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Summary: Values from a Calore user subroutine

### 9.2.54 Mixed

**Scope:** Equation System

Mixed Flux For Spherical_Harmonic [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Mixed [ Using Data Specification Data Spec Name ][ ]

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Summary: Applies the MIXED scalar flux from the spherical harmonic equations.
### 9.2.55 Capillary

**Scope:** Equation System

Capillary Flux For Momentum \[ \{\text{of|species|subindex}\} \text{Species}\ |\{\text{in|material_phase}\} \text{MaterialPhase} \] \{@at\text{for|in|on|over}\} \text{Mesh Extent Name} = \text{Capillary} [ \text{Using Data Specification Data Spec Name } ]

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**Summary**  
LS capillary momentum flux

### 9.2.56 Capillary Gap

**Scope:** Equation System

Capillary Gap Flux For Momentum \[ \{\text{of|species|subindex}\} \text{Species}\ |\{\text{in|material_phase}\} \text{MaterialPhase} \] \{@at\text{for|in|on|over}\} \text{Mesh Extent Name} = \text{Capillary Gap} [ \text{Using Data Specification Data Spec Name } ] \[ v_w = v_w \ | G = g \ | \Theta = \theta \]

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**Summary**  
Capillary gap momentum flux

### 9.2.57 Slip

**Scope:** Equation System

Slip Flux For Momentum \[ \{\text{of|species|subindex}\} \text{Species}\ |\{\text{in|material_phase}\} \text{MaterialPhase} \] \{@at\text{for|in|on|over}\} \text{Mesh Extent Name} = \text{Slip} [ \text{Using Data Specification Data Spec Name } ] \[ v_s_x = v_s_x \ | v_s_y = v_s_y \ | v_s_z = v_s_z \ | \beta = \beta \]

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</table>
9.2.58  **Ls Capillary**

**Scope:**  Equation System

Ls Capillary Flux For Momentum [ {of|species|subindex} *Species* |{in|material_phase} *MaterialPhase* ]{at|for|in|on|over} *Mesh Extent Name* = Ls_Capillary [ Using Data Specification *Data Spec Name* ]

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Summary  LS capillary momentum flux

9.2.59  **Open Flow Darcy**

**Scope:**  Equation System

Open Flow Darcy Flux For Momentum [ {of|species|subindex} *Species* |{in|material_phase} *MaterialPhase* ]{at|for|in|on|over} *Mesh Extent Name* = Open_Flow_Darcy [ Using Data Specification *Data Spec Name* ] [ Pressure = *pressure* |Total_Pressure = *total_pressure* ]

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Summary  Open flow momentum flux

9.2.60  **Symmetry Flow**

**Scope:**  Equation System

Symmetry Flow Flux For Momentum [ {of|species|subindex} *Species* |{in|material_phase} *MaterialPhase* ]{at|for|in|on|over} *Mesh Extent Name* = Symmetry_Flow [ Using Data Specification *Data Spec Name* ]

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Summary  Symmetry flow momentum flux
9.2.61  Constant Traction

Scope:  Equation System

Constant Traction Flux For Mesh [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ]@at|for|in|on|over} Mesh Extent Name = Constant_Traction [ Using Data Specification Data Spec Name ][ X = x | Y = y | Z = z ]

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</table>

Summary: Constant traction mesh flux

9.2.62  Transient Traction

Scope:  Equation System

Transient Traction Flux For Mesh [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ]@at|for|in|on|over} Mesh Extent Name = Transient_Traction [ Using Data Specification Data Spec Name ][ A_X = a_x | A_Y = a_y | A_Z = a_z | B_X = b_x | B_Y = b_y | B_Z = b_z ]

<table>
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Summary: Transient traction mesh flux

9.2.63  Electric Traction

Scope:  Equation System

Electric Traction Flux For Mesh [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ]@at|for|in|on|over} Mesh Extent Name = Electric_Traction [ Using Data Specification Data Spec Name ][ Sign = sign ]
### Pressure User Function

**Scope:** Equation System

Pressure User Function Flux For Mesh [{of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Pressure_User_Function [ Using Data Specification Data Spec Name ] [ Name = name | X = x | X_Multiplier = x_multiplier | Multiplier = multiplier | Toggle = toggle ]

### Kinematic

**Scope:** Equation System

Kinematic Rotated For Mesh [{of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Kinematic [ Using Data Specification Data Spec Name ] [ V0 = v0 | Shared_Normal = shared_normal | No_Mesh_Movement = no_mesh_movement ]

---

**Summary**
- Electric traction mesh flux
- Pressure user function mesh flux
- Kinematic model for rotated mesh boundary condition
9.2.66  Shear Free
Scope:  Equation System

Shear Free Flux For Mesh [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Shear_Free [ Using Data Specification Data Spec Name ]

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Summary  Shear-free traction mesh flux

9.2.67  Recession
Scope:  Equation System

Recession Rotated For Mesh [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Recession [ Using Data Specification Data Spec Name ][ P_Ref = p_ref | K = k | K_0 = k_0 | N = n | Pressure = pressure ]

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Summary  Recession model for rotated mesh boundary condition

9.2.68  Simple Interp Vector
Scope:  Equation System

Simple Interp Vector Flux For Cvfem_Lumped_Div_Projection [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Simple_Interp_Vector [ Using Data Specification Data Spec Name ][ Source = source ]

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Summary  CVFEM lumped divergence projection simple interpolation vector flux BC
9.2.69 Open
Scope: Equation System

Open Flux For Mass_Balance [{of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Open [ Using Data Specification Data Spec Name ] [ Pressure = pressure | Entrained_Value = entrained_value ]

<table>
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</table>

9.2.70 Fluid Robin Coupled One Region
Scope: Equation System

Fluid Robin Coupled One Region Flux For Mass_Balance [{of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Fluid_Robin_Coupled_One_Region [ Using Data Specification Data Spec Name ] [ Coeff_Scaling = coeff_scaling ]

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Summary Boundary condition for coupling of continuity to a free fluid in the same region.

9.2.71 Simple Interp
Scope: Equation System

Simple Interp Flux For Cvfem_Projection [{of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Simple_Interp [ Using Data Specification Data Spec Name ]

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</table>

Summary CVFEM projection simple interpolation flux BC

9.2.72 Open
Scope: Equation System

436
Open Flux For Cvfem_Projection [{of|species|subindex} Species |{in|material_phase} MaterialPhase ]@{at|for|in|on|over} Mesh Extent Name = Open [ Using Data Specification Data Spec Name ] [ Total_Pressure = total_pressure | Pressure = pressure ]

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Summary: CVFEM projection open flux BC

9.2.73 From Cht Temperature

Scope: Equation System

From Cht Temperature Disting For Porous_Enthalpy [{of|species|subindex} Species |{in|material_phase} MaterialPhase ]@{at|for|in|on|over} Mesh Extent Name = From_Cht_Temperature [ Using Data Specification Data Spec Name ] [ Interface_Name = Interface_Name | Twall_Field = Twall_Field | Tref_Field = Tref_Field | H_Field = H_Field ]

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Summary: Porous enthalpy distinguishing condition from conjugate heat transfer temperature

9.2.74 Kinematic

Scope: Equation System

Kinematic Rotated For Cvfem_Momentum [{of|species|subindex} Species |{in|material_phase} MaterialPhase ]@{at|for|in|on|over} Mesh Extent Name = Kinematic [ Using Data Specification Data Spec Name ] [ VO = v0 | Shared_Normal = shared_normal | No_Mesh_Movement = no_mesh_movement ]

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9.2.75 Symmetry Flow

Scope: Equation System

Symmetry Flow Flux For Cvem_Momentum [{of|species|subindex} Species |{in|material_phase} MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Symmetry_Flow [ Using Data Specification Data Spec Name ]

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Summary CVFEM momentum symmetry flow flux BC

9.2.76 Open Adv Flow

Scope: Equation System

Open Adv Flow Flux For Cvem_Momentum [{of|species|subindex} Species |{in|material_phase} MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Open_Adv_Flow [ Using Data Specification Data Spec Name ] [ Pressure = pressure | Total_Pressure = total_pressure | Coflow_Entrainment_Value_X = coflow_entrainment_value_x | Coflow_Entrainment_Value_Y = coflow_entrainment_value_y | Coflow_Entrainment_Value_Z = coflow_entrainment_value_z ]

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Summary CVFEM momentum open advective flow flux BC

9.2.77 Interface

Scope: Equation System

Interface Flux For Cvem_Momentum [{of|species|subindex} Species |{in|material_phase} MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Interface [ Using Data Specification Data Spec Name ]
### 9.2.78 Wall Function

**Scope:** Equation System

Wall Function Flux For Cvfem_Momentum [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Wall_Function [ Using Data Specification Data Spec Name ] [ Wall_Friction_Factor = wall_friction_factor | Wall_Velocity_X = wall_velocity_x | Wall_Velocity_Y = wall_velocity_y | Wall_Velocity_Z = wall_velocity_z ]

<table>
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**Summary**
CVFEM momentum wall function flux BC

### 9.2.79 Convective Outflow

**Scope:** Equation System

Convective Outflow Flux For Mass_Balance [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Convective_Outflow [ Using Data Specification Data Spec Name ] [ Ref_Frac = ref_frac ]

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**Summary**
CVFEM momentum wall function flux BC

### 9.2.80 Open Adv Flow

**Scope:** Equation System

Open Adv Flow Flux For Cvfem_Specific_Dissipation_Rate [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Open_Adv_Flow [ Using Data Specification Data Spec Name ] [ Pressure = pressure | Total_Pressure = total_pressure | Far_Field_Entrainment_Value = far_field_entrainment_value ]

439
### 9.2.81 Mass Flux

**Scope:** Equation System

\[
\text{Mass Flux Flux For Cvfem\_Specific\_Dissipation\_Rate} \ [\{\text{of|species|subindex}\} \text{Species} \ |\{\text{in|material\_phase}\} \text{MaterialPhase} \} @\{\text{at|for|in|on|over}\} \text{Mesh Extent Name} = \text{Mass Flux} \ [\text{Using Data Specification Data Spec Name}] [\text{Total_Mdot} = \text{total_mdot} \ |\text{Inflow_Phi} = \text{inflow\_phi}]
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### 9.2.82 Porous Robin One Region

**Scope:** Equation System

\[
\text{Porous Robin One Region Flux For Cvfem\_Continuity} \ [\{\text{of|species|subindex}\} \text{Species} \ |\{\text{in|material\_phase}\} \text{MaterialPhase} \} @\{\text{at|for|in|on|over}\} \text{Mesh Extent Name} = \text{Porous Robin One Region} \ [\text{Using Data Specification Data Spec Name}] [\ ]
\]

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**Summary**

Boundary condition for loose coupling of continuity to a free fluid region

### 9.2.83 Porous Robin Averaged Coeff

**Scope:** Equation System

\[
\text{Porous Robin Averaged Coeff Flux For Cvfem\_Continuity} \ [\{\text{of|species|subindex}\} \text{Species} \ |\{\text{in|material\_phase}\} \text{MaterialPhase} \} @\{\text{at|for|in|on|over}\} \text{Mesh Extent Name} = \text{Porous Robin Averaged Coeff} \ [\text{Using Data Specification Data Spec Name}] [\text{P\_Field} = \text{P\_Field} \ |\text{Massflux\_Field} = \text{Massflux\_Field} \ |\text{Oppositecoeff\_Field} = \text{Oppositecoeff\_Field}]
\]

440
### Parameter | Value | Default
---|---|---
Species | string | undefined
MaterialPhase | string | undefined
Mesh Extent Name | string | undefined
Data Spec Name | string | undefined
P_Field | "string" | undefined
MassFlux_Field | "string" | undefined
OppositeCoeff_Field | "string" | undefined

**Summary**  
Boundary condition for loose coupling of continuity to a free fluid region

#### 9.2.84 Mass Flux

**Scope:** Equation System

Mass Flux Flux For CvFem_Continuity [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@[at|for|in|on|over} Mesh Extent Name = Mass_Flux [ Using Data Specification Data Spec Name ][ Total_Mdot = total_mdot ]

**Parameter** | **Value** | **Default**
---|---|---
Species | string | undefined
MaterialPhase | string | undefined
Mesh Extent Name | string | undefined
Data Spec Name | string | undefined
total_mdot | real | undefined

**Summary**  
Mass flux CVFEM continuity flux

#### 9.2.85 Wall Function

**Scope:** Equation System

Wall Function Disting For Hfem_Turbulent_Kinetic_Energy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@[at|for|in|on|over} Mesh Extent Name = Wall_Function [ Using Data Specification Data Spec Name ][ Scale_Factor = scale_factor | Wall_Friction_Factor = Wall_Friction_Factor | Wall_Velocity_X = Wall_Velocity_X | Wall_Velocity_Y = Wall_Velocity_Y | Wall_Velocity_Z = Wall_Velocity_Z ]

**Parameter** | **Value** | **Default**
---|---|---
Species | string | undefined
MaterialPhase | string | undefined
Mesh Extent Name | string | undefined
Data Spec Name | string | undefined
scale_factor | real | undefined
Wall_Friction_Factor | real | undefined
Wall_Velocity_X | real | undefined
Wall_Velocity_Y | real | undefined
Wall_Velocity_Z | real | undefined

**Summary**  
HFEM wall function turbulent kinetic energy distinguishing condition
9.2.86  Open Adv Flow
Scope:  Equation System

Open Adv Flow Flux For Hfem_Specific_Dissipation_Rate [{of|species|subindex} Species | {in|material_phase} MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Open_Adv_Flow [ Using Data Specification Data Spec Name ][ Pressure = pressure |Total_Pressure = total_pressure |Far_Field_Entrainment_Value = far_field_entrainment_value ]

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Summary  HFEM specific dissipation rate open advective flow flux BC

9.2.87  Reconstructed Curvature
Scope:  Equation System

Reconstructed Curvature Flux For Momentum [{of|species|subindex} Species | {in|material_phase} MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Reconstructed_Curvature [ Using Data Specification Data Spec Name ][ Ls_Size = ls_size |Fixed_Curvature = fixed_curvature ]

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Summary  Reconstructed curvature momentum flux

9.2.88  Node Normal Capillary
Scope:  Equation System

Node Normal Capillary Flux For Momentum [{of|species|subindex} Species | {in|material_phase} MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Node_Normal_Capillary [ Using Data Specification Data Spec Name ]

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9.2.89 Generalized Nat Conv

Scope: Equation System

Generalized Nat Conv No_Coverage_Flux For Energy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Generalized_Nat_Conv [ Using Data Specification Data Spec Name ][ Multiplier = multiplier ]

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Summary Generalized natural convection energy flux

9.2.90 Vapor Recoil Pressure

Scope: Equation System

Vapor Recoil Pressure Flux For Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Vapor_Recoil_Pressure [ Using Data Specification Data Spec Name ][ Tboil = tboil ]

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Summary Vapor recoil pressure momentum flux

9.2.91 Porous Robin Coupled

Scope: Equation System

Porous Robin Coupled Flux For Cvem_Energy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Porous_Robin_Coupled [ Using Data Specification Data Spec Name ][ Data = data |Enthalpy_Field = enthalpy_field |Enthalpy_Diffusive_Flux = enthalpy_diffusive_flux_field |Averaged_Coeff_Field = averaged_coeff_field ]
Parameter | Value | Default
---|---|---
Species | string | undefined
MaterialPhase | string | undefined
Mesh Extent Name | string | undefined
Data Spec Name | string | undefined
data | "string" | undefined
enthalpy_field | "string" | undefined
enthalpy_diffusive_flux_field | "string" | undefined
averaged_coeff_field | "string" | undefined

Summary: Boundary condition for loose coupling of enthalpy to a porous region

### 9.2.92 Generalized Nat Conv

**Scope:** Equation System

Generalized Nat Conv Flux For Enthalpy [:Species :MaterialPhase :Mesh Extent Name = Generalized_Nat_Conv [ Using Data Specification Data Spec Name ][ Multiplier = multiplier ]

Parameter | Value | Default
---|---|---
Species | string | undefined
MaterialPhase | string | undefined
Mesh Extent Name | string | undefined
Data Spec Name | string | undefined
multiplier | real | undefined

Summary: Generalized natural convection enthalpy flux

### 9.2.93 Generalized Rad

**Scope:** Equation System

Generalized Rad Flux For Enthalpy [:Species :MaterialPhase :Mesh Extent Name = Generalized_Rad [ Using Data Specification Data Spec Name ][ Multiplier = multiplier ]

Parameter | Value | Default
---|---|---
Species | string | undefined
MaterialPhase | string | undefined
Mesh Extent Name | string | undefined
Data Spec Name | string | undefined
multiplier | real | undefined

Summary: Radiative energy flux, possibly defined with different models for emissivity, form factor and irradiation

### 9.2.94 Low Reynolds

**Scope:** Equation System
Low Reynolds Distinct For Cvem_Specific_Dissipation_Rate [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@at|for|in|on|over} Mesh Extent Name = Low_Reynolds [ Using Data Specification Data Spec Name ][ ]

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9.2.95  Ls Capillary
Scope:  Equation System

Ls Capillary Fluxbp For Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@at|for|in|on|over} Mesh Extent Name = Ls_Capillary [ Using Data Specification Data Spec Name ][ ]

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Summary  Momentum capillary flux BP BC using level set normal

9.2.96  Capillary Stabilization
Scope:  Equation System

Capillary Stabilization Fluxbp For Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@at|for|in|on|over} Mesh Extent Name = Capillary_Stabilization [ Using Data Specification Data Spec Name ][ Mu0 = mu0 |Multiplier = multiplier ]

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Summary  Momentum capillary stabilization flux BP BC

9.2.97  Ls Capillary Stabilization
Scope:  Equation System

Ls Capillary Stabilization Fluxbp For Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@at|for|in|on|over} Mesh Extent Name = Ls_Capillary_Stabilization [ Using Data Specification Data Spec Name ][ Mu0 = mu0 |Multiplier = multiplier ]
9.2.98 Normal Capillary Stabilization

Scope: Equation System

Normal Capillary Stabilization Fluxbp For Momentum [{of|species|subindex} Species |{in |material_phase} MaterialPhase ][@|at|for|in|on|over] Mesh Extent Name = Normal_Capillary_Stabilization [ Using Data Specification Data Spec Name ][ Mu0 = mu0 |Multiplier = multiplier ]

9.2.99 Free Open Flow

Scope: Equation System

Free Open Flow Flux For Momentum [{of|species|subindex} Species |{in|material_phase} MaterialPhase ][@|at|for|in|on|over] Mesh Extent Name = Free_Open_Flow [ Using Data Specification Data Spec Name ][ Pressure = pressure |Pressure User Sub PRESSURE User Sub ]

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Summary: Open flow for momentum; enforce normal gradient to be zero

9.2.101 Pressure

Scope: Equation System

Pressure Flux For Momentum [ of species subindex ] Species [ in material_phase ] MaterialPhase [ at for in on over ] Mesh Extent Name = Pressure [ Using Data Specification Data Spec Name ] [ Use_Bulk_Node = use_bulk_node | P = p | C_T = c_t | C_X = c_x | C_Y = c_y | C_Z = c_z | C_Cos_W = c_cos_w | C_Sin_W = c_sin_w | C1_Cos = c1_cos | C1_Sin = c1_sin ]

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Summary: Pressure momentum flux

9.2.102 Transient Traction

Scope: Equation System

Transient Traction Flux For Momentum [ of species subindex ] Species [ in material_phase ] MaterialPhase [ at for in on over ] Mesh Extent Name = Transient_Traction [ Using Data Specification Data Spec Name ] [ A_X = a_x | A_Y = a_y | A_Z = a_z | B_X = b_x | B_Y = b_y | B_Z = b_z ]
### 9.2.103 Extrapolated Pressure

**Scope:** Equation System

Extrapolated Pressure Flux For Continuity [{of|species|subindex} Species |{in|material_phase} MaterialPhase ][@|at|for|in|on|over] Mesh Extent Name = Extrapolated_Pressure [ Using Data Specification Data Spec Name ]

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**Summary**  
Extrapolated pressure for use with artificial compressibility methods

### 9.2.104 Pspg

**Scope:** Equation System

Pspg Flux For Continuity [{of|species|subindex} Species |{in|material_phase} MaterialPhase ][@|at|for|in|on|over] Mesh Extent Name = Pspg [ Using Data Specification Data Spec Name ] [ TauSurfaceFactor = TauSurfaceFactor | TauSurfaceJacobianFactor = TauSurfaceJacobianFactor ]

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**Summary**  
Continuity PSPG flux BC

### 9.2.105 Gaussian Spot Weld

**Scope:** Equation System

Gaussian Spot Weld

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9.2.106 Laser

Scope: Equation System

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### 9.2.107 Circle Weld

**Scope:** Equation System

Circle Weld Flux For Energy [ {of|species|subindex} Species {in|material_phase} MaterialPhase ] {at|for|in|on|over} Mesh Extent Name = Circle_Weld [ Using Data Specification Data Spec Name ] { Power_Output = power_output | Flux_Output = flux_output | Toggle = toggle | O_X = o_x | O_Y = o_y | O_Z = o_z | Ro_X = ro_x | Ro_Y = ro_y | Ro_Z = ro_z | A_X = a_x | A_Y = a_y | A_Z = a_z | W = w | Flux = flux | R = r | Max_Degrees = max_degrees | T_On = t_on | T_Off = t_off | Normal_Tolerance = normal_tolerance ]

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450
### 9.2.108 Distribution Factor

**Scope:** Equation System

Distribution Factor Flux For Energy \([ \{\text{of\ species|subindex}\} \ Species \ |\{\text{in\ material\ phase}\} \ MaterialPhase \}@|\text{at}\ |\text{for}\ |\text{in}\ |\text{on}\ |\text{over}\ \ Mesh\ Extent\ Name\ =\ Distribution\_Factor \ [\ \text{Using}\ Data\ Specification\ Data\ Spec\ Name\ ]\ [\ \text{Power\ Output}\ =\ power\_output\ |\ Flux\ Output\ =\ flux\_output\ |\ Toggle\ =\ toggle\ |\ Attribute\_Name\ =\ attribute\_name\ |\ Multiplier\ =\ multiplier\ ]\]

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### 9.2.109 Enclosure Radiation

**Scope:** Equation System

Enclosure Radiation Flux For Energy \([ \{\text{of\ species|subindex}\} \ Species \ |\{\text{in\ material\ phase}\} \ MaterialPhase \}@|\text{at}\ |\text{for}\ |\text{in}\ |\text{on}\ |\text{over}\ \ Mesh\ Extent\ Name\ =\ Enclosure\_Radiation \ [\ \text{Using}\ Data\ Specification\ Data\ Spec\ Name\ ]\ [\ \text{Power\ Output}\ =\ power\_output\ |\ Flux\ Output\ =\ flux\_output\ |\ Toggle\ =\ toggle\ |\ Ndof\ =\ ndof\ |\ Multiplier\ =\ multiplier\ |\ Enclosure\ =\ enclosure\ ]\]

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### 9.2.110 Gaussian Line Weld

**Scope:** Equation System

Gaussian Line Weld Flux For Energy \([ \{\text{of\ species|subindex}\} \ Species \ |\{\text{in\ material\ phase}\} \ MaterialPhase \}@|\text{at}\ |\text{for}\ |\text{in}\ |\text{on}\ |\text{over}\ \ Mesh\ Extent\ Name\ =\ Gaussian\_Line\_Weld \ [\ \text{Using}\ Data\ Specification\ Data\ Spec\ Name\ ]\ [\ \text{Power\ Output}\ =\ power\_output\ |\ Flux\ Output\ =\ flux\_output\ |\ Toggle\ =\ toggle\ |\ Src\_X\ =\ src\_x\ |\ Dir\_X\ =\ dir\_x\ |\ Vel\_X\ =\ vel\_x\ |\ Src\_Y\ =\ src\_y\ |\ Dir\_Y\ =\ dir\_y\ |\ Vel\_Y\ =\ vel\_y\ |\ Src\_Z\ =\ src\_z\ |\ Dir\_Z\ =\ dir\_z\ |\ Vel\_Z\ =\ vel\_z\ |\ R\ =\ r\ |\ T\_On\ =\ t\_on\ |\ T\_Off\ =\ t\_off\ |\ Flux\ =\ flux\ |\ R\_Eff\ =\ r\_eff\ |\ Alpha\ =\ alpha\ |\ Depth\_Enhanced\_Absorption\_Data\ =\ depth\_enhanced\_absorption\_data\ |\ Compute\_Visibility\_Field\ =\ compute\_visibility\_field\ ]\]

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### 9.2.111 From Cht Temperature

**Scope:** Equation System

From Cht Temperature Disting For Energy \{ of \{ species \{ subindex \} Species \{ in \{ material_phase \} MaterialPhase \} \} \{ at \| for \| in \| on \| over \} Mesh Extent Name = From_Cht_Temperature \[ Using Data Specification Data Spec Name \] \[ Interface_Name = Interface_Name \| Twall_Field = Twall_Field \| Tref_Field = Tref_Field \| H_Field = H_Field \]

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**Summary**

Energy distinguishing condition from conjugate heat transfer temperature

### 9.2.112 Calore User Sub

**Scope:** Equation System

Calore User Sub Flux For Energy \{ of \{ species \{ subindex \} Species \{ in \{ material_phase \} MaterialPhase \} \} \{ at \| for \| in \| on \| over \} Mesh Extent Name = Calore_User_Sub \[ Using Data Specification Data \]

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9.2.113 Capillary Force

Scope: Equation System

Capillary Force Flux For Lumped_Div_Projection [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Capillary_Force [ Using Data Specification Data Spec Name ][ Theta = theta ]

Summary: Capillary force model for lumped div projection flux

9.2.114 Recession

Scope: Equation System

Recession Disting For Solid_Y [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Recession [ Using Data Specification Data Spec Name ][ P_Ref = p_ref | K = k | K_0 = k_0 | N = n | Pressure = pressure ]
### Summary
Recession model for SOLID_x distinguishing condition

#### 9.2.115 Calore User Sub

**Scope:** Equation System

Calore User Sub Flux For Current [{of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@at|for|in|on|over} Mesh Extent Name = Calore_User_Sub [ Using Data Specification Data Spec Name ][ Toggle = toggle |Name = name |Type = type |Multiplier = multiplier |Material_Data_Block = material_data_block |Data = data |Scaling_Field = scaling_field ]

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**Summary**
Values from a Calore user subroutine

#### 9.2.116 Constant With Cutoff Voltage

**Scope:** Equation System

Constant With Cutoff Voltage Flux For Current [{of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@at|for|in|on|over} Mesh Extent Name = Constant_With_Cutoff_Voltage [ Using Data Specification Data Spec Name ][ Toggle = toggle |Value = value |Cutoff_Value = cutoff_value ]

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**Summary**

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Summary: Apply a constant current until a cutoff voltage is crossed anywhere on the surface. After that apply 0 current.

### 9.2.117 Resistive Load

**Scope:** Equation System

Resistive Load Flux For Current [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ][@|at|for|in|on|over] Mesh Extent Name = Resistive_Load [ Using Data Specification Data Spec Name ] [ Toggle = toggle | Area = Area | Reference Voltage = Reference Voltage ]

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Summary: Model a boundary attached to a resistive load. The local applied current density is \( V / (R A) \) where \( V \) is the local voltage, \( R \) is the specified resistance, and \( A \) is the area of the surface the BC is applied to.

### 9.2.118 Simple Interp

**Scope:** Equation System

Simple Interp Flux For Cvfem_Lumped_Projection [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ][@|at|for|in|on|over] Mesh Extent Name = Simple_Interp [ Using Data Specification Data Spec Name ]

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Summary: CVFEM lumped projection simple interpolation flux BC

### 9.2.119 Open

**Scope:** Equation System
Open Flux For Cvfem_Lumped_Projection [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@}|at|for|in|on|over} Mesh Extent Name = Open [ Using Data Specification Data Spec Name ][ Total_Pressure = total_pressure |Pressure = pressure ]

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Summary: CVFEM lumped projection open flux BC

9.2.120 Cht Flux

Scope: Equation System

Cht Flux Flux For Energy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@}|at|for|in|on|over} Mesh Extent Name = Cht_Flux [ Using Data Specification Data Spec Name ][ Power_Output = power_output |Flux_Output = flux_output |Toggle = toggle |Too_Field = Too_Field |
|H_Field = H_Field ]

<table>
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Summary: Applies H(T-Too) flux for conjugate heat transfer boundary. Optional parameters are "Too_Field" and "H_Field", which default to "nodal_cht_Too" and "nodal_cht_H" respectively.

9.2.121 Cht Robin

Scope: Equation System

Cht Robin Flux For Energy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@}|at|for|in|on|over} Mesh Extent Name = Cht_Robin [ Using Data Specification Data Spec Name ][ Power_Output = power_output |Flux_Output = flux_output |Toggle = toggle |Temperature_Field = temperature_field |
|Heat_Flux_Field = heat_flux_field |Coeff_Scaling = coeff_scaling ]

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### 9.2.122 Generalized Nat Conv

**Scope:** Equation System

Generalized Nat Conv Flux For Energy \{of\|species\|subindex\} \{in\|material\_phase\} MaterialPhase \{at\|for\|in\|on\|over\} Mesh Extent Name = Generalized_Nat_Conv \{ Using Data Specification Data Spec Name \} [ Power_Output = power_output \| Flux_Output = flux_output \| Toggle = toggle \| Multiplier = multiplier ]

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**Summary** Generalized natural convection energy flux

### 9.2.123 Fortran Conv

**Scope:** Equation System

Fortran Conv Flux For Energy \{of\|species\|subindex\} Species \{in\|material\_phase\} MaterialPhase \{at\|for\|in\|on\|over\} Mesh Extent Name = Fortran_Conv \{ Using Data Specification Data Spec Name \} [ Power_Output = power_output \| Flux_Output = flux_output \| Toggle = toggle \| Multiplier = multiplier \| Htc_Sub_Name = htc_sub_name \| Tref_Sub_Name = tref_sub_name \| Real_Data = real_data \| Int_Data = int_data \| Resource_Name = resource_name \| Data = data ]
### 9.2.124 User Vector Field

**Scope:** Equation System

User Vector Field Flux For Energy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = User_Vector_Field [ Using Data Specification Data Spec Name ]| Power_Output = power_output |Flux_Output = flux_output |Toggle = toggle |Name = name |Multiplier = multiplier |Scaling = scaling ]

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**Summary**  
User vector field energy flux

### 9.2.125 Scaled Nat Conv

**Scope:** Equation System

Scaled Nat Conv Flux For Energy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Scaled_Nat_Conv [ Using Data Specification Data Spec Name ]| Power_Output = power_output |Flux_Output = flux_output |Toggle = toggle |Multiplier = multiplier |Scaling_Field = scaling_field |Scaling = scaling |Global_Var = global_var ]

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**Summary**  
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Summary: Scaled natural convection energy flux

9.2.126 Fortran
Scope: Equation System

Fortran Flux For Energy [ \{of\} \{species\} \{subindex\} \{in\} \{material\} \{phase\} || \{of\} \{in\} \{on\} \{over\} \{Mesh\} \{Extent\} \{Name\} = Fortran [ Using Data Specification \{Data\} \{Spec\} \{Name\} ] [ Power_Output = power_output | Flux_Output = flux_output | Toggle = toggle | Multiplier = multiplier | Sub_Name = sub_name | Real_Data = real_data | Int_Data = int_data | Resource_Name = resource_name | Data = data ]

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Summary: Fortran subroutine energy flux

9.2.127 Face Field Scalar
Scope: Equation System

Face Field Scalar Flux For Energy [ \{of\} \{species\} \{subindex\} \{in\} \{material\} \{phase\} || \{of\} \{in\} \{on\} \{over\} \{Mesh\} \{Extent\} \{Name\} = Face_Field_Scalar [ Using Data Specification \{Data\} \{Spec\} \{Name\} ] [ Power_Output = power_output | Flux_Output = flux_output | Toggle = toggle | Name = name | Multiplier = multiplier | Advective_Bar = advective_bar | Command_Block_Name = command_block_name ]
Parameter | Value | Default
---|---|---
Species | string | undefined
MaterialPhase | string | undefined
Mesh Extent Name | string | undefined
Data Spec Name | string | undefined
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toggle | "string" | undefined
name | "string" | undefined
multiplier | real | undefined
advective_bar | "string" | undefined
command_block_name | "string" | undefined

Summary | FACE or EDGE Field energy flux

9.2.128  Fortran Rad
Scope: Equation System

Fortran Rad Flux For Energy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Fortran_Rad [ Using Data Specification Data Spec Name ] Power_Output = power_output | Flux_Output = flux_output | Toggle = toggle | Multiplier = multiplier | Emissivity_Sub_Name = emissivity_sub_name | Tref_Sub_Name = tref_sub_name | Form_Factor_Sub_Name = form_factor_sub_name | Real_Data = real_data | Int_Data = int_data | Resource_Name = resource_name | Data = data |

Parameter | Value | Default
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Mesh Extent Name | string | undefined
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real_data | "string" | undefined
int_data | "string" | undefined
resource_name | "string" | undefined
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Summary | Fortran subroutine radiation energy flux

9.2.129  Masked Nat Conv
Scope: Equation System

Masked Nat Conv Flux For Energy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Masked_Nat_Conv [ Using Data Specification Data
Spec Name ][ Power_Output = power_output | Flux_Output = flux_output | Toggle = toggle | Multiplier = multiplier | Masking_Field = masking_field | Scaling = scaling | Threshold = threshold ]

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Summary: Masked natural convection energy flux

9.2.130  Rad

Scope:  Equation System

Rad Flux For Cvfem_Energy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Rad [ Using Data Specification Data Spec Name ] [ Data = data | T_Ref = t_ref | Crad = crad ]

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Summary: CVFEM radiative energy flux defined with constant models for emissivity, form factor and reference temperature

9.2.131  Generalized Rad

Scope:  Equation System

Generalized Rad Flux For Cvfem_Energy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Generalized.Rad [ Using Data Specification Data Spec Name ] [ Data = data | Multiplier = multiplier ]
Summary: Radiative energy flux, possibly defined with different models for emissivity, form factor and irradiation.

9.2.132 Nat Conv

Scope: Equation System

Nat Conv Flux For Cvfem_Energy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Nat_Conv [ Using Data Specification Data Spec Name ] [ Data = data |H = h |T_Ref = t_ref ]

Summary: CVFEM convective energy flux defined with constant models for heat transfer coefficient and reference temperature.

9.2.133 Open Adv Flow

Scope: Equation System

Open Adv Flow Flux For Cvfem_Energy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Open_Adv_Flow [ Using Data Specification Data Spec Name ] [ Data = data |Far_Field_Entrainment_Value = Far_Field_Entrainment_Value | Pressure = pressure |Total_Pressure = total_pressure ]

Summary: CVFEM energy open advective flow flux.
9.2.134 Open Adv Flow From Temperature

Scope: Equation System

Open Adv Flow From Temperature Flux For Cvfem_Energy [{of|species|subindex} Species | {in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Open_Adv_Flow_From_Temperature [ Using Data Specification Data Spec Name ][ Data = data |Temperature = temperature |Pressure = pressure |Mass_Fraction_0 = mass_fraction_0 |Mass_Fraction_1 = mass_fraction_1 |Mass_Fraction_2 = mass_fraction_2 |Mass_Fraction_3 = mass_fraction_3 |Mass_Fraction_4 = mass_fraction_4 |Mass_Fraction_5 = mass_fraction_5 |Mass_Fraction_6 = mass_fraction_6 |Mass_Fraction_7 = mass_fraction_7 |Mass_Fraction_8 = mass_fraction_8 |Total_Pressure = total_pressure ]

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Summary
CVFEM energy open advective flow from temperature flux

9.2.135 Mass Flux

Scope: Equation System

Mass Flux Flux For Cvfem_Energy [{of|species|subindex} Species | {in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Mass_Flux [ Using Data Specification Data Spec Name ][ Data = data |Inflow_Phi = Inflow_Phi |Total_Mdot = total_mdot ]

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Summary
CVFEM energy mass flux flux BC
9.2.136  Mass Flux From Temperature

Scope:  Equation System

Mass Flux From Temperature Flux For Cvfem_Energy [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Mass_Flux_From_Temperature [ Using Data Specification Data Spec Name ][ Data = data |Temperature = temperature |Pressure = pressure |Mass_Fraction_0 = mass_fraction_0 |Mass_Fraction_1 = mass_fraction_1 |Mass_Fraction_2 = mass_fraction_2 |Mass_Fraction_3 = mass_fraction_3 |Mass_Fraction_4 = mass_fraction_4 |Mass_Fraction_5 = mass_fraction_5 |Mass_Fraction_6 = mass_fraction_6 |Mass_Fraction_7 = mass_fraction_7 |Mass_Fraction_8 = mass_fraction_8 |Total_Mdot = total_mdot ]

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Summary  CVFEM energy mass flux from temperature flux BC

9.2.137  Porous Robin Coupled One Region

Scope:  Equation System

Porous Robin Coupled One Region Flux For Cvfem_Energy [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Porous_Robin_Coupled_One_Region [ Using Data Specification Data Spec Name ][ Data = data ]

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Summary  Boundary condition for loose coupling of enthalpy to a porous region

9.2.138  Convective Outflow

Scope:  Equation System
Convective Outflow Flux For Enthalpy \{ of \{ species | subindex \} Species \{ in \{ material_phase \} MaterialPhase \} @ \{ at \{ for \{ in \{ on \{ over \} \} \} \} \} Mesh Extent Name = Convective_Outflow [ Using Data Specification Data Spec Name ][ ]

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9.2.139 Porous Robin Coupled With Solid Phase Convection One Region

Scope: Equation System

Porous Robin Coupled With Solid Phase Convection One Region Flux For Cvem_Energy \{ of \{ species | subindex \} Species \{ in \{ material_phase \} MaterialPhase \} @ \{ at \{ for \{ in \{ on \{ over \} \} \} \} \} Mesh Extent Name = Porous_Robin_Coupled_With_Solid_Phase_Convection_One_Region [ Using Data Specification Data Spec Name ][ Data = data | Multiplier = multiplier ]

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Summary Boundary condition for coupling of enthalpy to a porous region in the same region.

9.2.140 Porous Robin Coupled With Solid Phase Convection

Scope: Equation System

Porous Robin Coupled With Solid Phase Convection Flux For Cvem_Energy \{ of \{ species | subindex \} Species \{ in \{ material_phase \} MaterialPhase \} @ \{ at \{ for \{ in \{ on \{ over \} \} \} \} \} Mesh Extent Name = Porous_Robin_Coupled_With_Solid_Phase_Convection [ Using Data Specification Data Spec Name ][ Data = data | Multiplier = multiplier | Enthalpy_Field = enthalpy_field | Enthalpy_Diffusive_Flux_Field = enthalpy_diffusive_flux_field | Solid_Phase_Temperature_Field = solid_phase_temperature_field | Porosity_Field = porosity_field | Averaged_Coeff_Field = averaged_coeff | Volumetric_Heat_Transfer_Coeff_Field = volumetric_heat_transfer_coeff_field | Specific_Surface_Area_Field = specific_surface_area_field ]

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Parameter | Value | Default
--- | --- | ---
Species | string | undefined
MaterialPhase | string | undefined
Mesh Extent Name | string | undefined
Data Spec Name | string | undefined
data | "string" | undefined
multiplier | real | undefined
enthalpy_field | "string" | undefined
enthalpy_diffusive_flux_field | "string" | undefined
solid_phase_temperature_field | "string" | undefined
porosity_field | "string" | undefined
averaged_coeff_field | "string" | undefined
volumetric_heat_transfer_coeff_field | "string" | undefined
specific_surface_area_field | "string" | undefined

Summary: Boundary condition for loose coupling of enthalpy to a porous region

9.2.141 Free Open Flow
Scope: Equation System

Free Open Flow Flux For Cvfem_Turbulent_Kinetic_Energy [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ] {@|at|for|in|on|over} Mesh Extent Name = Free_Open_Flow [ Using Data Specification Data Spec Name ]

Parameter | Value | Default
--- | --- | ---
Species | string | undefined
MaterialPhase | string | undefined
Mesh Extent Name | string | undefined
Data Spec Name | string | undefined
total_mdot | real | undefined
inflow_phi | real | undefined

9.2.142 Mass Flux
Scope: Equation System

Mass Flux Flux For Cvfem_Turbulent_Kinetic_Energy [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ] {@|at|for|in|on|over} Mesh Extent Name = Mass_Flux [ Using Data Specification Data Spec Name ] [ Total_Mdot = total_mdot | Inflow_Phi = inflow_phi ]

Parameter | Value | Default
--- | --- | ---
Species | string | undefined
MaterialPhase | string | undefined
Mesh Extent Name | string | undefined
Data Spec Name | string | undefined
total_mdot | real | undefined
inflow_phi | real | undefined

9.2.143 Open Adv Flow
Scope: Equation System
Open Adv Flow Flux For Cvfem_Turbulent_Kinetic_Energy [ {of|species|subindex} Species | {in|material_phase} MaterialPhase (){at|for|in|on|over} Mesh Extent Name = Open_Adv_Flow [ Using Data Specification Data Spec Name ][ Pressure = pressure |Total_Pressure = total_pressure |Far_Field_Entrainment_Value = far_field_entrainment_value ]

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9.2.144  Nat E
Scope:  Equation System

Nat E Flux For Charge_Density [ {of|species|subindex} Species | {in|material_phase} MaterialPhase (){at|for|in|on|over} Mesh Extent Name = Nat_E [ Using Data Specification Data Spec Name ][ Sigma = sigma ]

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<td>sigma</td>
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Summary  Natural J = conductivity times Electric field BC that falls out of flux diffusion term

9.2.145  Nat Conv
Scope:  Equation System

Nat Conv Flux For Energy [ {of|species|subindex} Species | {in|material_phase} MaterialPhase (){at|for|in|on|over} Mesh Extent Name = Nat_Conv [ Using Data Specification Data Spec Name ][ Power_Output = power_output |Flux_Output = flux_output |Toggle = toggle |H = h |T_Ref = t_ref |Advective_Bar = advective_bar |Command_Block_Name = command_block_name ]

<table>
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Summary
Convective energy flux defined with constant models for heat transfer coefficient and reference temperature

9.2.146 From Cht Temperature Cpt

Scope: Equation System

From Cht Temperature Cpt Distinctive For Cvfem_Energy [ of | species | subindex ] Species | { in | material_phase } MaterialPhase ] | @ | at | for | in | on | over | Mesh Extent Name = From_Cht_Temperature_Cpt [ Using Data Specification Data Spec Name ] [ Twall_Field = Twall_Field | Variable = variable | Order = order | Variable_Offset = variable_offset | C0 = c0 | C1 = c1 | C2 = c2 | C3 = c3 | C4 = c4 | C5 = c5 | C6 = c6 | C7 = c7 | C8 = c8 ]

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Summary
CVFEM energy distinguishing condition from CHT temperature using Cp*T model

9.2.147 From Cht Temperature

Scope: Equation System

From Cht Temperature Distinguishing For Cvfem_Energy [ of | species | subindex ] Species | { in | material_phase } MaterialPhase ] | @ | at | for | in | on | over | Mesh Extent Name = From_Cht_Temperature [ Using Data Specification Data Spec Name ] [ Twall_Field = Twall_Field | Variable = variable | Order = order | Variable_Offset = variable_offset | C0 = c0 | C1 = c1 | C2 = c2 | C3 = c3 | C4 = c4 | C5 = c5 | C6 = c6 | C7 = c7 | C8 = c8 ]

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</table>

Summary CVFEM energy distinguishing condition from CHT temperature using Cantera

9.2.148 From Temperature

Scope: Equation System

From Temperature Disting For Cvem_Energy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@at|for|in|on|over} Mesh Extent Name = From_Temperature [ Using Data Specification Data Spec Name ][ Variable = variable |Order = order |Variable_Offset = variable_offset | C0 = c0 |C1 = c1 |C2 = c2 |C3 = c3 |C4 = c4 |C5 = c5 |C6 = c6 |C7 = c7 |C8 = c8 |T = t ]

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Summary From temperature distinguishing condition for CVFEM energy equation
9.2.149  Inflow
Scope:   Equation System

Inflow Flux For Hfem_Continuity [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@[at|for|in|on|over} Mesh Extent Name = Inflow [ Using Data Specification Data Spec Name ][]

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<tr>
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Summary  IBP continuity inflow; nodal interp

9.2.150  Suspension Traction
Scope:   Equation System

Suspension Traction Flux For Div_Projection [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@[at|for|in|on|over} Mesh Extent Name = Suspension_Traction [ Using Data Specification Data Spec Name ][]

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Summary  Suspension traction for suspension balance model

9.2.151  Curvature
Scope:   Equation System

Curvature Flux For Div_Projection [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@[at|for|in|on|over} Mesh Extent Name = Curvature [ Using Data Specification Data Spec Name ][ Theta = theta ]

<table>
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<th>Value</th>
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Summary  Curvature model for div projection flux

9.2.152  Capillary Force
Scope:   Equation System
Capillary Force Flux For Div_Projection [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Capillary_Force [ Using Data Specification Data Spec Name ] [ Theta = theta ]

<table>
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Summary
Capillary force model for div projection flux

9.2.153 Advection
Scope: Equation System

Advection Disting For Level_Set [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Advection [ Using Data Specification Data Spec Name ] [ ]

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Summary
Advection model for extension speed distinguishing condition

9.2.154 Recession
Scope: Equation System

Recession Disting For Level_Set [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Recession [ Using Data Specification Data Spec Name ] [ P_Ref = p_ref | K = k | K_0 = k_0 | N = n | Pressure = pressure ]

<table>
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<th>Value</th>
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Summary
Advection model for extension speed distinguishing condition
9.2.155  Kinematic

Scope:  Equation System

Kinematic Distinguishing For Mesh_Z [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Kinematic [ Using Data Specification Data Spec Name ] [ V0 = v0 | Shared_Normal = shared_normal | No_Mesh_Movement = no_mesh_movement ]

Parameter | Value | Default
--- | --- | ---
Species | string | undefined
MaterialPhase | string | undefined
Mesh Extent Name | string | undefined
Data Spec Name | string | undefined
v0 | real | undefined
shared_normal | "string" | undefined
no_mesh_movement | "string" | undefined

Summary  Kinematic model for mesh_z distinguishing condition

9.2.156  Recession

Scope:  Equation System

Recession Distinguishing For Mesh_Z [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Recession [ Using Data Specification Data Spec Name ] [ P_Ref = p_ref | K = k | K_0 = k_0 | N = n | Pressure = pressure ]

Parameter | Value | Default
--- | --- | ---
Species | string | undefined
MaterialPhase | string | undefined
Mesh Extent Name | string | undefined
Data Spec Name | string | undefined
p_ref | real | undefined
k | real | undefined
k_0 | real | undefined
n | real | undefined
pressure | "string" | undefined

Summary  Recession model for mesh_x distinguishing condition

9.2.157  Open Adv Flow

Scope:  Equation System

Open Adv Flow Flux For Hfem_Mixture_Fraction [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Open_Adv_Flow [ Using Data Specification Data Spec Name ] [ Pressure = pressure | Total_Pressure = total_pressure | Far_Field_Entrainment_Value = far_field_entrainment_value ]
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Summary: HFEM mixture fraction open advective flow flux BC

### 9.2.158 Open Flow

Scope: Equation System

Open Flow Flux For Cvfem_Dispersed_Continuity [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ][@at|for|in|on|over} Mesh Extent Name = Open_Flow [ Using Data Specification Data Spec Name ][ ]

<table>
<thead>
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Summary: Open bc for dispersed continuity

### 9.2.159 Open Flow

Scope: Equation System

Open Flow Flux For Cvfem_Dispersed_Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ][@at|for|in|on|over} Mesh Extent Name = Open_Flow [ Using Data Specification Data Spec Name ][ ]

<table>
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<th>Parameter</th>
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</table>

### 9.2.160 Open Adv Flow

Scope: Equation System

Open Adv Flow Flux For Cvfem_Dispersed_Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ][@at|for|in|on|over} Mesh Extent Name = Open_Adv_Flow [ Using Data Specification Data Spec Name ][ Coflow_Entrainment_Valuex = Coflow_Entrainment_ValueX |Coflow_Entrainment_Valuey = Coflow_Entrainment_ValueY |Coflow_Entrainment_Valuez = Coflow_Entrainment_ValueZ ]
Parameter | Value | Default
--- | --- | ---
Species | string | undefined
MaterialPhase | string | undefined
Mesh Extent Name | string | undefined
Data Spec Name | string | undefined
Coflow Entrainment ValueX | real | undefined
Coflow Entrainment ValueY | real | undefined
Coflow Entrainment ValueZ | real | undefined

9.2.161 Symmetry Flow

Scope: Equation System

Symmetry Flow Flux For Cvfem Dispersed Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase |{at|for|in|on|over} Mesh Extent Name = Symmetry_Flow [ Using Data Specification Data Spec Name ][ ]

Parameter | Value | Default
--- | --- | ---
Species | string | undefined
MaterialPhase | string | undefined
Mesh Extent Name | string | undefined
Data Spec Name | string | undefined

9.2.162 Simple Interp Vector

Scope: Equation System

Simple Interp Vector Flux For Cvfem Div Projection [ {of|species|subindex} Species |{in|material_phase} MaterialPhase |{at|for|in|on|over} Mesh Extent Name = Simple_Interp_Vector [ Using Data Specification Data Spec Name ][ Source = source ]

Parameter | Value | Default
--- | --- | ---
Species | string | undefined
MaterialPhase | string | undefined
Mesh Extent Name | string | undefined
Data Spec Name | string | undefined
source | "string" | undefined

Summary CVFEM divergence projection simple interpolation vector flux BC

9.2.163 Simple Interp Tensor

Scope: Equation System

Simple Interp Tensor Flux For Cvfem Div Projection [ {of|species|subindex} Species |{in|material_phase} MaterialPhase |{at|for|in|on|over} Mesh Extent Name = Simple_Interp_Tensor [ Using Data Specification Data Spec Name ][ Source = source ]
<table>
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<tr>
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Summary: CVFEM divergence projection simple interpolation tensor flux BC

### 9.2.164 Open Flow

**Scope:** Equation System

---

Open Flow Flux For Hfem_Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@at|for|in|on|over} Mesh Extent Name = Open_Flow [ Using Data Specification Data Spec Name ]([ Pressure = pressure |Total_Pressure = total_pressure ]

<table>
<thead>
<tr>
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<th>Default</th>
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</thead>
<tbody>
<tr>
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<tr>
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<td>undefined</td>
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<tr>
<td>total_pressure</td>
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</tbody>
</table>

Summary: Open flow for momentum; enforce normal gradient to be zero

### 9.2.165 Free Open Flow

**Scope:** Equation System

---

Free Open Flow Flux For Hfem_Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@at|for|in|on|over} Mesh Extent Name = Free_Open_Flow [ Using Data Specification Data Spec Name ]([ Pressure = pressure |Pressure User Sub PRESSURE User Sub ]

<table>
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<tr>
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</table>

Summary: Open flow for momentum; apply the full stress

### 9.2.166 Open Flow

**Scope:** Equation System

---

Open Flow Flux For Porous_Species [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@at|for|in|on|over} Mesh Extent Name = Open_Flow [ Using Data Specification Data Spec Name ][ ]
### 9.2.167 No Slip

**Scope:** Equation System

No Slip Disting For Solid [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@{at|for|in|on|over} Mesh Extent Name = No_Slip [ Using Data Specification Data Spec Name ][ V0 = v0 ]

<table>
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<td>Data Spec Name</td>
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<tr>
<td>v0</td>
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**Summary**
No slip model for SOLID distinguishing condition

### 9.2.168 Simple Interp

**Scope:** Equation System

Simple Interp Flux For Cvfem_Pressure_Edge_Gradient_Projection [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@{at|for|in|on|over} Mesh Extent Name = Simple_Interp [ Using Data Specification Data Spec Name ]

<table>
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<th>Value</th>
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<td>Data Spec Name</td>
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</table>

**Summary**
CVFEM edge-based projection simple interpolation flux BC

### 9.2.169 Open Flow

**Scope:** Equation System

Open Flow Flux For Hfem_Continuity [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@{at|for|in|on|over} Mesh Extent Name = Open_Flow [ Using Data Specification Data Spec Name ][ Pressure = pressure | Total_Pressure = total_pressure ]
Summary  Open bc for continuity

9.2.170 Normal Capillary Stabilization

Scope: Equation System

Normal Capillary Stabilization Fluxbp For Hfem_Momentum [{of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Normal_Capillary_Stabilization [ Using Data Specification Data Spec Name ][ Mu0 = mu0 |Multiplier = multiplier ]

Summary  Momentum capillary stabilization flux BP BC

9.2.171 Ls Capillary Stabilization

Scope: Equation System

Ls Capillary Stabilization Fluxbp For Hfem_Momentum [{of|species|subindex} Species |{in |material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Ls_Capillary_Stabilization [ Using Data Specification Data Spec Name ][ Mu0 = mu0 |Multiplier = multiplier ]

Summary  Momentum capillary stabilization flux BP BC using level set normal

9.2.172 Simple Interp

Scope: Equation System
Simple Interp Flux For Hfem_Projection [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Simple_Interp [ Using Data Specification Data Spec Name ]

<table>
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Summary: Simple interp model for projection flux

9.2.173 Open
Scope: Equation System

Open Flux For Hfem_Projection [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Open [ Using Data Specification Data Spec Name ]

<table>
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<th>Value</th>
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Summary: Open model for projection flux

9.2.174 Spring Force
Scope: Equation System

Spring Force Flux For Solid [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Spring_Force [ Using Data Specification Data Spec Name ]

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Summary: Spring force solid flux

9.2.175 Kinematic
Scope: Equation System
Kinematic Disting For Solid_X \{of\species\subindex\ \Species\ |{in|material\phase}\ Material\Phase\}\at\for\in\on\over\ Mesh\ Extent\ Name = Kinematic \[ Using\ Data\ Specification\ Data\ Spec\ Name \]\ V0 = v0 \ Shared\_Normal = \_\_\_shared\_normal \ No\_Mesh\_Movement = no\_mesh\_movement \]

<table>
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Summary: Kinematic model for SOLID\_x distinguishing condition

9.2.176 Recession

Scope: Equation System

Recession Disting For Solid_X \{of\species\subindex\ \Species\ |{in|material\phase}\ Material\Phase\}\at\for\in\on\over\ Mesh\ Extent\ Name = Recession \[ Using\ Data\ Specification\ Data\ Spec\ Name \]\ P\_Ref = p\_ref \ K = k \ K\_0 = k\_0 \ N = n \ Pressure = pressure \]

<table>
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Summary: Recession model for SOLID\_x distinguishing condition

9.2.177 Open Flow

Scope: Equation System

Open Flow Flux For Mass\_Balance \{of\species\subindex\ \Species\ |{in|material\phase}\ Material\Phase\}\at\for\in\on\over\ Mesh\ Extent\ Name = Open\_Flow \[ Using\ Data\ Specification\ Data\ Spec\ Name \]\]

<table>
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</table>

9.2.178 Mass Open

Scope: Equation System
Mass Open Flux For Mass Balance [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Mass_Open [ Using Data Specification Data Spec Name ][ Pressure = pressure ]

<table>
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9.2.179 Simple Interp
Scope: Equation System

Simple Interp Flux For CvFEM_Temperature_Edge_Gradient_Projection [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Simple_Interp [ Using Data Specification Data Spec Name ]

<table>
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</table>

Summary

CVFEM edge-based projection simple interpolation flux BC

9.2.180 Constant Velocity
Scope: Equation System

Constant Velocity Flux For Mass Balance [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Constant_Velocity [ Using Data Specification Data Spec Name ] [ Mult = Mult ]

<table>
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9.2.181 Co2 Convective Outflow
Scope: Equation System

Co2 Convective Outflow Flux For Mass Balance [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Co2_Convective_Outflow [ Using Data Specification Data Spec Name ] [ Ref_Frac = ref_frac ]
9.2.182  Kinematic

Scope:  Equation System

Summary  Kinematic model for SOLID_z distinguishing condition

9.2.183  Recession

Scope:  Equation System

Summary  Recession model for SOLID_x distinguishing condition

9.2.184  Curvature

Scope:  Equation System
Curvature Flux For Lumped Div Projection [ \{of|species|subindex\} \text{Species} | \{in|material\_phase\} \text{MaterialPhase} ][@|at|for|in|on|over\} \text{Mesh Extent Name} = \text{Curvature} [ \text{Using Data Specification Data Spec Name} ][ \Theta = \theta ]

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Summary Curvature model for lumped div projection flux

9.2.185 Outflow
Scope: Equation System

Outflow Disting For Level Set [ \{of|species|subindex\} \text{Species} | \{in|material\_phase\} \text{MaterialPhase} ][@|at|for|in|on|over\} \text{Mesh Extent Name} = \text{Outflow} [ \text{Using Data Specification Data Spec Name} ][ \text{V0} = v0 ]

<table>
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Summary Outflow model for level set distinguishing condition

9.2.186 Nat Conv
Scope: Equation System

Nat Conv Flux For Species [ \{of|species|subindex\} \text{Species} | \{in|material\_phase\} \text{MaterialPhase} ][@|at|for|in|on|over\} \text{Mesh Extent Name} = \text{Nat_Conv} [ \text{Using Data Specification Data Spec Name} ][ \text{K} = k | \text{Yinf} = yinf ]

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Summary natconv species flux

9.2.187 Free Open Flow
Scope: Equation System
Free Open Flow Flux For Cvfem_Mixture_Fraction [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ][@|at|for|in|on|over} Mesh Extent Name = Free_Open_Flow [ Using Data Specification Data Spec Name ]

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9.2.188 Recession

Scope: Equation System

Recession Rotated For Cvfem_Mesh [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ][@|at|for|in|on|over} Mesh Extent Name = Recession [ Using Data Specification Data Spec Name ][ P_Ref = p_ref | K = k | K_0 = k_0 | N = n | Pressure = pressure ]

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Summary Recession model for rotated CVFEM_MESH boundary condition

9.2.189 Kinematic

Scope: Equation System

Kinematic Rotated For Cvfem_Mesh [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ][@|at|for|in|on|over} Mesh Extent Name = Kinematic [ Using Data Specification Data Spec Name ][ V0 = v0 | Shared_Normal = shared_normal | No_Mesh_Movement = no_mesh_movement ]

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Summary Kinematic model for rotated CVFEM_MESH boundary condition
9.2.190  Open Flow
Scope:  Equation System

Open Flow Flux For Cvfem_Solvent_Continuity [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Open_Flow [ Using Data Specification Data Spec Name ][ Pressure = pressure |Total_Pressure = total_pressure ]

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Summary  Open bc for continuity

9.2.191  Inflow
Scope:  Equation System

Inflow Flux For Cvfem_Solvent_Continuity [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Inflow [ Using Data Specification Data Spec Name ]

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Summary  Inflow mass flux for continuity equation; nodal interpolation

9.2.192  Simple Inflow
Scope:  Equation System

Simple Inflow Flux For Cvfem_Solvent_Continuity [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Simple_Inflow [ Using Data Specification Data Spec Name ][ Density = Density |Velocity_X = Velocity_X |Velocity_Y = Velocity_Y |Velocity_Z = Velocity_Z ]

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9.2.193 Open Flow

Scope: Equation System

Open Flow Flux For CvFem_Solvent_Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ] @ {at|for|in|on|over} Mesh Extent Name = Open_Flow [ Using Data Specification Data Spec Name ] [ Pressure = pressure | Total_Pressure = total_pressure ]

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Summary CVFEM solvent momentum open flow flux BC

9.2.194 Kinematic

Scope: Equation System

Kinematic Disting For Mesh_X [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ] @ {at|for|in|on|over} Mesh Extent Name = Kinematic [ Using Data Specification Data Spec Name ] [ V0 = v0 | Shared_Normal = shared_normal | No_Mesh_Movement = no_mesh_movement ]

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Summary Kinematic model for mesh_x distinguishing condition

9.2.195 Recession

Scope: Equation System

Recession Disting For Mesh_X [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ] @ {at|for|in|on|over} Mesh Extent Name = Recession [ Using Data Specification Data Spec Name ] [ P_Ref = p_ref | K = k | K_0 = k_0 | N = n | Pressure = pressure ]

485
### 9.2.196 Pressure

**Scope:** Equation System

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**Summary**  
Recession model for mesh_x distinguishing condition

---

### 9.2.197 No Slip

**Scope:** Equation System

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**Summary**  
Pressure mesh flux

---

486
### 9.2.198 Open Flow

**Scope:** Equation System

Open Flow Flux For Cvfem_Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Open_Flow [ Using Data Specification Data Spec Name ][ Pressure = pressure |Total_Pressure = total_pressure ]

<table>
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**Summary**  
Open flow for momentum; enforce normal gradient to be zero

### 9.2.199 Mass Flux

**Scope:** Equation System

Mass Flux Flux For Cvfem_Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Mass_Flux [ Using Data Specification Data Spec Name ][ Total_Mdot = total_mdot |Temperature = temperature |Pressure = pressure |Mass_Fraction_0 = mass_fraction_0 |Mass_Fraction_1 = mass_fraction_1 |Mass_Fraction_2 = mass_fraction_2 |Mass_Fraction_3 = mass_fraction_3 |Mass_Fraction_4 = mass_fraction_4 |Mass_Fraction_5 = mass_fraction_5 |Mass_Fraction_6 = mass_fraction_6 |Mass_Fraction_7 = mass_fraction_7 |Mass_Fraction_8 = mass_fraction_8 ]

**Summary**  
Open flow for momentum; enforce normal gradient to be zero
9.2.200  Normal Capillary Stabilization
Scope:  Equation System

Normal Capillary Stabilization Fluxbp For Cvfem_Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@at|for|in|on|over} Mesh Extent Name = Normal_Capillary_Stabilization [ Using Data Specification Data Spec Name ][ Mu0 = mu0 |Multiplier = multiplier ]

<table>
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Summary  Momentum capillary stabilization flux BP BC

9.2.201  Capillary Stabilization
Scope:  Equation System

Capillary Stabilization Fluxbp For Cvfem_Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@at|for|in|on|over} Mesh Extent Name = Capillary_Stabilization [ Using Data Specification Data Spec Name ][ Mu0 = mu0 |Multiplier = multiplier ]

<table>
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Summary  Momentum capillary stabilization flux BP BC
9.2.202  Ls Capillary Stabilization
Scope:  Equation System

Ls Capillary Stabilization Fluxbp For Cvem_Momentum  
Species | 
MaterialPhase |  
Mesh Extent Name = Ls_Capillary_Stabilization  
[ Using Data Specification Data Spec Name ]  
[ Mu0 = mu0 | Multiplier = multiplier ]

<table>
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Summary  Momentum capillary stabilization flux BP BC using level set normal

9.2.203  Free Open Flow
Scope:  Equation System

Free Open Flow Flux For Cvem_Momentum  
Species | 
MaterialPhase |  
Mesh Extent Name = Free_Open_Flow  
[ Using Data Specification Data Spec Name ]  
[ Pressure = pressure | Pressure User Sub PRESSURE User Sub ]

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Summary  Open flow for momentum; apply the full stress

9.2.204  Constant Traction
Scope:  Equation System

Constant Traction Flux For Cvem_Momentum  
Species | 
MaterialPhase |  
Mesh Extent Name = Constant_Traction  
[ Using Data Specification Data Spec Name ]  
[ X = x | Y = y | Z = z ]

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9.2.205 Simple Inflow
Scope: Equation System

Simple Inflow Flux For Cvfem_Dispersed_Continuity [ {of | species | subindex} Species | {in | material_phase} MaterialPhase ] @ at | for | in | on | over Mesh Extent Name = Simple_Inflow [ Using Data Specification Data Spec Name ] [ Velocity_X = Velocity_X | Velocity_Y = Velocity_Y | Velocity_Z = Velocity_Z ]

<table>
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Summary: Inflow mass flux for dispersed continuity equation; specified density and velocity

9.2.206 Convective Outflow
Scope: Equation System

Convective Outflow Flux For Porous_Enthalpy [ {of | species | subindex} Species | {in | material_phase} MaterialPhase ] @ at | for | in | on | over Mesh Extent Name = Convective_Outflow [ Using Data Specification Data Spec Name ] [ Ref_H = ref_h ]

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9.2.207 User Vector Field
Scope: Equation System

User Vector Field Flux For Porous_Enthalpy [ {of | species | subindex} Species | {in | material_phase} MaterialPhase ] @ at | for | in | on | over Mesh Extent Name = User_Vector_Field [ Using Data Specification Data Spec Name ] [ Name = name | Multiplier = multiplier ]

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Summary: User vector field enthalpy flux
9.2.208  Open
Scope:  Equation System

Open Flux For Porous_Enthalpy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Open [ Using Data Specification Data Spec Name ][ Pressure = pressure |Entrained_Value = entrained_value ]

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9.2.209  Fluid Coupled
Scope:  Equation System

Fluid Coupled Flux For Porous_Enthalpy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Fluid_Coupled [ Using Data Specification Data Spec Name ][ Pressure_Field = pressure_field |Density_Field = density_field |Enthalpy_Field = enthalpy_field ]

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Summary  Boundary condition for loose coupling of enthalpy to a free fluid region

9.2.210  Generalized Nat Conv
Scope:  Equation System

Generalized Nat Conv Flux For Porous_Enthalpy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Generalized_Nat_Conv [ Using Data Specification Data Spec Name ][ Multiplier = multiplier ]

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Summary  Generalized natural convection flux BC
9.2.211 Fluid Solid Convection Coupled

Scope: Equation System

Fluid Solid Convection Coupled Flux For Porous_Enthalpy [ Species |in| MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Fluid_Solid_Convection_Coupled

Using Data Specification Data Spec Name [ Multiplier = multiplier | Fluid_Temperature_Field = fluid_temperature_field ]

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Summary  Boundary condition for loose coupling of enthalpy to a free fluid region

9.2.212 Fluid Robin Coupled

Scope: Equation System

Fluid Robin Coupled Flux For Porous_Enthalpy [ Species |in| MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Fluid_Robin_Coupled

Using Data Specification Data Spec Name [ Enthalpy_Field = enthalpy_field | Enthalpy_Diffusive_Flux_Field = enthalpy_diffusive_flux_field | Coeff_Field = coeff_field ]

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Summary  Boundary condition for loose coupling of enthalpy to a free fluid region

9.2.213 Fluid Robin Coupled One Region

Scope: Equation System

Fluid Robin Coupled One Region Flux For Porous_Enthalpy [ Species |in| MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Fluid_Robin_Coupled_OneRegion

Using Data Specification Data Spec Name [ ]

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9.2.214 Fluid Robin Coupled With Solid Convection

Scope: Equation System

Fluid Robin Coupled With Solid Convection Flux For Porous_Enthalpy [ {of|species|subindex} 
Species |{in|material_phase} MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Fluid_Robin_Coupled_With_Solid_Convection
[ Using Data Specification Data Spec Name ][ Enthalpy_Field = enthalpy_field |Enthalpy_Diffusive_Flux_Field = enthalpy_diffusive_flux_field |Coeff_Field = coeff_field ]

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Summary: Boundary condition for loose coupling of enthalpy to a free fluid region

9.2.215 Wall Function

Scope: Equation System

Wall Function Disting For Cvfem_Specific_Dissipation_Rate [ {of|species|subindex} 
Species |{in|material_phase} MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Wall_Function
[ Using Data Specification Data Spec Name ][ Wall_Friction_Factor = Wall_Friction_Factor | Wall_Velocity_X = Wall_Velocity_X |Wall_Velocity_Y = Wall_Velocity_Y |Wall_Velocity_Z = Wall_Velocity_Z ]

<table>
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Summary: CVFEM specific dissipation rate distinguishing condition

9.2.216 Wall Function

Scope: Equation System

Wall Function Disting For Hfem_Specific_Dissipation_Rate [ {of|species|subindex} 
Species |{in|material_phase} MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Wall_Function
[ Using Data Specification Data Spec Name ][ Scale_Factor = scale_factor |Wall_Friction_Factor

493
Wall_Friction_Factor | Wall_Velocity_X = Wall_Velocity_X | Wall_Velocity_Y = Wall_Velocity_Y | Wall_Velocity_Z = Wall_Velocity_Z

<table>
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Summary: HFEM specific dissipation rate distinguishing condition

9.2.217  Low Reynolds

Scope: Equation System

Low Reynolds Distinguishing Condition for Hfem_Specific_Dissipation_Rate [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Low_Reynolds [ Using Data Specification Data Spec Name ][ Scale_Factor = scale_factor ]

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Summary: HFEM specific dissipation rate distinguishing condition

9.2.218  Interface

Scope: Equation System

Interface Flux For Cvfm_Continuity [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Interface [ Using Data Specification Data Spec Name ]

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Summary: INTERFACE CVFEM continuity flux

9.2.219  Simple Interp

Scope: Equation System
Simple Interp Flux For Cvfem_Sdr_Edge_Gradient_Projection [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ] {at|for|in|on|over} Mesh Extent Name = Simple_Interp [ Using Data Specification Data Spec Name ][ ]

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Summary CVFEM edge-based projection simple interpolation flux BC

9.2.220 Inflow
Scope: Equation System

Inflow Flux For Cvfem_Dispersed_Continuity [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ] {at|for|in|on|over} Mesh Extent Name = Inflow [ Using Data Specification Data Spec Name ][ ]

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Summary Inflow mass flux for dispersed phase continuity equation; nodal interpolation

9.2.221 Simple Interp
Scope: Equation System

Simple Interp Flux For Cvfem_Density_Edge_Gradient_Projection [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ] {at|for|in|on|over} Mesh Extent Name = Simple_Interp [ Using Data Specification Data Spec Name ][ ]

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Summary CVFEM edge-based projection simple interpolation flux BC

9.2.222 Rte Sp
Scope: Equation System

Rte Sp Flux For Energy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ] {at|for|in|on|over} Mesh Extent Name = Rte_Sp [ Using Data Specification Data Spec Name ]

495
Parameter | Value | Default
--- | --- | ---
Species | string | undefined
MaterialPhase | string | undefined
Mesh Extent Name | string | undefined
Data Spec Name | string | undefined
power_output | "string" | undefined
flux_output | "string" | undefined
toggle | "string" | undefined
order | integer | undefined

Summary  
RTE flux for energy equation with Simplified Spherical Harmonic RTE model

### 9.2.223 Rte Field

**Scope:** Equation System

Rte Field Flux For Energy [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ] @ {at|for|in|on|over} Mesh Extent Name = Rte_Field [ Using Data Specification Data Spec Name ] [ Power_Output = power_output | Flux_Output = flux_output | Toggle = toggle ]

Parameter | Value | Default
--- | --- | ---
Species | string | undefined
MaterialPhase | string | undefined
Mesh Extent Name | string | undefined
Data Spec Name | string | undefined
power_output | "string" | undefined
flux_output | "string" | undefined
toggle | "string" | undefined

Summary  
RTE energy flux from a field

### 9.2.224 Cht Dirichlet Robin

**Scope:** Equation System

Cht Dirichlet Robin Flux For Energy [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ] @ {at|for|in|on|over} Mesh Extent Name = Cht_Dirichlet_Robin [ Using Data Specification Data Spec Name ] [ Power_Output = power_output | Flux_Output = flux_output | Toggle = toggle | Temperature_Field = temperature_field | Heat_Flux_Field = heat_flux_field | Density_Field = density_field | Specific_Heat_Field = specific_heat_field | Thermal_Conductivity_Field = thermal_conductivity_field ]
Parameter | Value | Default
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Summary
Applies a Dirichlet-Robin style BC for the energy equation: $F_{local} = F_{bc} + \alpha \times (T_{local} - T_{bc})$, where $\alpha$ is computed based on a 1D analytical solution.

9.2.225 Kinematic

Scope: Equation System

Summary
Kinematic model for rotated momentum

9.2.226 Simple Spring Force

Scope: Equation System

Summary
Simple Spring Force Flux For Momentum [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Simple_Spring_Force [ Using Data Specification Data Spec Name ][ Kx = kx | Ky = ky | Kz = kz | X0 = x0 | Y0 = y0 | Z0 = z0 ]
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Summary: Spring force for momentum boundary condition

### 9.2.227 Spring Force

**Scope:** Equation System

```latex
Spring Force Flux For Momentum [ \{\text{of} | \text{species} | \text{subindex}\} \text{Species} | \{\text{in} | \text{material\_phase}\} \text{MaterialPhase} |\{@ | \text{at} | \text{for} \text{|in}\text{|on}\text{|over}} \text{Mesh Extent Name} = \text{Spring\_Force} [ \text{Using Data Specification Data Spec Name} ][ \text{Mult} = \text{Mult} ]
```

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<tr>
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Summary: Spring force for momentum boundary condition

### 9.2.228 Capillary Stabilization

**Scope:** Equation System

```latex
Capillary Stabilization Fluxbp For Hfem\_Momentum [ \{\text{of} | \text{species} | \text{subindex}\} \text{Species} | \{\text{in} | \text{material\_phase}\} \text{MaterialPhase} |\{@ | \text{at} | \text{for} \text{|in}\text{|on}\text{|over}} \text{Mesh Extent Name} = \text{Capillary\_Stabilization} [ \text{Using Data Specification Data Spec Name} ][ \text{Mu0} = \text{mu0} | \text{Multiplier} = \text{multiplier} ]
```

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Summary: Momentum capillary stabilization flux BP BC

### 9.2.229 Non Ibp Pressure

**Scope:** Equation System
Non Ibp Pressure Flux For Momentum \{of\{species\subindex\} Species \{in\material\_phase\}
MaterialPhase \}@\at\for\in\on\over Mesh Extent Name = Non_Ibp_Pressure [ Using Data Specification
Data Spec Name ][ Pressure = pressure ]

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9.2.230  Rt Pressure Darcy

Scope:  Equation System

Rt Pressure Darcy Flux For Momentum \{of\{species\subindex\} Species \{in\material\_phase\}
MaterialPhase \}@\at\for\in\on\over Mesh Extent Name = Rt_Pressure_Darcy [ Using Data Specification
Data Spec Name ][ P = p | C_T = c_t | C_X = c_x | C_Y = c_y | C_Z = c_z ]

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9.2.231  Pressure Darcy

Scope:  Equation System

Pressure Darcy Flux For Momentum \{of\{species\subindex\} Species \{in\material\_phase\}
MaterialPhase \}@\at\for\in\on\over Mesh Extent Name = Pressure_Darcy [ Using Data Specification
Data Spec Name ][ Use_Bulk_Node = use_bulk_node | P = p | C_T = c_t | C_X = c_x | C_Y = c_y
| C_Z = c_z | C_Cos_W = c_cos_w | C_Sin_W = c_sin_w | C1_Cos = c1_cos | C1_Sin = c1_sin ]

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9.2.232 Constant Traction

Scope:  Equation System

Constant Traction Flux For Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@at|for|in|on|over|Mesh Extent Name = Constant_Traction [ Using Data Specification Data Spec Name ][ X = x |Y = y |Z = z ]

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</table>

9.2.233 Darcy Slip

Scope:  Equation System

Darcy Slip Flux For Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@at|for|in|on|over|Mesh Extent Name = Darcy_Slip [ Using Data Specification Data Spec Name ][ Vs_X = vs_x |Vs_Y = vs_y |Vs_Z = vs_z |K = k ]

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Summary  Darcy slip momentum flux
9.2.234 Electric Traction

Scope: Equation System

Electric Traction Flux For Momentum [ \{of\} Species \{of\} subindex Species | \{in\} MaterialPhase MaterialPhase ]@\{at\} for\{in\} on\{over\} \{Mesh Extent Name\} = Electric_Traction [ Using Data Specification Data Spec Name ][ \{ Sign \} = sign ]

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Summary Electric traction momentum flux

9.2.235 Projected Capillary

Scope: Equation System

Projected Capillary Flux For Momentum [ \{of\} Species \{of\} subindex Species | \{in\} MaterialPhase MaterialPhase ]@\{at\} for\{in\} on\{over\} \{Mesh Extent Name\} = Projected_Capillary [ Using Data Specification Data Spec Name ][ ]

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Summary Projected capillary momentum flux

9.2.236 Flow Hydrostatic

Scope: Equation System

Flow Hydrostatic Flux For Momentum [ \{of\} Species \{of\} subindex Species | \{in\} MaterialPhase MaterialPhase ]@\{at\} for\{in\} on\{over\} \{Mesh Extent Name\} = Flow_Hydrostatic [ Using Data Specification Data Spec Name ][ P\_ref = p\_ref | Gx = gx | Gy = gy | Gz = gz ]

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Summary Flow hydrostatic momentum flux
9.2.237  Ls Oriented Slip
Scope:  Equation System

Ls Oriented Slip Flux For Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@at|for|in|on|over} Mesh Extent Name = Ls_Oriented_Slip [ Using Data Specification Data Spec Name ][ Vs_X = vs_x |Vs_Y = vs_y |Vs_Z = vs_z |Beta_Normal_A = beta_normal_a | Beta_Tangent_A = beta_tangent_a |Beta_Normal_B = beta_normal_b |Beta_Tangent_B = beta_tangent_b ]

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Summary  Oriented slip momentum flux

9.2.238  Oriented Slip
Scope:  Equation System

Oriented Slip Flux For Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@at|for|in|on|over} Mesh Extent Name = Oriented_Slip [ Using Data Specification Data Spec Name ][ Vs_X = vs_x |Vs_Y = vs_y |Vs_Z = vs_z |Beta_Normal = beta_normal | Beta_Tangent = beta_tangent ]

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Summary  Oriented slip momentum flux

9.2.239  Symmetry Flow
Scope:  Equation System

Symmetry Flow Flux For Cvfem_Solvent_Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@at|for|in|on|over} Mesh Extent Name = Symmetry_Flow [ Using Data Specification Data Spec Name ][ Symmetry_Flow_Name = Symmetry_Flow_Name ]

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Summary  Symmetry flow momentum flux
MaterialPhase ){@|at|for|in|on|over} Mesh Extent Name = Symmetry_Flow [ Using Data Specification Data Spec Name ]

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Summary CVFEM solvent momentum symmetry flow flux BC

9.2.240 Mass Flux
Scope: Equation System

Mass Flux Flux For Cvfem_Mass_Fraction [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Mass_Flux [ Using Data Specification Data Spec Name ][ Total_Mdot = total_mdot |Inflow_Phi = inflow_phi ]

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9.2.241 Open Adv Flow
Scope: Equation System

Open Adv Flow Flux For Cvfem_Mass_Fraction [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Open_Adv_Flow [ Using Data Specification Data Spec Name ][ Pressure = pressure |Total_Pressure = total_pressure |Far_Field_Entrainment_Value = far_field_entrainment_value ]

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9.2.242 Porous Robin Coupled
Scope: Equation System

Porous Robin Coupled Flux For Cvfem_Mass_Fraction [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Porous_Robin_Coupled
[ Using Data Specification Data Spec Name ][ Mass_Fraction_Field = mass_fraction_field | Mass_Fraction_Diffusive_Flux_Field = mass_fraction_diffusive_flux_field | Averaged_Coeff_Field = averaged_coeff_field ]

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Summary: Boundary condition for loose coupling of scalars to a free fluid region

9.2.243 Porous Robin Coupled One Region

Scope: Equation System

Porous Robin Coupled One Region Flux For Cvfem_Mass_Fraction [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ] @at|for|in|on|over { Mesh Extent Name = Porous_Robin_Coupled_One_Region [ Using Data Specification Data Spec Name ][ ]

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Summary: Boundary condition for loose coupling of scalars to a free fluid region

9.2.244 Open Adv Flow

Scope: Equation System

Open Adv Flow Flux For Cvfem_Solvent_Momentum [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ] @at|for|in|on|over Mesh Extent Name = Open_Adv_Flow [ Using Data Specification Data Spec Name ][ Pressure = pressure | Total_pressure = total_pressure | Coflow_Entrainment_Value_X = coflow_entrainment_value_x | Coflow_Entrainment_Value_Y = coflow_entrainment_value_y | Coflow_Entrainment_Value_Z = coflow_entrainment_value_z ]

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</tbody>
</table>

Summary: CVFEM solvent momentum open advective flow flux BC
9.2.245 Smoothed Capillary

Scope: Equation System

Smoothed Capillary Fluxbp For Momentum [{of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Smoothed_Capillary [ Using Data Specification Data Spec Name ]

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Summary: Momentum capillary flux BP BC

9.2.246 Capillary

Scope: Equation System

Capillary Fluxbp For Momentum [{of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Capillary [ Using Data Specification Data Spec Name ]

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Summary: Momentum capillary flux BP BC

9.2.247 No Slip

Scope: Equation System

No Slip Disting For Momentum [{of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = No_Slip [ Using Data Specification Data Spec Name ] [ V0 = v0 ]

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</table>

Summary: No slip model for momentum distinguishing condition

9.2.248 Electroosmotic Velocity

Scope: Equation System
Electroosmotic Velocity Disting For Momentum [ \{\text{of|species|subindex}\} \text{Species}\ |\{\text{in|material_phase}\} \text{MaterialPhase}\ ](\{@|\text{at|for|in|on|over}\} \text{Mesh Extent Name}\ = \text{Electroosmotic Velocity}\ [\text{Using}\ \text{Data} \ \text{Specification}\ \text{Data Spec Name}]\ [C = c]

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<tbody>
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Summary: Electro-osmotic velocity model for momentum distinguishing condition

9.2.249 Wetting Speed Blake Ls
Scope: Equations System

Wetting Speed Blake Ls Disting For Momentum [ \{\text{of|species|subindex}\} \text{Species}\ |\{\text{in|material_phase}\} \text{MaterialPhase}\ ](\{@|\text{at|for|in|on|over}\} \text{Mesh Extent Name}\ = \text{Wetting_Speed_Blake_Ls}\ [\text{Using}\ \text{Data} \ \text{Specification}\ \text{Data Spec Name}]\ [V_W = v_w \ G = g \ \Theta = \theta \ \text{Width} = width \ \text{Tau} = \tau]

<table>
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Summary: Blake level set wetting speed model for momentum distinguishing condition

9.2.250 Darcy Leak
Scope: Equations System

Darcy Leak Disting For Momentum [ \{\text{of|species|subindex}\} \text{Species}\ |\{\text{in|material_phase}\} \text{MaterialPhase}\ ](\{@|\text{at|for|in|on|over}\} \text{Mesh Extent Name}\ = \text{Darcy_Leak}\ [\text{Using}\ \text{Data} \ \text{Specification}\ \text{Data Spec Name}]\ [P_{\text{Ref}} = p_{\text{ref}} \ L = l \ K = k \ \text{Pressure} = pressure]

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<td>\text{pressure}</td>
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</table>
### 9.2.251 Vector User Function Disting

**Scope:** Equation System

Vector User Function Disting Disting For Momentum [ of | species | subindex | Species | { in | material_phase } | MaterialPhase | ] @ | at | for | in | on | over Mesh Extent Name = Vector_User_Function_Disting 
[ Using Data Specification | Data Spec Name ] | Name_X = name_x | Name_Y = name_y | Name_Z = name_z | X = x | Multiplier = multiplier | Mult = Mult

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**Summary**
Vector user function model for momentum distinguishing condition

### 9.2.252 Inflow

**Scope:** Equation System

Inflow Flux For Cv fem Continuity [ of | species | subindex | Species | { in | material_phase } | MaterialPhase | ] @ | at | for | in | on | over Mesh Extent Name = Inflow 
[ Using Data Specification | Data Spec Name ]

<table>
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<tr>
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</table>

**Summary**
IBP continuity inflow; nodal interp

### 9.2.253 Open Flow

**Scope:** Equation System

Open Flow Flux For Cv fem Continuity [ of | species | subindex | Species | { in | material_phase } | MaterialPhase | ] @ | at | for | in | on | over Mesh Extent Name = Open_Flow 
[ Using Data Specification | Data Spec Name ]

<table>
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<th>Value</th>
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<td>Total_Pressure</td>
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</table>
### Parameter | Value  | Default
--- | --- | ---
Species | string | undefined
MaterialPhase | string | undefined
Mesh Extent Name | string | undefined
Data Spec Name | string | undefined
pressure | real | undefined
total_pressure | real | undefined

**Summary**
Open bc for continuity

#### 9.2.254 Simple Inflow

**Scope:** Equation System

Simple Inflow Flux For Cvfem_Continuity [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ]@at|for|in|on|over Mesh Extent Name = Simple_Inflow [ Using Data Specification Data Spec Name ][ Density = Density | Velocity_X = Velocity_X | Velocity_Y = Velocity_Y | Velocity_Z = Velocity_Z ]

### Parameter | Value  | Default
--- | --- | ---
Species | string | undefined
MaterialPhase | string | undefined
Mesh Extent Name | string | undefined
Data Spec Name | string | undefined
Density | real | undefined
Velocity_X | real | undefined
Velocity_Y | real | undefined
Velocity_Z | real | undefined

**Summary**
IBP continuity inflow; specified dens and u

#### 9.2.255 Open Adv Pen Flow

**Scope:** Equation System

Open Adv Pen Flow Flux For Cvfem_Level_Set [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ]@at|for|in|on|over Mesh Extent Name = Open_Adv_Pen_Flow [ Using Data Specification Data Spec Name ][ Penalty_Factor = penalty_factor | Far_Field_Entrainment_Value = far_field_entrainment_value ]

### Parameter | Value  | Default
--- | --- | ---
Species | string | undefined
MaterialPhase | string | undefined
Mesh Extent Name | string | undefined
Data Spec Name | string | undefined
penalty_factor | real | undefined
far_field_entrainment_value | real | undefined

#### 9.2.256 Open Nc Adv Flow

**Scope:** Equation System
Open Nc Adv Flow Flux For Cvfem_Level_Set [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Open_Nc_Adv_Flow [ Using Data Specification Data Spec Name ][ Far_Field_Entrainment_Value = far_field_entrainment_value ]

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9.2.257 Open Adv Flow
Scope: Equation System

Open Adv Flow Flux For Cvfem_Level_Set [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Open_Adv_Flow [ Using Data Specification Data Spec Name ][ Pressure = pressure |Total_Pressure = total_pressure |Far_Field_Entrainment_Value = far_field_entrainment_value ]

<table>
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</table>

9.2.258 Latent Heat
Scope: Equation System

Latent Heat Flux For Enthalpy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Latent_Heat [ Using Data Specification Data Spec Name ][ Yinf = Yinf ]

<table>
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Summary Latent heat enthalpy flux

9.2.259 Nat Conv
Scope: Equation System

Nat Conv Flux For Enthalpy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Nat_Conv [ Using Data Specification Data Spec Name}
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<td>t_ref</td>
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</table>

Summary: Natural convection BC for enthalpy flux

### 9.2.260 Simple Interp Tensor

**Scope:** Equation System

Simple Interp Tensor Flux For Cvfem_Lumped_Div_Projection [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Simple_Interp_Tensor [ Using Data Specification Data Spec Name ][ Source = source ]

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Summary: CVFEM lumped divergence projection simple interpolation tensor flux BC

### 9.2.261 Rad

**Scope:** Equation System

Rad Flux For Enthalpy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Rad [ Using Data Specification Data Spec Name ][ T_Ref = t_ref |Crad = crad ]

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</table>

Summary: Radiative energy flux defined with constant models for emissivity, form factor and reference temperature

### 9.2.262 Kinematic

**Scope:** Equation System
Kinematic Rotated For Mesh_Colloc [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Kinematic [ Using Data Specification Data Spec Name ][ VO = v0 |Shared_Normal = shared_normal |No_Mesh_Movement = no_mesh_movement ]

<table>
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Summary  
Kinematic model for rotated collocated mesh boundary condition

9.2.263 Kuntz
Scope:  Equation System

Kuntz Flux For Energy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Kuntz [ Using Data Specification Data Spec Name ][ Power_Output = power_output |Flux_Output = flux_output |Toggle = toggle |Multiplier = multiplier |File = file |Units = units ]

<table>
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Summary  
Interpolated value from tabulated data for the energy flux

9.2.264 Generalized Rad
Scope:  Equation System

Generalized Rad Flux For Energy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Generalized_Rad [ Using Data Specification Data Spec Name ][ Power_Output = power_output |Flux_Output = flux_output |Toggle = toggle |Multiplier = multiplier ]

511
Summary Radiative energy flux, possibly defined with different models for emissivity, form factor and irradiation

9.2.265 Latent Heat

Scope: Equation System

Latent Heat Flux For Energy [{of|species|subindex} Species |{in|material_phase} MaterialPhase ]@{at|for|in|on|over} Mesh Extent Name = Latent_Heat [ Using Data Specification Data Spec Name ] [ Power_Output = power_output | Flux_Output = flux_output | Toggle = toggle | Yinf = Yinf ]

Summary Latent heat energy flux

9.2.266 Sharp Line Weld

Scope: Equation System

Sharp Line Weld Flux For Energy [{of|species|subindex} Species |{in|material_phase} MaterialPhase ]@{at|for|in|on|over} Mesh Extent Name = Sharp_Line_Weld [ Using Data Specification Data Spec Name ] [ Power_Output = power_output | Flux_Output = flux_output | Toggle = toggle | Src_X = src_x | Dir_X = dir_x | Vel_X = vel_x | Src_Y = src_y | Dir_Y = dir_y | Vel_Y = vel_y | Src_Z = src_z | Dir_Z = dir_z | Vel_Z = vel_z | R = r | T_On = t_on | T_Off = t_off | Flux = flux ]
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vel_y | real | undefined
src_z | real | undefined
dir_z | real | undefined
vel_z | real | undefined
r | real | undefined
t_on | real | undefined
t_off | real | undefined
flux | real | undefined

9.2.267 Laser Weld
Scope: Equation System

Laser Weld Flux For Energy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Laser_Weld [ Using Data Specification Data Spec Name ][ Power_Output = power_output | Flux_Output = flux_output | Toggle = toggle | Flux = flux | R = r | Normal_Tolerance = normal_tolerance | Path_Function = path_function ]

Parameter | Value | Default
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dir_z | real | undefined
vel_z | real | undefined
r | real | undefined
t_on | real | undefined
t_off | real | undefined
flux | real | undefined

9.2.268 Rad
Scope: Equation System

Rad Flux For Energy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{at|for|in|on|over} Mesh Extent Name = Rad [ Using Data Specification Data Spec Name ][ Power_Output = power_output | Flux_Output = flux_output | Toggle = toggle | T_Ref = t_ref | Crad = crad ]

513
### 9.2.269 Sharp Spot Weld

**Scope:** Equation System

Radiative energy flux defined with constant models for emissivity, form factor and reference temperature

---

#### Summary

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#### 9.2.269 Sharp Spot Weld

**Equation System**

\[
\text{Sharp Spot Weld Flux For Energy [ \{of|species|subindex\} Species \{in|material_phase\} MaterialPhase ] @ \{at|for|in|on|over\} Mesh Extent Name = Sharp_Spot_Weld [ Using Data Specification Data Spec Name ] [ Power_Output = power_output | Flux_Output = flux_output | Toggle = toggle | Src_X = src_x | Dir_X = dir_x | Src_Y = src_y | Dir_Y = dir_y | Src_Z = src_z | Dir_Z = dir_z | R = r | T_On = t_on | T_Off = t_off | Flux = flux ]
\]

---

### 9.2.270 Vapor Cooling

**Scope:** Equation System

Vapor Cooling Flux For Energy [ \{of|species|subindex\} Species \{in|material_phase\} MaterialPhase ] @ \{at|for|in|on|over\} Mesh Extent Name = Vapor_Cooling [ Using Data Specification Data Spec Name ] [ Power_Output = power_output | Flux_Output = flux_output | Toggle = toggle | Tboil = tboil ]
### 9.2.271 Thermal Latent Heat

**Scope:** Equation System

Thermal Latent Heat Flux For Energy \[ {of|species|subindex} \text{Species} | {in|material\_phase} \text{MaterialPhase} \] \{@\at|for|in|on|over\} \text{Mesh Extent Name} = \text{Thermal\_Latent\_Heat} [ Using Data Specification \text{Data Spec Name} ][ \text{Power\_Output} = \text{power\_output} | \text{Flux\_Output} = \text{flux\_output} | \text{Toggle} = \text{toggle} ]

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**Summary** Vapor cooling energy flux

### 9.2.272 Wall Function From Temperature

**Scope:** Equation System

Wall Function From Temperature Flux For Cvfem\_Energy \[ {of|species|subindex} \text{Species} | {in|material\_phase} \text{MaterialPhase} \] \{@\at|for|in|on|over\} \text{Mesh Extent Name} = \text{Wall\_Function\_From\_Temperature} [ Using Data Specification \text{Data Spec Name} ][ \text{Data} = \text{data} | T = t | \text{Wall\_Friction\_Factor} = \text{wall\_friction\_factor} ]

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**Summary** CVFEM energy wall function from temperature flux BC
### 9.2.273 Wall Function From Cht Temperature

**Scope:** Equation System

Wall Function From Cht Temperature Flux For Cvfem_Energy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ][@|at|for|in|on|over} Mesh Extent Name = Wall_Function_From_Cht_Temperature [ Using Data Specification Data Spec Name ][ Data = data |Wall_Friction_Factor = wall_friction_factor ]

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**Summary**
CVFEM energy wall function from conjugate heat transfer temperature flux BC

### 9.2.274 Simple Interp

**Scope:** Equation System

Simple Interp Flux For Hfem_Bf_Projection [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ][@|at|for|in|on|over} Mesh Extent Name = Simple_Interp [ Using Data Specification Data Spec Name ][ ]

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**Summary**
Simple interp model for BF projection flux

### 9.2.275 Melting

**Scope:** Equation System

Melting Disting For Extension_Speed [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ][@|at|for|in|on|over} Mesh Extent Name = Melting [ Using Data Specification Data Spec Name ][ ]

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**Summary**
Melting model for extension speed distinguishing condition
9.2.276  Recession

Scope:    Equation System

Recession Disting For Extension_Speed [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Recession [ Using Data Specification Data Spec Name ][ P_Ref = p_ref |K = k |K_0 = k_0 |N = n |Pressure = pressure ]

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Summary    Advection model for extension speed distinguishing condition

9.2.277  Advection

Scope:    Equation System

Advection Disting For Extension_Speed [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Advection [ Using Data Specification Data Spec Name ][ ]

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Summary    Advection model for extension speed distinguishing condition

9.2.278  Solidification

Scope:    Equation System

Solidification Disting For Extension_Speed [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Solidification [ Using Data Specification Data Spec Name ][ ]

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Summary    Solidification model for extension speed distinguishing condition
9.2.279  Simple Interp

Scope:  Equation System

Simple Interp Flux For Hfem_Bf_Lumped_Projection [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Simple_Interp [ Using Data Specification Data Spec Name ][ ]

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Summary  Simple interp model for BF projection flux

9.2.280  Open Adv Flow

Scope:  Equation System

Open Adv Flow Flux For Cvfem_Turbulence_Dissipation_Rate [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Open_Adv_Flow [ Using Data Specification Data Spec Name ][ Pressure = pressure |Total_Pressure = total_pressure |Far_Field_Entrainment_Value = far_field_entrainment_value ]

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9.2.281  Mass Flux

Scope:  Equation System

Mass Flux Flux For Cvfem_Turbulence_Dissipation_Rate [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Mass_Flux [ Using Data Specification Data Spec Name ][ Total_Mdot = total_mdot |Inflow_Phi = inflow_phi ]

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9.2.282  Free Open Flow

Scope:  Equation System

Free Open Flow Flux For Cvfem_Turbulence_Dissipation_Rate [ {of|species|subindex} Species |
{in|material_phase} MaterialPhase ][@|at|for|in|on|over] Mesh Extent Name = Free_Open_Flow
[ Using Data Specification Data Spec Name ][ ]

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9.2.283  Wall Function

Scope:  Equation System

Wall Function Disting For Cvfem_Turbulent_Kinetic_Energy [ {of|species|subindex} Species |
{in|material_phase} MaterialPhase ][@|at|for|in|on|over] Mesh Extent Name = Wall_Function
[ Using Data Specification Data Spec Name ][ Wall_Friction_Factor = Wall_Friction_Factor |
Wall_Velocity_X = Wall_Velocity_X | Wall_Velocity_Y = Wall_Velocity_Y | Wall_Velocity_Z = Wall_Velocity_Z ]

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Summary  CVFEM wall function turbulent kinetic energy distinguishing condition

9.2.284  Equilibrium Capillary Gap

Scope:  Equation System

Equilibrium Capillary Gap Flux For Momentum [ {of|species|subindex} Species |
{in|material_phase} MaterialPhase ][@|at|for|in|on|over] Mesh Extent Name = Equilibrium_Capillary_Gap
[ Using Data Specification Data Spec Name ][ Theta = theta ]

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Summary  Equilibrium gap capillary momentum flux
9.2.285 Pspg
Scope: Equation System

Pspg Fluxbp For Continuity [ \{of\{species\{subindex\}\}\} Species \{in\{material\_phase\}\} MaterialPhase ] \{\@\at\for\in\on\over\} Mesh Extent Name = Pspg [ Using Data Specification Data Spec Name ] [ Tausurfacefactor = TauSurfaceFactor | Tausurfacejacobianfactor = TauSurfaceJacobianFactor ]

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Summary Continuity PSPG flux BP BC

9.2.286 Kinematic
Scope: Equation System

Kinematic Rotated For Momentum_Colloc [ \{of\{species\{subindex\}\}\} Species \{in\{material\_phase\}\} MaterialPhase ] \{\@\at\for\in\on\over\} Mesh Extent Name = Kinematic [ Using Data Specification Data Spec Name ] [ V0 = v0 | Shared_Normal = shared_normal | No_Mesh_Movement = no_mesh_movement ]

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Summary Kinematic model for rotated collocated momentum boundary condition

9.2.287 Open Flow Convection
Scope: Equation System

Open Flow Convection Flux For Mixture_Fraction [ \{of\{species\{subindex\}\}\} Species \{in\{material\_phase\}\} MaterialPhase ] \{\@\at\for\in\on\over\} Mesh Extent Name = Open_Flow_Convection [ Using Data Specification Data Spec Name ] [ Reference_Value = Reference_Value ]

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9.2.288  Simple Inflow

Scope:  Equation System

Simple Inflow Flux For Hfem_Continuity [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ][@|at|for|in|on|over] Mesh Extent Name = Simple_Inflow [ Using Data Specification Data Spec Name ][ Density = Density |Velocity_X = Velocity_X |Velocity_Y = Velocity_Y |Velocity_Z = Velocity_Z ]

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</table>

Summary  IBP continuity inflow; specified dens and u

9.2.289  Interface

Scope:  Equation System

Interface Flux For Hfem_Continuity [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ][@|at|for|in|on|over] Mesh Extent Name = Interface [ Using Data Specification Data Spec Name ][ ]

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Summary  INTERFACE HFEM continuity flux

9.2.290  Open

Scope:  Equation System

Open Flux For Hfem_Lumped_Projection [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ][@|at|for|in|on|over] Mesh Extent Name = Open [ Using Data Specification Data Spec Name ][ Total_Pressure = total_pressure |Pressure = pressure ]
### 9.2.291 Simple Interp

**Scope:** Equation System

Simple Interp Flux For Hfem_Lumped_Projection [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Simple_Interp [ Using Data Specification Data Spec Name ]

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**Summary**

Simple interp model for projection flux

### 9.2.292 Generalized Nat Conv

**Scope:** Equation System

Generalized Nat Conv Flux For Cvem_Energy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Generalized_Nat_Conv [ Using Data Specification Data Spec Name ]

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**Summary**

CVFEM energy generalized natural convection flux BC

### 9.2.293 Simple Interp

**Scope:** Equation System

Simple Interp Flux For Darcy_Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Simple_Interp [ Using Data Specification Data Spec Name ]

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### Summary

Darcy momentum simple interpolation flux BC

#### 9.2.294 Simple Interp

**Scope:** Equation System

**Parameter**
- **Species:** string undefined
- **MaterialPhase:** string undefined
- **Mesh Extent Name:** string undefined
- **Data Spec Name:** string undefined

**Summary**
CVFEM edge-based projection simple interpolation flux BC

#### 9.2.295 Simple Dp Interp

**Scope:** Equation System

**Parameter**
- **Species:** string undefined
- **MaterialPhase:** string undefined
- **Mesh Extent Name:** string undefined
- **Data Spec Name:** string undefined

**Summary**
Simple DP interpolation model for HFEM velocity pressure projection flux

#### 9.2.296 Kinematic

**Scope:** Equation System

**Parameter**
- **Species:** string undefined
- **MaterialPhase:** string undefined
- **Mesh Extent Name:** string undefined
- **Data Spec Name:** string undefined

**Summary**
Kinematic Distinct For Mesh_Y [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]{@|at|for|in|on|over} Mesh Extent Name = Kinematic [ Using Data Specification Data Spec Name ][ V0 = v0 |Shared_Normal = shared_normal |No_Mesh_Movement = no_mesh_movement ]
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Summary: Kinematic model for mesh_y distinguishing condition

### 9.2.297 Recession
Scope: Equation System

Recession Distinguishing For Mesh_Y [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ] {at|for|in|on|over} Mesh Extent Name = Recession [ Using Data Specification Data Spec Name ] [ P_Ref = p_ref | K = k | K_0 = k_0 | N = n | Pressure = pressure ]

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Summary: Recession model for mesh_x distinguishing condition

### 9.2.298 Well Outflow
Scope: Equation System

Well Outflow Flux For Mass_Balance [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ] {at|for|in|on|over} Mesh Extent Name = Well_Outflow [ Using Data Specification Data Spec Name ] [ Pi = pi | Pref = pref ]

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### 9.2.299 Ls Capillary
Scope: Equation System
Ls Capillary Fluxbp For Hfem_Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ] @ at for in on over Mesh Extent Name = Ls_Capillary [ Using Data Specification Data Spec Name ] [ ]

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Summary: Momentum capillary flux BP BC using level set normal

9.2.300 Capillary
Scope: Equation System

Capillary Fluxbp For Hfem_Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ] @ at for in on over Mesh Extent Name = Capillary [ Using Data Specification Data Spec Name ] [ ]

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Summary: Momentum capillary flux BP BC

9.2.301 Interface
Scope: Equation System

Interface Flux For Hfem_Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ] @ at for in on over Mesh Extent Name = Interface [ Using Data Specification Data Spec Name ] [ ]

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Summary: INTERFACE HFEM momentum flux

9.2.302 Symmetry Flow
Scope: Equation System

Symmetry Flow Flux For Hfem_Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ] @ at for in on over Mesh Extent Name = Symmetry_Flow [ Using Data Specification Data Spec Name ] [ ]

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### 9.2.303 Open Adv Flow

**Scope:** Equation System

Open Adv Flow Flux For Hfem_Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ] @at|for|in|on|over} Mesh Extent Name = Open_Adv_Flow [ Using Data Specification Data Spec Name ][ Pressure = pressure |Total_Pressure = total_pressure |Coflow_Entrainment_Value_X = coflow_entrainment_value_x |Coflow_Entrainment_Value_Y = coflow_entrainment_value_y |Coflow_Entrainment_Value_Z = coflow_entrainment_value_z ]

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**Summary**
HFEM momentum open advective flow flux BC

### 9.2.304 Balanced Force

**Scope:** Equation System

Balanced Force Flux For Hfem_Momentum [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ] @at|for|in|on|over} Mesh Extent Name = Balanced_Force [ Using Data Specification Data Spec Name ][ ]

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**Summary**
HFEM momentum balanced force flux BC

### 9.2.305 Mass Fluid Robin Coupled

**Scope:** Equation System

Mass Fluid Robin Coupled

Summary: HFEM momentum symmetry flow flux BC

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Mass Fluid Robin Coupled Flux For Mass_Balance [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ] @ at | for | in | on | over} Mesh Extent Name = Mass_Fluid_Robin_Coupled [ Using Data Specification Data Spec Name ] [ Coeff_Scaling = coeff_scaling | P_Field = P_Field | MassFlux_Field = MassFlux_Field ]

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Summary: Boundary condition for loose coupling of continuity to a free fluid region

9.2.306 Mass Fraction Fluid Robin Coupled One Region
Scope: Equation System

Mass Fraction Fluid Robin Coupled One Region Flux For Mass_Balance [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ] @ at | for | in | on | over} Mesh Extent Name = Mass_Fraction_Fluid_Robin_Coupled One Region [ Using Data Specification Data Spec Name ]

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Summary: Boundary condition for coupling of scalars to a free fluid region in the same region.

9.2.307 Mass Fraction Fluid Robin Coupled
Scope: Equation System

Mass Fraction Fluid Robin Coupled Flux For Mass_Balance [ {of|species|subindex} Species | {in|material_phase} MaterialPhase ] @ at | for | in | on | over} Mesh Extent Name = Mass_Fraction_Fluid_Robin_Coupled [ Using Data Specification Data Spec Name ] [ Mass_Fraction_Field = mass_fraction_field | Mass_Fraction_Diffusive_Flux_Field = mass_fraction_diffusive_flux_field | Coeff_Field = coeff_field ]

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Summary: Boundary condition for loose coupling of scalars to a free fluid region
9.2.308 Generalized Rad

Scope: Equation System

Generalized Rad Flux For Porous_Enthalpy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Generalized_Rad [ Using Data Specification Data Spec Name ][ Multiplier = multiplier ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>MaterialPhase</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Mesh Extent Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Data Spec Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>multiplier</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Radiative porous enthalpy flux, possibly defined with different models for emissivity, form factor and irradiation

9.2.309 Enclosure Radiation

Scope: Equation System

Enclosure Radiation Flux For Porous_Enthalpy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Enclosure_Radiation [ Using Data Specification Data Spec Name ][ Ndof = ndof |Multiplier = multiplier |Enclosure = enclosure ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>MaterialPhase</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Mesh Extent Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Data Spec Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>ndof</td>
<td>integer</td>
<td>undefined</td>
</tr>
<tr>
<td>multiplier</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>enclosure</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

9.2.310 Fluid Robin Coupled With Solid Convection One Region

Scope: Equation System

Fluid Robin Coupled With Solid Convection One Region Flux For Porous_Enthalpy [ {of|species|subindex} Species |{in|material_phase} MaterialPhase ]@|at|for|in|on|over} Mesh Extent Name = Fluid_Robin_Coupled_With_Solid_Convection_One_Region [ Using Data Specification Data Spec Name ][ ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>MaterialPhase</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Mesh Extent Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Data Spec Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Boundary condition for loose coupling of enthalpy to a free fluid region
9.2.311 Fluid Solid Convection Coupled One Region

**Scope:** Equation System

Fluid Solid Convection Coupled One Region Flux For Porous_Enthalpy [{of|species|subindex} Species |{in|material_phase} MaterialPhase }{at|for|in|on|over} Mesh Extent Name = Fluid_Solid_Convection_Coupled_One_Region [ Using Data Specification Data Spec Name ] [ Multiplier = multiplier ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>MaterialPhase</td>
<td>string</td>
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<td>Mesh Extent Name</td>
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<td>undefined</td>
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<tr>
<td>Data Spec Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>multiplier</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Boundary condition for loose coupling of enthalpy to a free fluid region

9.2.312 Bc Bulk Node Coupling For Density = K Factor Flow

**Scope:** Equation System

9.2.313 Bc Bulk Node Coupling For Density = K Factor Flow With Choking

**Scope:** Equation System

Bc Bulk Node Coupling For Density = K Factor Flow With Choking [ Critical_Pressure_Ratio = critical_pressure_ratio ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>critical_pressure_ratio</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

9.2.314 Bc Bulk Node Coupling For Species = K Factor Flow

**Scope:** Equation System

9.2.315 Bc Bulk Node Coupling For Species = K Factor Flow With Choking

**Scope:** Equation System

Bc Bulk Node Coupling For Species = K Factor Flow With Choking [ Critical_Pressure_Ratio = critical_pressure_ratio ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>critical_pressure_ratio</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

9.2.316 Bc Interface Colloc Disting For Lubrication Height = Deforming

**Scope:** Equation System

Bc Interface Colloc Disting For Lubrication Height = Deforming [ Hmin = hmin ]
Parameter | Value | Default
--- | --- | ---
hmin | real | undefined

Summary
Collocated distinguishing condition for the lubrication height that deforms by the displacement of the adjoining volume region.

### 9.3 Periodic BC Overview

Aria provides a periodic boundary condition capability. The underlying implementation is based upon augmenting the basic governing equations with multi-point constraints on nodal DOF at the periodic boundaries. The implementation is equally applicable to scalar and vector DOF. Since the enforcement employs nodal DOF, the capability is limited to grids with conforming surface nodes, i.e. the mesh is periodic as well.

### 9.4 Coupling BCs for Organic Material Decomposition

Aria provides the capability of coupling between porous (see section 5.11.2) and fluid blocks using flux-style boundary conditions. The boundary conditions described here are for a single region computation; for the most part, these boundary conditions can be used for a multi-region computation with the phrase ‘one_region’ omitted. These boundary conditions are imposed as Robin boundary conditions,

\[
\begin{align*}
F_P &= \rho_F u_F \cdot n + \beta(p_g - p_F) \\
F_F &= \rho_g u_g \cdot n + \beta(p_F - p_g)
\end{align*}
\]  

where \( F_P \) is the flux into the porous region and \( F_F \) is the flux into the fluid region. The boundary conditions for the pressure mass balance equation on the porous side and the corresponding flux condition on the fluid side CVFEM continuity equation are written as

BC flux for mass_balance in gas_phase on surface = fluid_robin_coupled_one_region
BC flux for cvfem_continuity on surface = porous_robin_one_region

This boundary condition uses a penalty coefficient of the form

\[
\beta = \frac{c_{scaling} \rho_g K}{\mu_g h}
\]

(9.42)

where \( h \) is the mesh width adjacent to the interface (need to check what happens with a non-uniform mesh). The default value of \( c_{scaling} \) is 0.001. If this parameter is too large, the increase in pressure might be underpredicted.

A distinguishing boundary condition is used on the fluid momentum equation,

\[
u_j - (u_{j,n}^D + u_{j,t}^D) = 0,
\]

(9.43)

where \( u_{j,n}^D \) is the imposed normal component of velocity and \( u_{j,t}^D \) is the imposed tangential component of velocity. The normal component is computed directly from the continuity flux at the interface,

\[
u_{j,n}^D = \frac{F_F}{\rho_F} n_j.
\]

(9.44)
The tangential component is based on a variation of the classical Beavers-Joseph-Saffman condition [21, 22] for the slip velocity which has been extended to non-planar surfaces in multidimensional flow [23], which defines a provisional model velocity

\[ u_{j}^{\text{BJS}} = -\sqrt{\frac{\bar{K}}{\alpha\mu}} (n_i \tau_{ij}) \]  

(9.45)

where \( \bar{K} \) is the permeability of the porous region at the interface, \( \mu \) is the viscosity of the local fluid at the interface, \( \tau_{ij} \) is the viscous stress tensor of the fluid at the interface, and \( \alpha \) is a dimensionless model parameter that is a function of the microstructure of the porous material, which has been found to have typical values near 0.1 [22]. The tangential component of this vector quantity is used as the tangential component of the distinguishing condition velocity, and is computed as

\[ u_{j,t}^{D} = u_{j}^{\text{BJS}} - (u_{k}^{\text{BJS}} n_k) n_j. \]  

(9.46)

which is written in the input file as

BC Distinct for cvfem_momentum on surface = porous_robin_one_region

The flux of a species \( k \) across a porous-fluid interface is

\[ J_{P}^{Y} = J_{F}^{\text{diff}} + P \rho g Y_{k,PF} + \left( \frac{D_k \rho_f}{h} \right) (Y_{k,P} - Y_{k,F}) \]

(9.47)

where \( P \) refers to porous side, \( F \) refers to the fluid side and \( Y_{k,PF} \) is the upwinded interface mass fraction, equivalent to \( h_{PF} \) from the enthalpy coupling. The mass fraction coupling boundary condition on the porous side is written as

BC flux for mass_balance in phase on surface = fluid_robin_coupled_one_region

The enthalpy coupling boundary conditions in the porous region are

BC flux for porous_enthalpy in phase on surface = fluid_robin_coupled_with_solid_convection_one_region
BC flux for porous_enthalpy in phase on surface = fluid_solid_convection_coupled_one_region
Chapter 10

Distinguishing Conditions

10.1 An Introduction to Distinguishing Conditions

Aria’s distinguishing condition (DC) feature is an essential ingredient in solving many coupled physics problems. A distinguishing condition is really just another equation specification except that it typically replaces a regular equation on a subset of the domain such as a surface.

For example, in solving fluids problems with a free surface where the mesh boundary moves with the material, e.g., an ALE simulation, a kinematic condition is used to tie the mesh to the fluid on the free boundary. In this example, one of the mesh coordinates, say \( \text{MESH\_DISPLACEMENTS\_X} \), is unknown. So, the equation that is normally used to solve for that component (the \( x \)-component of the \( \text{MESH} \) equation) is replaced with the kinematic condition:

\[
\mathbf{n} \cdot \left( \mathbf{v} - \dot{\mathbf{d}} \right) - v_o = 0.
\]

An additional feature of distinguishing conditions is that multiple DCs for a given degree of freedom on a given surface are added together. This additive feature allows users to build up their own conditions from primitive ones.

An important thing to know about these conditions is that they are satisfied weakly. That is, the DC is multiplied by a finite element weight function and integrated over the surface. Consequently, the condition is only satisfied weakly and to within the tolerance of the nonlinear solver.

The remainder of this chapter contains a description of the primitive DCs that are available in Aria. Using the user plugin feature described in chapter 18 users can add their own, more complicated or specialized conditions. Input specification of distinguishing conditions on portions of the FEM mesh can be simplified by aggregation of mesh parts into Mesh Groups 6.26.

10.2 BC DISTING

<table>
<thead>
<tr>
<th>BC DISTING for</th>
<th>Description</th>
<th>Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>( E\text{QUANE} ) ____&lt;Subindex&gt;\____&lt;Phase&gt;</td>
<td>Replaces the equations for ( E\text{QUANE} ) on ( M\text{ESH_PART} ) with a distinguishing condition implemented by ( \text{MODEL} ).</td>
<td>Prior to the assembly of the distinguishing conditions, the “normal” matrix and RHS entries are zeroed. So, these conditions do not rely on penalty parameters. Also, if multiple distinguishing conditions are supplied they are added together so the sum of the conditions is satisfied – that is, they are not individually satisfied. This allows you to construct complex expressions based on the available primitives and/or any plugins. See the ( \text{POLYNOMIAL} ) model below for an example combining two distinguishing conditions. Finally, it is worth noting</td>
</tr>
</tbody>
</table>
that these conditions are satisfied weakly and so they are only satisfied to within the tolerance of the nonlinear solver.

### 10.2.1 DARCY_LEAK

**Description**  This model implements a vector distinguishing condition that allows fluid to “leak” through a boundary according to a Darcy’s Law like relationship:

\[
v - \dot{x} \cdot nn - \frac{K}{L \mu} (P - P_{\text{ref}}) n = 0. \tag{10.1}
\]

the velocity, \( \dot{x} \) is the time derivative of the mesh boundary position and \( P \) is the fluid pressure. \( K \) is the permeability of the boundary, \( \mu \) is the viscosity, \( P_{\text{ref}} \) is a reference pressure (defaults to zero) and \( L \) is a length scale for the pressure drop (defaults to unity).

Note that the second term in 10.1 ensures that the resulting velocity is relative to any boundary motion that arises in moving mesh problems. For problems on fixed meshes this term is automatically excluded.

An alternate pressure field can be used by provided the optional parameter `PRESSURE`.

**Parameters**  
\[
K = \text{REAL} \\
[P_{\text{REF}} = \text{REAL}] \\
[L = \text{REAL}] \\
[PRESSURE = \text{STRING}]
\]

**Example**  
BC Disting For Momentum_A On surface_2 = Darcy_Leak K=1
BC Disting For Momentum_B On surface_2 = Darcy_Leak K=1e-7

### 10.2.2 KINEMATIC

**Description**  This model implements the kinematic condition of the form

\[
n \cdot (v - \dot{x}) - v_0 = 0 \tag{10.2}
\]

where \( n \) is the outward unit normal to the boundary, \( v \) is the velocity, \( \dot{x} \) is the time derivative of the mesh boundary position and \( v_0 \) is the mass flux per unit mass across the interface, viz. a “leak” velocity. The leak velocity can be supplied by the `V0` parameter which defaults to zero.

**Parameters**  
\[
V_0 = \text{REAL}
\]

**Example**  
BC Disting For Mesh_X On surface_2 = Kinematic V0=0

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10.2.3 POLYNOMIAL

Parameters
- VARIABLE = STRING
- ORDER = INT
- [C0 = REAL]
- [C1 = REAL]
- ...
- [CN = REAL]

Example
BC Disting For Mesh_Y On surface_2 = Polynomial Variable=Time Order=1 C1=0.5
BC Disting For Mesh_Y On surface_2 = Polynomial Variable=Mesh_Dispacement_Y Order=1 C1=-1

Description
Arbitrary order polynomial function of a specified scalar variable.

\[ \sum_{i=0}^{N} C_i X^i = 0 \] (10.3)

Here, \( N \) is the order of the polynomial provided by the ORDER parameter and \( X \) is the variable supplied by the VARIABLE parameter and \( C_i \) are the supplied coefficients. Coefficients that are not supplied default to a value of zero. The VARIABLE argument can be TIME or any internal Expression that evaluates to a scalar. For the latter case, the format of the VARIABLE argument is described in section 2.6.1.

In the example given above, two distinguishing conditions are combined to give a composite function. In that example, the resulting conditions is

\[ \frac{1}{2} t - d_y = 0 \] (10.4)

or

\[ d_y = \frac{1}{2} t \] (10.5)

10.2.4 WETTING_SPEED_BLAKE_LS

Parameters
- V_W = REAL
- G = REAL
- THETA = REAL
- WIDTH = REAL
- [TAU = REAL]

Example
BC Disting for Momentum_A on surface_3 = Wetting_Speed_Blake_LS V_w=1e-1 g=1 Width=1 Theta=60

Description

\[ v - \delta(F) v_w \sinh \left( g \left( \cos \theta_s - \cos \theta \right) \right) t_w + \delta(F) \tau \dot{v} = 0 \] (10.6)

where \( v \) is the fluid velocity and \( t_w \) is the tangent to the wall. The function \( \delta(F) \), where \( F \) is the level set distance field, is given by

\[ \delta(F) = \frac{1}{2} \left( 1 + \cos \frac{\pi F}{\alpha} \right) \] (10.7)
when \(|F| < \alpha\) and zero elsewhere. Here, \(\alpha\) is the half of the \texttt{WIDTH} parameter. The term involving \(\tau\) is a transient relaxation term. By default, \(\tau = 0\).

This distinguishing condition is a function of the static and observed contact angles. The static contact angle \(\theta_s\), supplied by the \texttt{THETA} parameter, is fixed. The observed contact angle \(\theta\) is computed from the current state of the solution as illustrated in the following diagram. The important point here is that the contact angle is measured through the negative side of the distance function which is denoted \texttt{PHASE_A} in Aria.

10.2.5 \textbf{ELECTROOSMOTIC_VELOCITY}

\begin{itemize}
  \item \textbf{Parameters} \hspace{1cm} \(C = \text{REAL}\)
  \item \textbf{Example} \hspace{1cm} \texttt{BC Disting for Momentum on surface_3 = Electroosmotic_Velocity C=0.3}
  \item \textbf{Description} \hspace{1cm} This distinguishing condition sets the velocity to simply be a constant multiple of the electric field,
    \[ \nu - cE = 0 \] \hspace{1cm} (10.8)
    where \(\nu\) is the fluid velocity, \(E\) is the electric field and \(c\) is the constant \(C\) supplied by the user.
\end{itemize}
Chapter 11

Volumetric Sources

This chapter documents the Source line commands within the current version of Aria. The Source line defines a model for a volumetric source term for a given equation 6. It is important to note that with the exception of sources defined in Volume Source command blocks for thermal problems 33.1 the source term must be explicitly defined with a Source command line. For both the single line and command block syntax, input specification of volumetric sources on portions of the FEM mesh can be simplified by aggregation of element blocks into Mesh Groups 6.26.

Source command lines appear in the Region scope of the input file as illustrated below.

```plaintext
Begin Procedure My_Aria_Procedure
.
.
Begin Aria Region My_Region
.
.
EQ Energy for Temperature on block_2 with Q1 using DIFF SRC
Source for energy on block_2 = constant value = 0.2
.
.
End
.
.
End
.
.
```

### 11.1 POINT SOURCE FOR...

| POINT SOURCE FOR `eqname` ON `mesh_part` VALUE = `q` `x = x` `y = y` `z = z` |
| Description | Arbitrary point source contributions. |
| Summary | This adds a point source with value `q` at the specified position. Currently limited to constant and scalar sources. Only supported by the voltage equation (though extending it to other equations is simple, just ask). This line command syntax needs to change since the `ON mesh_part` piece doesn’t really make sense since you already supply the coordinates. You have to supply as many coordinate positions as the problem has dimensions. |
Mathematically, the point source is represented as

\[ q = Q\delta(x - X) \]  \hspace{1cm} (11.1)

where \( \delta() \) is the Dirac delta function which is zero everywhere except at the point \( x = X \) where \( x \) is the physical coordinate and \( X \) is the position provided via the input. In finite elements, this source is integrated

\[ \int_V Q\delta(x - X)\phi^i \, dV = Q\phi^i(X) \in \text{Elem}_X. \]  \hspace{1cm} (11.2)

Here we employ the local support of the basis functions \( \phi \) so that this is only evaluated in the elements containing the point \( X, \text{Elem}_X \).

Example POINT SOURCE FOR Voltage ON block_1 Value = 1 X = 0 Y = 0 Z = 0

### 11.2 VECTOR POINT SOURCE FOR ...

VECTOR POINT SOURCE FOR \( \text{EQNAME} \) ON \( \text{MESH} \_\text{PART} \) V_X=REAL V_Y=REAL V_Z=REAL at X=REAL Y=REAL Z=REAL

Description Arbitrary vector point source contribution.

Summary This adds a vector point source with value \((v_x, v_y, v_z)\) at the specified position. Currently limited to a constant value.

Mathematically, the point source is represented as

\[ q = Q\delta(x - X) \]  \hspace{1cm} (11.3)

where \( \delta() \) is the Dirac delta function which is zero everywhere except at the point \( x = X \) where \( x \) is the physical coordinate and \( X \) is the position provided via the input. In finite elements, this source is integrated

\[ \int_V Q\delta(x - X)\phi^i \, dV = Q\phi^i(X) \in \text{Elem}_X. \]  \hspace{1cm} (11.4)

Here we employ the local support of the basis functions \( \phi \) so that this is only evaluated in the elements containing the point \( X, \text{Elem}_X \).

Example VECTOR POINT SOURCE FOR Momentum ON block_1 V_X=1 V_Y=1 V_Z=1 at X=0 Y=0 Z=0
11.3 SOURCE FOR ENERGY

SOURCE FOR ENERGY ON MESH_PART = MODEL[param1 = val1, param2 = val2 ...]

Description  Arbitrary source contributions for the energy equation.

Summary  Adds the source provided by MODEL to the energy equation.

Parent Block(s)  ARIA_REGION

11.3.1 SOURCE FOR ENERGY = CALORE_USER_SUB

Parameters  
NAME = STRING

TYPE = STRING

[MULTIPLIER = REAL]

[NR = INT]

[R0 = INT]

[R1 = INT(etc.)]

[NI = INT]

[I0 = INT]

[I1 = INT(etc.)]

Example  
Source for Energy on block_10 = Calore_User_Sub name=w80afftuser
type=element NR=2 R0=1000 R1=0.7

Description  
NAME is the name of the user subroutine was registered with. NR and NI are the numbers of real and integer parameters, respectively. Each real (int) parameter is supplied by the Rn (In) parameter. Note: the parameters use zero based counting. The TYPE parameter denotes the API the user subroutine uses; currently Aria only supports types ELEMENT and NODE.

The optional MULTIPLIER parameter (defaults to 1.0) can be supplied to scale the output of the user subroutine, i.e., the flux $f$ will be $f = mf_{user}$ where $m$ is the multiplier and $f_{user}$ is the output of the user subroutine.

See section 9.2.2 for a more complete example of a Calore user subroutine.

11.3.2 SOURCE FOR ENERGY = CHEMEQ_HEATING

Parameters  
MODEL = STRING

[POWER_OUTPUT = STRING]

Example  
Source for Energy on block_10 = CHEMEQ_Heating model = reaction_model
power_output = VAR_NAME

Description  
Defines a heating source caused by chemical reaction. Here MODEL pertains to the paired CHEMEQ MODEL and CHEMEQ SOLVER command blocks that define the chemical system and how it can evolve. Chemical source contributions to the energy equation are further controlled through parameters in the CHEMEQ SOLVER command block.
When the POWER_OUTPUT option is present the energy source will be integrated over the associated element blocks and saved to the user specified global variable, VAR_NAME.

11.3.3 SOURCE FOR ENERGY = MELTING

Parameters

Ts = REAL
Tl = REAL
[latent_heat = REAL]

Example

Source for Energy on block_1 = Melting Ts = 400 Tl = 410 latent_heat = 1e5
Source for Energy on block_1 = Melting Ts = 700 Tl = 725 latent_heat = 1e8
Source for Energy on block_1 = Melting Ts = 900 Tl = 950

Description
Defines a heating source caused by a phase change. The phase change is modeled using a normal distribution spread over a temperature range defined by Ts and Tl. The distribution is sized so that 99% of the energy release occurs between Ts and Tl. The latent heat associated with the phase change can either be specified in-line with the optional “latent_heat” argument or by specifying the latent heat expression in the material model. If both are provided, the value specified in-line is used. The latent heat should be specified units of energy per mass (e.g. J/kg).

This model is preferable to the “Specific Heat = Use_Phase_Change” model because it does not limit the time step or create a discontinuous specific heat. It is also guaranteed to correctly capture the total energy source or sink regardless of the time step size.

Multiple phase transitions can be specified by simply adding multiple melting sources with different temperatures and latent heats.

11.3.4 SOURCE FOR ENERGY = COMPRESSIVE_WORK

Parameters

[MULTIPLIER = REAL]

Example

Source For Energy on block_1 = Compressive_Work

Description
This source accounts for the heat generation or consumption due to compression:

$$q = -m p : \nabla \cdot v$$

(11.5)

where $p$ is the pressure, $\nabla \cdot v$ is the divergence of the velocity field and $m$ is a multiplier (MULTIPLIER) that defaults to 1.

11.3.5 SOURCE FOR ENERGY = CONSTANT

Parameters

VALUE = REAL

Example

SOURCE FOR Energy ON block_1 = Constant Value=1041.3
11.3.6 SOURCE FOR ENERGY = CURING_FOAM_HEAT_OF_RXN

Parameters

VFRAC_SUBINDEX = INT
EXTENT_SUBINDEX = INT
H_RXN = REAL

Example

Source For Energy on block_1 = Curing_Foam_H of_Rxn Vfrac_Subindex=1 Extent_Subindex=2 H_rxn=250

Description

This source accounts for the heat of reaction of a curing epoxy foam, specifically:

$$q = \rho (1 - \phi) \Delta H_{rxn} \frac{\partial \xi}{\partial t}$$  \hspace{1cm} (11.6)

where $\rho$ is the density of the fluid, $\phi$ is the volume fraction, $\Delta H_{rxn}$ is the heat of reaction and $\xi$ is the extent of reaction.

**NOTE:** The volume fraction is assumed to be a SPECIES field with the subindex provided by the VFRAC_SUBINDEX parameter. Likewise, the extent of reaction field is assumed to be a SPECIES field with the subindex provided by the EXTENT_SUBINDEX parameter.

11.3.7 SOURCE FOR ENERGY = CURING_FOAM_LATENT_HEAT

Parameters

VFRAC_SUBINDEX = INT
H_EVAP = REAL

Example

Source For Energy on block_1 = Curing_Foam_Latent_Heat Vfrac_Subindex=1 H_evap=15

Description

This source accounts for the loss of energy due to evaporation of a curing epoxy foam, specifically:

$$q = \rho H_{evap} \frac{\partial \phi}{\partial t}$$  \hspace{1cm} (11.7)

where $\rho$ is the density of the fluid, $H_{evap}$ is the latent heat of evaporation and $\phi$ is the volume fraction.

**NOTE:** The volume fraction is assumed to be a SPECIES field with the subindex provided by the VFRAC_SUBINDEX parameter.

11.3.8 SOURCE FOR ENERGY = CURING_FOAM_SPECIFIC_HEAT

Parameters

VFRAC_SUBINDEX = INT
[CP_FG = REAL]
[CP_E = REAL]
[PHI_ZERO = REAL]

Example

Source For Energy on block_1 = Curing_Foam_Specific_Heat Vfrac_Subindex=1 Cp_fG=1 Cp_e=1 phi_zero=0.2
This source accounts for the loss of energy due to the variable specific heat for the special case where the specific heat material model is CURING_FOAM. See 4.48.6. Specifically, this source term is

\[
q = -\rho T \left( \frac{\partial C_p}{\partial t} + v \cdot \nabla C_p \right)
\]  
\[
= -\rho T b \left( \frac{\partial \phi}{\partial t} + v \cdot \nabla \phi \right)
\]  

where \( \rho \) is the density of the fluid, \( T \) is the temperature, \( v \) is the velocity, \( \phi \) is the volume fraction and \( b \) is as defined in CURING_FOAM specific heat material model (see 4.48.6).

**NOTE:** The volume fraction is assumed to be a SPECIES field with the subindex provided by the VFRAC_SUBINDEX parameter.

### 11.3.9 SOURCE FOR ENERGY = ENCORE_FUNCTION

**Parameters**

- **NAME = STRING**
  - [EVAL_TYPE = STRING]

**Example**

```plaintext```
SOURCE FOR Energy ON block_1 = Encore_Function Name=My_Function
```

**Description**

Using the Encore library, you can define your own functions in a number of ways, such as analytically in the input file, based on the values of a Sierra field or your own compiled functions. See the Encore documentation for more details.

As an example, this snippet of input can be added to the domain level of your input file:

```plaintext```
Begin String Function My_Primitive
  Value is "200 + 1.0\times t"
End
```

Then, this function can be used as shown in the example above.

The optional argument EVAL_TYPE can be used to select the particular function on the Encore function object. Valid options include VALUE, DOT, GRADIENT, FLUX and STRESS. If this option is not supplied Aria will make a sensible choice depending on the resulting expressions – VALUE for NO_OP expressions, DOT for DT_OP expressions and GRADIENT for GRAD_OP expressions.

### 11.3.10 SOURCE FOR ENERGY = JOULE_HEATING

**Parameters**

- [VOLTAGE_SUBINDEX = INT]

**Example**

```plaintext```
SOURCE FOR Energy ON block_1 = Joule_Heating
```

**Description**

This source term adds the volumetric heat source due to Joule heating, a.k.a., Ohmic heating or resistance heating.

\[
H_V = J \cdot E
\]  

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where \( J \) is the current density vector and \( E \) is the electric field vector.

When thermoelectric effects are negligible volumetric heating is effectively given by (see, Section 5.3)

\[
H_V = J^2 R \\
= (-\sigma_e \nabla V) \cdot (-\sigma_e \nabla V) R \\
= (-\sigma_e \nabla V) \cdot (-\sigma_e \nabla V) \frac{1}{\sigma_e} \\
= \sigma_e (\nabla V \cdot \nabla V)
\]

(11.11)

(11.12)

(11.13)

(11.14)

where \( J \) is the current density, \( R \) is the resistivity, \( \sigma_e = 1/R \) is the electrical conductivity and \( V \) is the voltage.

If thermoelectric effects are significant then a thermoelectric current density 4.6.4 should be used instead

\[
J = -\sigma_e (\nabla V + \alpha \nabla T)
\]

(11.15)

where \( \alpha \) is the Seebeck coefficient and \( T \) is the temperature.

### 11.3.11 SOURCE FOR ENERGY = POLYNOMIAL

#### Parameters

- VARIABLE = STRING
- ORDER = INT
  - [VARIABLE_OFFSET = REAL]
  - [C0 = REAL]
  - [C1 = REAL]
  - ...
  - [CN = REAL]

#### Example

```
SOURCE For Energy on block_1= Polynomial Variable=Temperature Order=1
C0=401.0 C1=88.5
```

#### Description

Arbitrary order polynomial function of a specified scalar variable.

\[
H_V = \sum_{i=0}^{N} C_i (X + X_o)^i
\]

(11.16)

Here, \( N \) is the order of the polynomial provided by the ORDER parameter and \( X \) is the variable supplied by the VARIABLE parameter, \( X_o \) is an optional offset (VARIABLE_OFFSET, default is zero) and \( C_i \) are the supplied coefficients. Coefficients that are not supplied default to a value of zero. The VARIABLE argument can be TIME or any internal Expression that evaluates to a scalar. For the latter case, the format of the VARIABLE argument is described in section 2.6.1.

### 11.3.12 SOURCE FOR ENERGY = TBC_JOULE_HEATING

#### Parameters

- Ua = REAL
- Ub = REAL
- V_NS = INT
- Ti = REAL
- CURRENT_LOAD = REAL
  - [VOLTAGE_SUBINDEX = INT]
Example

Example

Example

Description

This source term adds the volumetric heat source due to Joule heating in a thermal battery cell volumetric heating is given by (see, Section 5.3)

\[ H_V = \left( U_o - V - T_i \frac{\partial U_o}{\partial T} \right) I_o \]  

(11.17)

Here, \( U_o \) is the open circuit potential which is given as a linear function in temperature \( T \) and \( U_a \) and \( U_b \) are the coefficients of that function. \( V \) is the cell potential which is taken as the voltage at the node given in the single-node nodeset number \( V_NS \) and \( I_o \) is cell load.

For more details, see [24].

11.3.13 SOURCE FOR ENERGY = VISCOUS_DISSIPATION

Parameters

[MULTIPLIER = REAL]

Example

Example

Example

Description

This source accounts for the heat generation due to viscous dissipation:

\[ q = m \tau : \nabla v \]  

(11.18)

where \( \tau \) is the viscous stress tensor, \( \nabla v \) is the gradient of the velocity and \( m \) is a multiplier (MULTIPLIER) that defaults to 1. The exact form of \( \tau \) will depend on the MOMENTUM_STRESS model choice(s) for the material.

11.3.14 SOURCE FOR ENERGY = USER_FUNCTION

Parameters

NAME = STRING

X = STRING

Example

Example

Example

Description

Source magnitude is defined by a tabular function of the independent variable \( X \). The user specified tabular function NAME as a function of \( X \) scalar solution variable or time. The \( X \) argument can be TIME or any internal Expression that evaluates to a scalar. For the latter case, the format of the \( X \) argument is described in section 2.6.1. Although described here for the energy equation, this source term may be defined on other scalar equations.

11.3.15 SOURCE FOR ENERGY = ELECTRODE_OBJECT

See 11.5.3 for complete description.
11.4 SOURCE FOR MOMENTUM

SOURCE FOR MOMENTUM ON MESH_PART = MODEL [param_1 = val_1, param_2 = val_2 ...]

Description  Arbitrary source contributions for the momentum equation.

Summary    Adds the source provided by MODEL to the momentum equation.

Parent Block(s)  ARIA_REGION

11.4.1 SOURCE FOR MOMENTUM = CONSTANT_VECTOR

Parameters  
[X = REAL]
[Y = REAL]
[Z = REAL]

Example  SOURCE FOR momentum ON block_1 = CONSTANT_VECTOR Z=-9.8

Description  This source term only applies to the momentum equation though it is automatically applied to the continuity equation in PSPG formulations.

X, Y, Z are the components of the vector source. This vector source is not multiplied by the density.

11.4.2 SOURCE FOR MOMENTUM = HYDROSTATIC

Parameters  
GX = REAL
GY = REAL
GZ = REAL
[REF_DENSITY = REAL]

Example  SOURCE FOR momentum ON block_1 = HYDROSTATIC gx = 0 gy = 0 gz = -980

Description  This source term only applies to the momentum equation though it is automatically applied to the continuity equation in PSPG formulations.

GX, GY, GZ are the components of the gravity vector \( \mathbf{g} \) and REF_DENSITY is an constant, uniform reference density \( \rho_\circ \). With density \( \rho \), the hydrostatic source is defined as \( (\rho - \rho_\circ)\mathbf{g} \). The default value of the reference density is 0.

11.4.3 SOURCE FOR MOMENTUM = POTENTIAL

Parameters  None

Example  SOURCE FOR momentum ON block_1 = POTENTIAL
This source term applies the gradient of the potential field, defined by the POTENTIAL equation, as a momentum source. It is most commonly used when the potential field is defined as the hydrostatic potential, and is used in place of the HYDROSTATIC model discussed above.

### 11.4.4 SOURCE FOR MOMENTUM = ROTATING_BODY_FORCE

**Parameters**

- \( G = \text{REAL} \)
- \( \text{FREQUENCY} = \text{REAL} \)
- \([\text{PHASE_SHIFT} = \text{REAL}]\)
- \([\text{REF_DENSITY} = \text{REAL}]\)

**Example**

Source for Momentum on block_1 = Rotating_Body_Force \( g=9.8 \) frequency=2.5
phase_shift=90

**Description**

This source term only applies to the momentum equation though it is automatically applied to the continuity equation in PSPG formulations.

\( GX, GY, GZ \) are the components of the gravity vector \( g \) and \( \text{REF_DENSITY} \) is an constant, uniform reference density \( \rho_o \). With density \( \rho \), the hydrostatic source is defined as \( (\rho - \rho_o)g \).

The default value of the reference density is 0.

### 11.4.5 SOURCE FOR MOMENTUM = BOUSSINESQ

**Parameters**

- \( \text{TEMP_REF} = \text{REAL} \)
- \( \text{VOL_EXP} = \text{REAL} \)
- \( GX = \text{REAL} \)
- \( GY = \text{REAL} \)
- \( GZ = \text{REAL} \)

**Example**

Source for Momentum on block_1 = BOUSSINESQ vol_exp=0.1 temp_ref=298.15 gx = 0 gy = 0 gz = -980

**Description**

This source term only applies to the momentum equation though it is automatically applied to the continuity equation in PSPG formulations.

\( GX, GY, GZ \) are the components of the gravity vector \( g \). \( \text{VOL_EXP} \) is the volume expansion coefficient \( \alpha \) and \( \text{TEMP_REF} \) is the reference temperature \( T_{ref} \). With density \( \rho \) and temperature \( T \) the Boussinesq source is defined as \( \rho g \alpha (T - T_{ref}) \).

### 11.5 SOURCE FOR CURRENT

SOURCE FOR CURRENT ON mesh_part = model [param1 = val1, param2 = val2 ...]

**Description**

Arbitrary source contributions for the current equation.

**Summary**

Adds the source provided by \textit{model} to the current equation.

**Parent Block(s)**

ARIA_REGION
11.5.1 SOURCE FOR CURRENT = BUTLER_VOLMER_SIMPLE

Parameters

\[ A = \text{REAL} \]
\[ C_A = \text{REAL} \]
\[ C_C = \text{REAL} \]
\[ U = \text{REAL} \]
\[ \text{Sign} = \text{INT} \]
\[ [V1\_\text{SUBINDEX} = \text{INT}] \]
\[ [V2\_\text{SUBINDEX} = \text{INT}] \]

Example

\[
\text{SOURCE FOR CURRENT}_1 \text{ ON block}_1 = \text{Butler_Volmer_Simple} \ A=1.0 \ C_a=1.0 \ C_c=-1.0 \ U=0.2 \ \text{Sign}=-1
\]

\[
\text{SOURCE FOR CURRENT}_2 \text{ ON block}_1 = \text{Butler_Volmer_Simple} \ A=1.0 \ C_a=1.0 \ C_c=-1.0 \ U=0.0 \ \text{Sign}=+1
\]

Description

This model implements a very simple form of the Butler-Volmer reaction kinetics. It is intended for developmental and demonstrational purposes only.

This source term has the following form (see, also, equation 5.35)

\[
R_{V,k} = A \left( e^{c_a(V_1-V_2-U)} - e^{-c_c(V_1-V_2-U)} \right) \quad (11.19)
\]

where \( V_1 \) is the first electric potential field and \( V_2 \) is the second electric potential field. By default \( V_1 \) is \( \text{VOLTAGE}_1 \) and \( V_2 \) is \( \text{VOLTAGE}_2 \) but the subindices may be changed using the optional \( V1\_\text{SUBINDEX} \) and \( V2\_\text{SUBINDEX} \) options.

11.5.2 SOURCE FOR CURRENT = POLYNOMIAL

Parameters

\[ \text{VARIABLE} = \text{STRING} \]
\[ \text{ORDER} = \text{INT} \]
\[ [C0 = \text{REAL}] \]
\[ [C1 = \text{REAL}] \]
\[ \ldots \]
\[ [CN = \text{REAL}] \]

Example

\[
\text{SOURCE For Current on block}_1 = \text{Polynomial} \ \text{Variable}=\text{Temperature} \ \text{Order}=1
\]

\[ C0=401.0 \ C1=88.5 \]

Description

Arbitrary order polynomial function of a specified scalar variable.

\[
H_V = \sum_{i=0}^{N} C_i X^i \quad (11.20)
\]

Here, \( N \) is the order of the polynomial provided by the \text{ORDER} parameter and \( X \) is the variable supplied by the \text{VARIABLE} parameter and \( C_i \) are the supplied coefficients. Coefficients that are not supplied default to a value of zero. The \text{VARIABLE} argument can be \text{TIME} or any internal Expression that evaluates to a scalar. For the latter case, the format of the \text{VARIABLE} argument is described in section 2.6.1.
11.5.3 SOURCE FOR CURRENT = ELECTRODE_OBJECT

Parameters

- electrodeFile = STRING
  
- [F = REAL]

- [kmolConversion = REAL]

- [meterConversion = REAL]

- [JConversion = REAL]

Example

SOURCE FOR CURRENT in SOLID_PHASE ON block_1 = ELECTRODE_OBJECT F=96485
electrodeFile=cathode.inp kmolConversion=1000

Description

This model uses the Cantera Electrode object to compute source terms for battery models. Generally this source term will need to be applied to the following equations:

- CURRENT in SOLID_PHASE
- CURRENT in LIQUID_PHASE
- POROUS_SPECIES of (SPECIES_NAME) in LIQUID_PHASE
- [ENERGY]

electrodeFile specifies the input file that will be used to initialize the Electrode objects. F specifies the value of Faraday’s constant in the unit system of the simulation. kmolConversion specifies the conversion factor from kmol to the unit system of the simulation. meterConversion specifies the conversion factor from meters to the unit system of the simulation. JConversion specifies the conversion factor from Joules to the unit system of the simulation.

11.6 SOURCE FOR SPECIES

SOURCE FOR SPECIES ON MESH_PART = MODEL[param_1 = val_1, param_2 = val_2 ...

Description

Arbitrary source contributions for the species equation.

Summary

Adds the source provided by MODEL to the current equation.

Parent Block(s)

ARIA_REGION

11.6.1 SOURCE FOR SPECIES = CURING_FOAM_EXTENT

Parameters

- K = REAL

- E = REAL

- R = REAL

- N = REAL

Example

Source For Species_2 on block_1 = Curing_Foam_Extent k=1.145e5 E=10
R=8.314472E3 n=1.3
Description
This source accounts for the reaction of a curing epoxy foam, specifically:

\[ q = ke^{E/RT} (1 - \xi)^n \]  \hspace{1cm} (11.21)

where \( T \) is the temperature and \( \xi \) is the extent of reaction.

11.6.2 SOURCE FOR SPECIES = CURING_FOAM_VFRAC

Parameters
- \( A = \text{REAL} \)
- \( B = \text{REAL} \)
- \( C = \text{REAL} \)
- \( T_{\text{BOILING}} = \text{REAL} \)

Example
Source For Species_2 on block_1 = Curing_Foam_Vfrac a=1 b=0 c=2e-3
T_BOILING=473.15

Description
This source accounts for the change in volume fraction with temperature.

\[ q = (a + bT + cT^2) \exp \frac{\partial T}{\partial t} \]  \hspace{1cm} (11.22)

where \( T \) is the temperature. This source term is only active when \( T >= T_{\text{boiling}} \).

11.6.3 SOURCE FOR SPECIES = POLYNOMIAL

Parameters
- \( \text{VARIABLE} = \text{STRING} \)
- \( \text{ORDER} = \text{INT} \)
- \([C_0 = \text{REAL}] \)
- \([C_1 = \text{REAL}] \)
- \( \ldots \)
- \([C_N = \text{REAL}] \)

Example
Source For Species on block_1 = Polynomial Variable=Temperature Order=1
C0=401.0 C1=88.5

Description
Arbitrary order polynomial function of a specified scalar variable.

\[ H_V = \sum_{i=0}^{N} C_i X^i \]  \hspace{1cm} (11.23)

Here, \( N \) is the order of the polynomial provided by the \text{ORDER} parameter and \( X \) is the variable supplied by the \text{VARIABLE} parameter and \( C_i \) are the supplied coefficients. Coefficients that are not supplied default to a value of zero. The \text{VARIABLE} argument can be \text{TIME} or any internal Expression that evaluates to a scalar. For the latter case, the format of the \text{VARIABLE} argument is described in section 2.6.1.

11.7 SOURCE FOR POROUS_SPECIES

SOURCE FOR POROUS_SPECIES ON MESH_PART = MODEL[param1 = val1, param2 = val2 ...]
Description: Arbitrary source contributions for the porous_species equation.

Summary: Adds the source provided by \textit{MODEL} to the current equation.

Parent Block(s): ARIA_REGION

### 11.7.1 SOURCE FOR POROUS_SPECIES = ELECTRODE_OBJECT

See 11.5.3 for complete description.

### 11.8 SOURCE FOR VOLTAGE

\texttt{SOURCE FOR VOLTAGE ON MESH\_PART = MODEL[param_1 = val_1, param_2 = val_2 ...]}

Description: Arbitrary source contributions for the voltage equation.

Summary: Adds the source provided by \textit{MODEL} to the current equation.

Parent Block(s): ARIA_REGION

### 11.8.1 SOURCE FOR VOLTAGE = POLYNOMIAL

**Parameters**

- \texttt{VARIABLE = STRING}
- \texttt{ORDER = INT}
- \texttt{[C0 = REAL]}
- \texttt{[C1 = REAL]}
- \ldots
- \texttt{[CN = REAL]}

**Example**

Source For Voltage on block_1 = Polynomial Variable=Temperature Order=1
C0=401.0 C1=88.5

Description: Arbitrary order polynomial function of a specified scalar variable.

\begin{equation}
H_V = \sum_{i=0}^{N} C_i X^i
\end{equation}

(11.24)

Here, \(N\) is the order of the polynomial provided by the \texttt{ORDER} parameter and \(X\) is the variable supplied by the \texttt{VARIABLE} parameter and \(C_i\) are the supplied coefficients. Coefficients that are not supplied default to a value of zero. The \texttt{VARIABLE} argument can be \texttt{TIME} or any internal Expression that evaluates to a scalar. For the latter case, the format of the \texttt{VARIABLE} argument is described in section 2.6.1.
11.9 SOURCE FOR POTENTIAL

SOURCE FOR POTENTIAL ON \texttt{MESH\_PART} = \texttt{MODEL}[\texttt{param}_1 = \texttt{val}_1, \texttt{param}_2 = \texttt{val}_2 \ldots]

Description Arbitrary source contributions for the potential equation.

Summary Adds the source provided by \texttt{MODEL} to the potential equation.

Parent Block(s) ARIA\_REGION

11.9.1 SOURCE FOR POTENTIAL = HYDROSTATIC

Parameters \begin{align*}
GX &= \texttt{REAL} \\
GY &= \texttt{REAL} \\
GZ &= \texttt{REAL} \\
[\texttt{REF\_DENSITY} &= \texttt{REAL}] 
\end{align*}

Example SOURCE FOR Potential ON block\_1 = HYDROSTATIC gx = 0 gy = 0 gz = -980

Description This source term only applies a hydrostatic potential source term to the POTENTIAL equation, defining the potential. It can then be applied to the momentum equation through a separate potential source term in the momentum equation.

\texttt{GX}, \texttt{GY}, \texttt{GZ} are the components of the gravity vector \textit{g} and \texttt{REF\_DENSITY} is an constant, uniform reference density \textit{\rho}_0. With density \textit{\rho}, the hydrostatic source is defined as \((\textit{\rho} - \textit{\rho}_0)\textit{g}\).

The default value of the reference density is 0.
Chapter 12

Constraint Conditions

12.1 CONSTRRAIN

CONSTRRAIN AVERAGE_VOLUME_FRACTION phi = C_0

Description
Integral constraint for the average particle volume fraction in a SUSPENSION problem.

Summary
Constrains $\phi$ throughout the problem domain to achieve the specified value of average volume particle fraction in accordance with the relation

$$C_0 = \left[ \int_{V_o} dV \right]^{-1} \int_{V_o} \phi dV \quad (12.1)$$

over element blocks where the SUSPENSION equation is defined.

Parent Block(s) ARIA_REGION
Chapter 13

Element Death

Aria employs an element death capability supported by the Sierra Framework. Here the basic Framework capability is used to define conditions under which elements will be excluded from the normal set of calculations. The death capability is implemented in two different flavors, one where elements are explicitly removed (Standard Element Death) and another (CDFEM) in which elements are subdivided in a manner consistent with the element death criterion. Generally speaking there is computational economy in the usage of standard element death over CVFEM death at the expense of resolution in front tracking. With a few exceptions both element death implementations utilize essentially the same command directives.

Usage of the death capability is comprised of two parts, definition of criterion and definition of scope. Each death criterion definition is must be named uniquely (DeathName) and its scope will be a specific set of mesh entities, surfaces or blocks of the input mesh. Element death is a dynamic process in which a variable (DEATH_STATUS) on existing elements is modified to denote whether an element is currently active, DEATH_STATUS=1 or inactive, DEATH_STATUS=0.

Visualization of a model with the "dead" elements removed is made possible by use of the DEATH_STATUS variable. Since DEATH_STATUS=0 denotes dead elements, the DEATH_STATUS variable can then be used in conjunction with any visualization thresholding utility to animate element death process.

During a simulation in which element death is active one can monitor the extent of element death through two values output to the log file, KILLED_ELEMENTS and TOTAL_DEAD_ELEMENTS. KILLED_ELEMENTS reports the number of elements eliminated after completion of the time step while TOTAL_DEAD_ELEMENTS is the cumulative number of elements deleted. Both KILLED_ELEMENTS and TOTAL_DEAD_ELEMENTS are also available for output to the ExodusII database as global variables under the same name (e.g. global variables = killed_elements).

Since the death criteria specification will normally involve an internally stored Field, one must use the complete name of that Field when defining element death. As an example, element death named criteria DEATH_TEMP8 based upon the average temperature solution on BLOCK_3 exceeding 322K might be defined as:

```
Begin Element Death death_temp8
  add volume block_3
  Criterion is Avg nodal value of solution->Temperature > 322.0
End
```

In most cases the element death criteria will be evaluated based upon values computed within Aria. Here the death criteria are evaluated at the end of a solution step. However, it is also possible to evaluate the criteria based upon externally supplied "transfer" variables. In this case one can force the death criteria to be evaluated at the beginning of time step using the TRANSFER ELEMENT DEATH command line.
13.1 Element Death

Scope: Aria Region

---

```
Begin Element Death Death Name

Add Volume Block Name...

Criterion {=|are|is} DeathCriterionType Variable Name DeathCriterionCompareType Threshold Value

End
```

---

Summary Allows the specification of an element death criterion to one or more element volumes. The group of element volumes plus a death criterion make up a "death instance".

13.1.1 Add Volume

Scope: Element Death

---

```
Add Volume Block Name...

Parameter Value Default
Block Name string... undefined
```

---

Summary Specifies the names identifying an element volume to which an element death criterion is to be applied. Line can be repeated as needed to add multiple volumes. A group of element volumes plus a death criterion make up a "death instance".

13.1.2 Criterion

Scope: Element Death

---

```
Criterion {=|are|is} DeathCriterionType Variable Name DeathCriterionCompareType Threshold Value

Parameter Value Default
DeathCriterionType {avg nodal value of|death criterion type undefined|element value of|max nodal value undefined|min nodal value of}
Variable Name string undefined
DeathCriterionCompareType {<|<=|=|>|>=|death criterion compare type undefined}
Threshold Value real undefined
```

---

Summary The death criterion for one or more element volumes. Currently designed to support the specification of a threshold for a scalar variable defined on either a node or element. For nodal variables, various operations such as Min., Max. and Avg. are provided to reduce the nodal values to a single element value. For an element variable no operations are given as there is only one value per element. A group of element volumes plus a death criterion make up a "death instance". Currently only one criterion is allowed per death instance.
13.1.3 Transfer Element Death

Scope:

Summary  Cause execution of element death to occur at beginning of a time step rather than at the end of the step.
Chapter 14

Solution Control Reference

14.1 Overview

Aria uses the solution control (SC) library from the SIERRA Framework to orchestrate execution of simulations. All Aria input files must include a Solution Control Description block in the Procedure section of the input file. This description contains directives for executing either a steady-state (sequential) or transient analysis either of which can include nested nonlinear iteration or subcycling. Within the description one selects a named solution control system where the details of execution are more clearly spelled out. Because there are similarities between the Sequential, Transient, Nonlinear Iteration and Subcycling many operations are shared between these directives. However, each of these segments must be uniquely named internally so they can be properly managed under solution control.

Within each SC system, execution of a problem defined at the Region level corresponds to an Advance directive. Thus a steady-state analysis could conceivably be carried out with a single Advance directive. For transient analysis the system can contain several time blocks, each with a corresponding Advance directive. Examples of different control structures are given below.

An example the solution control command block for steady-state analysis would reflect the structure indicated below:

```plaintext
Begin Sierra myJob
  . Materials, Solvers, Finite Element Model
Begin Procedure myProcedure
  Begin Solution Control Description
    Use System Main
    Begin System Main
      Begin Sequential MySolveBlock
        Advance myRegion
      End
    End
  End
Begin Aria Region myRegion
  . ICs, BCs, equations
  . myRegion output
End Aria Region myRegion
```

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A solution control command block for steady-state analysis containing nonlinear iteration for Aria and Adagio would reflect the structure indicated below:

```
Begin Sierra myJob
.
. Materials, Solvers, Finite Element Model

Begin Procedure myProcedure

Begin Solution Control Description
  Use System Main
  Begin System Main
    Begin Sequential MySolveBlock
      Begin Nonlinear Iteration
        Advance myAriaRegion
        Advance myAdagioRegion
        transfer adagio_to_aria
      End Nonlinear Iteration
    End
  End
End

Begin transfer adagio_to_aria
.
  transfer commands
.
End transfer adagio_to_aria

Begin Aria Region myAriaRegion
.
  ICs, BCs, equations
  myAriaRegion output
.
End Aria Region myAriaRegion

Begin Adagio Region myAdagioRegion
.
  ICs, BCs, equations
  myAdagioRegion output
.
End Adagio Region myAdagioRegion

End Procedure myProcedure
.
End Sierra myJob
```

In the case of transient analysis the solution control command block will contain specification of times for which the analysis will be carried out. Additionally parameters defining the time integration must also
be supplied by the user. The details concerning time integration parameters are included in the section on Time Integration 16. A simple example the solution control command block for transient analysis would resemble the structure indicated below: Thus an example solution control for transient analysis is deferred to the section on Time Integration 16.

```
Begin Sierra myJob
  
  . Materials, Solvers, Finite Element Model
  
  Begin Procedure My_Aria_Procedure

  Begin Solution Control Description

  Use System Main

  Begin System Main
  
  Simulation Start Time = 0.0
  Simulation Termination Time = 10.0
  Simulation Max Global Iterations = 1000

  Begin Transient Time_Block_1
    Advance My_Aria_Region
  End

  Begin Transient Time_Block_2
    Advance My_Aria_Region
  End

  End

Begin Parameters For Transient Time_Block_1
  Start Time = 0.0
  Number of steps = 8
  Begin Parameters For Aria Region My_Aria_Region
    Time Step Variation = Fixed
    Initial Time Step Size = 0.001
  End

Begin Parameters For Transient Time_Block_2
  Begin Parameters For Aria Region My_Aria_Region
    Time Step Variation = Adaptive
    Initial Time Step Size = 0.001
    Predictor-Corrector Tolerance = 1e-3
    Minimum Time Step Size = 1e-6
  End

  End

End Procedure My_Aria_Procedure
  
  End Sierra myJob
```
Similarly subcycled iterations in a one-way coupling between Aria and Presto could also be carried out in a transient analysis.

Begin Sierra myJob
.
Begin Procedure My_Aria_Procedure

Begin Solution Control Description

Use System Main

Begin System Main
Simulation Start Time = 0.0
Simulation Termination Time = 10.0
Simulation Max Global Iterations = 1000

Begin Transient Time_Block_1
Transfer Presto_to_Aria
Advance My_Aria_Region
Begin Subcycle PrestoSubcycle
  Advance PrestoRegion
End
End

Begin Parameters For Transient Time_Block_1
Start Time = 0.0
Number of steps = 8

Begin Parameters For Aria Region My_Aria_Region
  Time Step Variation = Fixed
  Initial Time Step Size = 0.001
End

Begin Parameters for Presto Region PrestoRegion
  initial time step = 1.0e-6
  # time step scale factor = 1.0
  time step increase factor = 10.
  # step interval = 500
End

End
.
.
Begin Aria Region myAriaRegion
.
End Aria Region myAriaRegion

Begin Presto Region myPrestoRegion
End Presto Region myPrestoRegion

End Procedure myProcedure

End Sierra myJob

It is important to note that Solution Control can orchestrate the execution of one Region or the execution of many Regions. Within a loosely-coupled code analysis SC is also used to control the movement of data between the coupled codes using the Transfer subsystem. In what follows it is implied that optional \[\text{whenExpression}\]s represent evaluations which must be interpreted at parse time. Hence the \[\text{whenExpression}\] must be enclosed in quotes within the input file.

In the case of transient analysis it is sometimes necessary to initialize a distribution of values before the analysis actually begins. As an example, one may want to initialize a Field that will be transferred to another Region with a distribution of values with the goal of setting a reference state. For this purpose solution control provides a means of initialization, Initialize.

Begin Sierra myJob
  . Materials, Solvers, Finite Element Model
  . Begin Procedure My_Aria_Procedure
    Begin Initialize
      Transfer var1_Region_to_var2_My_Aria_Region
    End Initialize
    Begin Solution Control Description
      Use system Initialize
      Use System Main
      Begin System Main
        Simulation Start Time = 0.0
        Simulation Termination Time = 10.0
        Simulation Max Global Iterations = 1000
        Begin Transient Time_Block_1
          Advance My_Aria_Region
        End
        Begin Transient Time_Block_2
          Advance My_Aria_Region
        End
      End
      Begin Parameters For Transient Time_Block_1
        Start Time = 0.0
        Number of steps = 8
        Begin Parameters For Aria Region My_AriaRegion
          Time Step Variation = Fixed
          Initial Time Step Size = 0.001
        End
      End
    End
Begin Parameters For Transient Time_Block_2
  Begin Parameters For Aria Region My_Aria_Region
    Time Step Variation = Adaptive
    Initial Time Step Size = 0.001
    Predictor-Corrector Tolerance = 1e-3
    Minimum Time Step Size = 1e-6
  End
End

End Procedure My_Aria_Procedure

End Sierra myJob

14.2 Solution Control Description

Scope: Procedure

Begin Solution Control Description Name
  Use System Name
  Begin Initialize Name
  End

  Begin Parameters For
  End

  Begin System Name
  End

End

Summary
Contains the commands needed to execute an analysis using the arpeggio procedure that utilizes Solver Control.

14.2.1 Use System
Scope: Solution Control Description

Use System Name

<table>
<thead>
<tr>
<th>Parameter Name</th>
<th>Value string</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
This set the name of which system to use.
14.3 System

Scope: Solution Control Description

Begin System Name
  Adapt Region_name... Using Field_name... [ When When-expression ]
  Compute Indicator On Region_name... Using Indicator_name... [ When When-expression ]
  Event Name... [ When When-expression ]
  Execute Postprocessor Group Group_name... On Region_name... [ When When-expression ]
  Indicatemarkeadapt Region_name Using Indicator Marker [ When When-expression ]
  Mark Region_name... Using Marker_name... [ When When-expression ]
  Markadapt Region_name Using Marker [ When When-expression ]
  Output Name [ When When-expression ]
  Simulation Max Global Iterations {=|are|is} Number
  Simulation Start Time {=|are|is} Number
  Simulation Termination Time {=|are|is} Number
  Transfer Name [ When When-expression ]
  Use Initialize Name
  Begin Adaptivity Name
  End

  Begin Sequential Name
  End

  Begin Transient Name
  End

End

Summary
This block wraps a solver system for a given name. The NAME parameter is the name used to define the system. There can be more than one system block in the Solver Control Description block. The "use system NAME" line command controls which one is to be used.

14.3.1 Adapt

Scope: System

Begin System Name
  Adapt Region_name... Using Field_name... [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td>Field_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Used within a Solver Control block to indicate a mesh adaptation on the specific block should be performed.
14.3.2 Compute Indicator On
Scope: System

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td>Indicator_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Used within a Solver Control block to indicate a mesh adaptation on the specific block should be performed.

14.3.3 Event
Scope: System

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Used within a Solver Control block to indicate a single step that has no time associated with it. It can cause a solution transfer between regions or cause something to print.

14.3.4 Execute Postprocessor Group
Scope: System

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td>Region_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Used within a Solver Control block to cause the group named group_name to be executed on region region_name.

14.3.5 Indicatemarkadapt
Scope: System

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region_name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Indicator</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Marker</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Shortcut line command... equivalent to: Compute Indicator On ... Mark ... Adapt ...
14.3.6 Mark
Scope: System

Mark Region_name... Using Marker_name... [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td>Marker_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Used within a Solver Control block to indicate a mesh adaptation on the specific block should be performed.

14.3.7 Markadapt
Scope: System

Markadapt Region_name Using Marker [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region_name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Marker</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Shortcut line command... equivalent to: Mark ... Adapt ...

14.3.8 Output
Scope: System

Output Name [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: A Solver Control Output line command which execute a perform I/O on the region.

14.3.9 Simulation Max Global Iterations
Scope: System

Simulation Max Global Iterations {=|are|is} Number

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: The Total number of Solves.

14.3.10 Simulation Start Time
Scope: System
Simulation Start Time \{=|are|is\} \textit{Number}

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{Number}</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Simulation starting time. (by default 0.0)

14.3.11 Simulation Termination Time

Scope: System

Simulation Termination Time \{=|are|is\} \textit{Number}

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{Number}</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: The drop dead time.

14.3.12 Transfer

Scope: System

\texttt{Transfer \textit{Name} \ [ When \textit{When-expression} ]}

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{Name}</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: A Solver Control Transfer line command which executes all transfers defined from the specified region. All transfers with a send region of 'name' will be executed.

14.3.13 Use Initialize

Scope: System

\texttt{Use Initialize \textit{Name}}

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{Name}</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: This set the name of which initialization to use.

14.4 Transient

Scope: System

\texttt{Begin Transient \textit{Name}}

\begin{itemize}
\item \texttt{Adapt Region\_name... Using Field\_name... \ [ When \textit{When-expression} ]}
\item \texttt{Advance \textit{Name}... \ [ When \textit{When-expression} ]}
\end{itemize}
Computes Indicator On Region_name... Using Indicator_name... [ When When-expression ]

Event Name... [ When When-expression ]

Execute Postprocessor Group Group_name... On Region_name... [ When When-expression ]

IndicateMarkAdapt Region_name Using Indicator Marker [ When When-expression ]

Invoke Name

Mark Region_name... Using Marker_name... [ When When-expression ]

MarkAdapt Region_name Using Marker [ When When-expression ]

Output Name [ When When-expression ]

Transfer Name [ When When-expression ]

Begin Adaptivity Name
End

Begin Nonlinear Name
End

Begin Subcycle Name
End

End

Summary   This block is used to wrap a time loop.

14.4.1 Adapt
Scope: Transient

Adapt Region_name... Using Field_name... [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td>Field_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary   Used within a Solver Control block to indicate a mesh adaptation on the specific block should be performed.

14.4.2 Advance
Scope: Transient

Advance Name... [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary   Used within a Solver Control block to indicate a single step that advances the solution. The name is that matches the physics.
14.4.3 Compute Indicator On
Scope: Transient

Compute Indicator On Region_name... Using Indicator_name... [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td>Indicator_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Used within a Solver Control block to indicate a mesh adaptation on the specific block should be performed.

14.4.4 Event
Scope: Transient

Event Name... [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Used within a Solver Control block to indicate a single step that has no time associated with it. It can cause a solution transfer between regions or cause something to print.

14.4.5 Execute Postprocessor Group
Scope: Transient

Execute Postprocessor Group Group_name... On Region_name... [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td>Region_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Used within a Solver Control block to cause the group named group_name to be executed on region region_name.

14.4.6 Indicatemarkadapt
Scope: Transient

Indicatemarkadapt Region_name Using Indicator Marker [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region_name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Indicator</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Marker</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Shortcut line command... equivalent to: Compute Indicator On ... Mark ... Adapt ...
### 14.4.7 Involve

**Scope:** Transient

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Name</em></td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** Specify a physics participant to a coupled problem solved using matrix-free nonlinear.

### 14.4.8 Mark

**Scope:** Transient

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Region name</em></td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td><em>Marker name</em></td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** Used within a Solver Control block to indicate a mesh adaptation on the specific block should be performed.

### 14.4.9 Markadapt

**Scope:** Transient

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Region name</em></td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td><em>Marker</em></td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** Shortcut line command... equivalent to: Mark ... Adapt ...

### 14.4.10 Output

**Scope:** Transient

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Name</em></td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** A Solver Control Output line command which execute a perform I/O on the region.

### 14.4.11 Transfer

**Scope:** Transient
Transfer Name [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  A Solver Control Transfer line command which executes all transfers defined from the specified region. All transfers with a send region of 'name' will be executed.

14.5  Nonlinear
Scope:  Sequential

```plaintext
Begin Nonlinear Name
    Adapt Region_name... Using Field_name... [ When When-expression ]
    Advance Name... [ When When-expression ]
    Compute Indicator On Region_name... Using Indicator_name... [ When When-expression ]
    Event Name... [ When When-expression ]
    Execute Postprocessor Group Group_name... On Region_name... [ When When-expression ]
    Indicatemarkadapt Region_name Using Indicator Marker [ When When-expression ]
    Involve Name
    Mark Region_name... Using Marker_name... [ When When-expression ]
    Markadapt Region_name Using Marker [ When When-expression ]
    Output Name [ When When-expression ]
    Transfer Name [ When When-expression ]
    Begin Subcycle Name
End
```

Summary  This block is used to wrap a nonlinear solve loop.

14.5.1  Adapt
Scope:  Nonlinear

```plaintext
Adapt Region_name... Using Field_name... [ When When-expression ]
```

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td>Field_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Used within a Solver Control block to indicate a mesh adaptation on the specific block should be performed.
14.5.2 Advance
Scope: Nonlinear

**Advance Name... [ When When-expression ]**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Used within a Solver Control block to indicate a single step that advances the solution. The name is that matches the physics.

14.5.3 Compute Indicator On
Scope: Nonlinear

**Compute Indicator On Region_name... Using Indicator_name... [ When When-expression ]**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td>Indicator_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Used within a Solver Control block to indicate a mesh adaptment on the specific block should be performed.

14.5.4 Event
Scope: Nonlinear

**Event Name... [ When When-expression ]**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Used within a Solver Control block to indicate a single step that has no time associated with it. It can cause a solution transfer between regions or cause something to print.

14.5.5 Execute Postprocessor Group
Scope: Nonlinear

**Execute Postprocessor Group Group_name... On Region_name... [ When When-expression ]**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td>Region_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Used within a Solver Control block to cause the group named group_name to be executed on region region_name.
14.5.6 Indicatemarkadapt
Scope: Nonlinear

Indicatemarkadapt Region_name Using Indicator Marker [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region_name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Indicator</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Marker</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Shortcut line command... equivalent to: Compute Indicator On ... Mark ... Adapt ...

14.5.7 Involve
Scope: Nonlinear

Involve Name

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Specifiy a physics participant to a coupled problem solved using matrix-free nonlinear.

14.5.8 Mark
Scope: Nonlinear

Mark Region_name... Using Marker_name... [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td>Marker_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Used within a Solver Control block to indicate a mesh adaptment on the specific block should be performed.

14.5.9 Markadapt
Scope: Nonlinear

Markadapt Region_name Using Marker [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region_name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Marker</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Shortcut line command... equivalent to: Mark ... Adapt ...

574
14.5.10  Output  
Scope:  Nonlinear

Output Name [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  A Solver Control Output line command which execute a perform I/O on the region.

14.5.11  Transfer  
Scope:  Nonlinear

Transfer Name [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  A Solver Control Transfer line command which executes all transfers defined from the specified region. All transfers with a send region of 'name' will be executed.

14.6  Subcycle  
Scope:  Nonlinear

Begin Subcycle Name

Adapt Region_name... Using Field_name... [ When When-expression ]
Advance Name... [ When When-expression ]
Compute Indicator On Region_name... Using Indicator_name... [ When When-expression ]
Event Name... [ When When-expression ]
Execute Postprocessor Group Group_name... On Region_name... [ When When-expression ]
Indicatemarkadapt Region_name Using Indicator Marker [ When When-expression ]
Involve Name
Mark Region_name... Using Marker_name... [ When When-expression ]
Markadapt Region_name Using Marker [ When When-expression ]
Output Name [ When When-expression ]
Transfer Name [ When When-expression ]
End

Summary  This block is used to wrap a subcycle time loop.
14.6.1 Adapt
Scope: Subcycle

Adapt Region_name... Using Field_name... [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td>Field_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Used within a Solver Control block to indicate a mesh adaptment on the specific block should be performed.

14.6.2 Advance
Scope: Subcycle

Advance Name... [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Used within a Solver Control block to indicate a single step that advances the solution. The name is that matches the physics.

14.6.3 Compute Indicator On
Scope: Subcycle

Compute Indicator On Region_name... Using Indicator_name... [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td>Indicator_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Used within a Solver Control block to indicate a mesh adaptment on the specific block should be performed.

14.6.4 Event
Scope: Subcycle

Event Name... [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Used within a Solver Control block to indicate a single step that has no time associated with it. It can cause a solution transfer between regions or cause something to print.
14.6.5 Execute Postprocessor Group

Scope: Subcycle

Execute Postprocessor Group **Group_name...** On **Region_name...** [ When **When-expression** ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Group_name</strong></td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td><strong>Region_name</strong></td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Used within a Solver Control block to cause the group named **group_name** to be executed on region **region_name**.

14.6.6 Indicatemarkadapt

Scope: Subcycle

Indicatemarkadapt **Region_name** Using **Indicator Marker** [ When **When-expression** ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Region_name</strong></td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td><strong>Indicator</strong></td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td><strong>Marker</strong></td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Shortcut line command... equivalent to: Compute Indicator On ... Mark ... Adapt ...

14.6.7 Involve

Scope: Subcycle

Involve **Name**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Name</strong></td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify a physics participant to a coupled problem solved using matrix-free nonlinear.

14.6.8 Mark

Scope: Subcycle

Mark **Region_name...** Using **Marker_name...** [ When **When-expression** ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Region_name</strong></td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td><strong>Marker_name</strong></td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Used within a Solver Control block to indicate a mesh adaptation on the specific block should be performed.
14.6.9  Markadapt
Scope:  Subcycle

Markadapt Region_name Using Marker [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region_name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Marker</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Shortcut line command... equivalent to: Mark ... Adapt ...

14.6.10  Output
Scope:  Subcycle

Output Name [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  A Solver Control Output line command which execute a perform I/O on the region.

14.6.11  Transfer
Scope:  Subcycle

Transfer Name [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  A Solver Control Transfer line command which executes all transfers defined from the specified region. All transfers with a send region of ‘name’ will be executed.

14.7  Sequential
Scope:  System

Begin Sequential Name
  Adapt Region_name... Using Field_name... [ When When-expression ]
  Advance Name... [ When When-expression ]
  Compute Indicator On Region_name... Using Indicator_name... [ When When-expression ]
  Event Name... [ When When-expression ]
  Execute Postprocessor Group Group_name... On Region_name... [ When When-expression ]
  Indicatemarkadapt Region_name Using Indicator Marker [ When When-expression ]
Involve Name
Mark Region_name... Using Marker_name... [ When When-expression ]
Markadapt Region_name Using Marker [ When When-expression ]
Output Name [ When When-expression ]
Transfer Name [ When When-expression ]
Begin Adaptivity Name
End

Begin Nonlinear Name
End

Summary This block is used to wrap a sequential solution. It is used to wrap a sequence of Non-Linear or pseudo time solve step solves.

### 14.7.1 Adapt
**Scope:** Sequential

Adapt Region_name... Using Field_name... [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td>Field_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Used within a Solver Control block to indicate a mesh adaptment on the specific block should be performed.

### 14.7.2 Advance
**Scope:** Sequential

Advance Name... [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Used within a Solver Control block to indicate a single step that advances the solution. The name is that matches the physics.

### 14.7.3 Compute Indicator On
**Scope:** Sequential

Compute Indicator On Region_name... Using Indicator_name... [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td>Indicator_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Summary: Used within a Solver Control block to indicate a mesh adaptation on the specific block should be performed.

14.7.4 Event
Scope: Sequential

Event Name... [When When-expression]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Used within a Solver Control block to indicate a single step that has no time associated with it. It can cause a solution transfer between regions or cause something to print.

14.7.5 Execute Postprocessor Group
Scope: Sequential

Execute Postprocessor Group Group_name... On Region_name... [When When-expression]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td>Region_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Used within a Solver Control block to cause the group named group_name to be executed on region region_name.

14.7.6 Indicatemarkadapt
Scope: Sequential

Indicatemarkadapt Region_name Using Indicator Marker [When When-expression]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region_name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Indicator</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Marker</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Shortcut line command... equivalent to: Compute Indicator On ... Mark ... Adapt ...

14.7.7 Involve
Scope: Sequential

Involve Name

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify a physics participant to a coupled problem solved using matrix-free nonlinear.
14.7.8 Mark
Scope: Sequential

Mark Region_name... Using Marker_name... [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td>Marker_name</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Used within a Solver Control block to indicate a mesh adaptment on the specific block should be performed.

14.7.9 Markadapt
Scope: Sequential

Markadapt Region_name Using Marker [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region_name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Marker</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Shortcut line command... equivalent to: Mark ... Adapt ...

14.7.10 Output
Scope: Sequential

Output Name [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: A Solver Control Output line command which execute a perform I/O on the region.

14.7.11 Transfer
Scope: Sequential

Transfer Name [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: A Solver Control Transfer line command which executes all transfers defined from the specified region. All transfers with a send region of 'name' will be executed.
14.8 Initialize

Scope: Solution Control Description

Begin Initialize Name
  Advance Name... [ When When-expression ]
  Event Name... [ When When-expression ]
  Involve Name
    Transfer Name [ When When-expression ]
End

Summary This block wraps a initializer for a given name. The NAME parameter is the name used to define the initialization block. There can be more than one initialize block in the Solver Control Description block. The "use initialize NAME" line command controls which one is to be used.

14.8.1 Advance

Scope: Initialize

Advance Name... [ When When-expression ]

Parameter | Value | Default  
---|---|---
Name | string... | undefined |

Summary Used within a Solver Control block to indicate a single step that advances the solution. The name is that matches the physics.

14.8.2 Event

Scope: Initialize

Event Name... [ When When-expression ]

Parameter | Value | Default  
---|---|---
Name | string... | undefined |

Summary Used within a Solver Control block to indicate a single step that has no time associated with it. It can cause a solution transfer between regions or cause something to print.

14.8.3 Involve

Scope: Initialize

Involve Name

Parameter | Value | Default  
---|---|---
Name | string | undefined |
14.8.4 Transfer
Scope: Initialize

Transfer Name [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
A Solver Control Transfer line command which executes all transfers defined from the specified region. All transfers with a send region of 'name' will be executed.

14.9 Parameters For
Scope: Solution Control Description

Begin Parameters For

Converged When Convergence-expression
Incremental Number Of Steps {=|are|is} Number
Initial Deltat {=|are|is} Number
Number Of Adaptivity Steps {=|are|is} Number
Number Of Steps {=|are|is} Number
Reinitialize Transient
Start Time {=|are|is} Number
Termination Time {=|are|is} Number
Time Step Quantum {=|are|is} TimeStepQuantum
Time Step Style TimeStepStyle...
Total Change In Time {=|are|is} Number
Begin Parameters For Aria Region RegionName
End

Summary
A Solver Control PARAMETERS block to set up control data for the SC_type parameter. Inside this block one sets the time step parameters or nonlinear parameters.

14.9.1 Converged When
Scope: Parameters For

Converged When Convergence-expression

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Convergence-expression</td>
<td>(expression)</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Set the convergence expression.
14.9.2  Incremental Number Of Steps
Scope: Parameters For

Incremental Number Of Steps {=|are|is} Number

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  The incremental number steps to run the time for nonlinear loop. Number of time steps to run after restarting. NUMBER OF STEPS is total number of steps to run

14.9.3  Initial Deltat
Scope: Parameters For

Initial Deltat {=|are|is} Number

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Assign an initial delta T

14.9.4  Number Of Adaptivity Steps
Scope: Parameters For

Number Of Adaptivity Steps {=|are|is} Number

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  The number steps to run the time or nonlinear loop

14.9.5  Number Of Steps
Scope: Parameters For

Number Of Steps {=|are|is} Number

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  The number steps to run the time for nonlinear loop

14.9.6  Reinitialize Transient
Scope: Parameters For

Summary  Reset time and re-initialize regions each step of the adaptivity loop.
### 14.9.7 Start Time

**Scope:** Parameters For

| Start Time \(\text{=} | \text{are} | \text{is} \) \(\text{Number}\) |
|----------------|
| **Parameter** | **Value** |
| \(\text{Number}\) | \(\text{real}\) |
| **Default** | **undefined** |

**Summary**
Assign a start time.

### 14.9.8 Termination Time

**Scope:** Parameters For

| Termination Time \(\text{=} | \text{are} | \text{is} \) \(\text{Number}\) |
|----------------|
| **Parameter** | **Value** |
| \(\text{Number}\) | \(\text{real}\) |
| **Default** | **undefined** |

**Summary**
Assign a final time to stop.

### 14.9.9 Time Step Quantum

**Scope:** Parameters For

| Time Step Quantum \(\text{=} | \text{are} | \text{is} \) \(\text{TimeStepQuantum}\) |
|----------------|
| **Parameter** | **Value** |
| \(\text{TimeStepQuantum}\) | \(\text{real}\) |
| **Default** | **undefined** |

**Summary**
Set the time stepping quantum time for SNAP style stepping.

### 14.9.10 Time Step Style

**Scope:** Parameters For

<table>
<thead>
<tr>
<th>Time Step Style (\text{TimeStepStyle}\ldots)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Parameter</strong></td>
</tr>
<tr>
<td>(\text{TimeStepStyle})</td>
</tr>
<tr>
<td><strong>Default</strong></td>
</tr>
</tbody>
</table>

**Summary**
Set the time stepping style.

When CLIP is specified, the time step size will be clipped at the last step of the transient loop so that it ends at the transient loop’s end time. If clip is not specified, the last time is allowed to exceed to the transient loop’s end time and the following transient loop will start at the exceeded end time.

When SNAP is specified, the time step is broken down into "quantum" time units. By default this quantum time is 12 orders of magnitude down from the difference between the start and end time for the transient loop. This value can be overridden using the TIME STEP QUANTUM line command. All time values are "snapped" to multiples of the quantum time by rounding to the nearest quantum multiple.
### 14.9.11 Total Change In Time

**Scope:** Parameters For

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>Number</code></td>
<td><code>real</code></td>
<td><code>undefined</code></td>
</tr>
</tbody>
</table>

**Summary**
Use this number and the initial time to compute termination time.

### 14.9.12 Advance

**Scope:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>Name</code></td>
<td><code>string...</code></td>
<td><code>undefined</code></td>
</tr>
</tbody>
</table>

**Summary**
Used within a Solver Control block to indicate a single step that advances the solution. The name is that matches the physics.

### 14.9.13 Converged When

**Scope:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>Convergence-expression</code></td>
<td><code>(expression)</code></td>
<td><code>undefined</code></td>
</tr>
</tbody>
</table>

**Summary**
Set the convergence expression.

### 14.9.14 Event

**Scope:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>Name</code></td>
<td><code>string...</code></td>
<td><code>undefined</code></td>
</tr>
</tbody>
</table>

**Summary**
Used within a Solver Control block to indicate a single step that has no time associated with it. It can cause a solution transfer between regions or cause something to print.

### 14.9.15 Initial Deltat

**Scope:**

---

586
Initial Delta {=|are|is} \textit{Number}

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{Number}</td>
<td>\text{real}</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Assign an initial delta T

14.9.16 Involve

Scope:

\text{Involve} \textit{Name}

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{Name}</td>
<td>\text{string}</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify a physics participant to a coupled problem solved using matrix-free nonlinear.

14.9.17 Number Of Adaptivity Steps

Scope:

Number Of Adaptivity Steps {=|are|is} \textit{Number}

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{Number}</td>
<td>\text{integer}</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: The number steps to run the time or nonlinear loop

14.9.18 Number Of Steps

Scope:

Number Of Steps {=|are|is} \textit{Number}

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{Number}</td>
<td>\text{integer}</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: The number steps to run the time for nonlinear loop

14.9.19 Output

Scope:

Output \textit{Name} [ When \textit{When-expression} ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{Name}</td>
<td>\text{string}</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: A Solver Control Output line command which execute a perform I/O on the region.
14.9.20 Reinitialize Transient
Scope:

Summary Reset time and re-initialize regions each step of the adaptivity loop.

14.9.21 Simulation Max Global Iterations
Scope:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Simulation Max Global Iterations</td>
<td>= Number</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary The Total number of Solves.

14.9.22 Simulation Start Time
Scope:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Simulation Start Time</td>
<td>= Number</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Simulation starting time. (by default 0.0)

14.9.23 Simulation Termination Time
Scope:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Simulation Termination Time</td>
<td>= Number</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary The drop dead time.

14.9.24 Start Time
Scope:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Start Time</td>
<td>= Number</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Assign a start time.
14.9.25 Termination Time
Scope:

| Termination Time {=|are|is} Number |
|-----------------------------------|
| Parameter | Value | Default |
| Number     | real  | undefined |

Summary Assign a final time to stop

14.9.26 Time Step Quantum
Scope:

| Time Step Quantum {=|are|is} TimeStepQuantum |
|---------------------------------------------|
| Parameter | Value | Default |
| TimeStepQuantum | real   | undefined |

Summary Set the time stepping quantum time for SNAP style stepping.

14.9.27 Time Step Style
Scope:

<table>
<thead>
<tr>
<th>Time Step Style TimeStepStyle...</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter</td>
</tr>
<tr>
<td>TimeStepStyle</td>
</tr>
</tbody>
</table>

Summary Set the time stepping style.

When CLIP is specified, the time step size will be clipped at the last step of the transient loop so that it ends at the transient loop’s end time. If clip is not specified, the last time is allowed to exceed to the transient loop’s end time and the following transient loop will start at the exceeded end time.

When SNAP is specified, the time step is broken down into "quantum" time units. By default this quantum time is 12 orders of magnitude down from the difference between the start and end time for the transient loop. This value can be overridden using the TIME STEP QUANTUM line command. All time values are "snapped" to multiples of the quantum time by rounding to the nearest quantum multiple.

14.9.28 Total Change In Time
Scope:

| Total Change In Time {=|are|is} Number |
|---------------------------------------|
| Parameter | Value | Default |
| Number     | real  | undefined |

Summary Use this number and the initial time to compute termination time.
14.9.29 Transfer

Scope:

Transfer Name [ When When-expression ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: A Solver Control Transfer line command which executes all transfers defined from the specified region. All transfers with a send region of 'name' will be executed.

14.9.30 Use Initialize

Scope:

Use Initialize Name

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: This set the name of which initialization to use.

14.9.31 Use System

Scope:

Use System Name

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: This set the name of which system to use.
Chapter 15

Transfer Reference

15.1 Overview

The Sierra Transfer utility provides the means by which to communicate data between two Sierra application Regions. The utility is fairly flexible as it provides the ability to move data directly onto another problem domain either by direct copy or by interpolation. Analysts without prior experience with transfer are often uncertain as to which type of transfer to use. The two capabilities function exactly as their names imply but understanding which method to use requires a basic understanding of how each method works.

Copy transfer assumes that the discretization for applications involved in the transfer are identical. Moreover, copy transfer also assumes that the mesh is identical so that global IDs of nodes and elements within each mesh are the same. Under these assumptions a geometric search of source to destination locations is not necessary and a simple algorithm is able to perform the data transfer in a straightforward manner.

Interpolation transfer is much more general than copy transfer since it assumes only that data from one application must be geometrically mapped for use in another application. A mathematical definition of this mapping is made possible using the results from a geometric search of points on the destination mesh and their image on the sending mesh. With regard to code performance copy transfer will always more efficient than interpolation transfer but is rarely applicable in mainstream simulations. Interpolate transfer is designed to deal with complications that arise in mapping data from one application to the other and is more reliable. As a rule, one should always use interpolation transfer and not copy transfer. At the same time an analyst should strategize model construction so as to offset some of the performance costs of interpolation transfer.

Even with a basic understanding of transfer users of what transfer operations should be defined. Several proper transfer source and destinations are illustrated in Figure 15.1, here the numbers on the figures correspond to the ExodusII global IDs of nodes or elements.

Problematic transfer source and destination configurations are illustrated in Figure 15.2. Once again the numbers on the figures correspond to the ExodusII global IDs of nodes or elements.

In using the transfer utility one must clearly define the sending region (where the data resides) and the the receiving region (the data destination). Additionally one must also specify the general geometric location of data sender and receiver based upon existing mesh entities (blocks or surfaces). Sender and receiver need not be of same topology but the source and target destinations should overlap geometrically. Clearly the definition mesh entities influences time spent in the geometric search process and should be a key consideration in model construction.

Both Aria solution Fields and User Fields 2.9 can be transferred to/from other Sierra application Regions. Transfer of data into Aria requires that provision be made for a Field storage on the Aria Region. Storage for solution Field (DOF) transfer into Aria is implicitly provided by the presence of an Aria equation command line with term type of XFER (Chapter 6). Storage for non-DOF Fields is implicitly provided through the presence of a User Field command line 2.10. These non-DOF Fields are internally used or modified by virtue
of being referenced in other command lines such as material models, boundary conditions, initial conditions or source terms.

The following section outlines the commands to be used in setting up transfer operations. Special attention should be paid to the syntax of the SEND command line since it differs between COPY and INTERPOLATION transfer.

Since several different uses of transfer can arise and several of those examples for steady problems are included below. The same basic setup of transfer would apply to transient problems as well.

A skeleton outline of one-way transfer from Region_1 to Region_2 in a steady-state problem would be:

```
Begin Sierra
  Begin Transfer my_transfer
    transfer commands for first_region to second_region
  End
  Begin Procedure My_Aria_Procedure
    Begin Solution Control Description
      Use System Main
      Begin System Main
        Begin Sequential MySolveBlock
          ...
        End
      End
    End
  End
End
```
Figure 15.2. Invalid Transfer Operation

Advance first_region
transfer my_transfer
Advance second_region
End
End
End

Begin Aria Region first_region
   .
   eq energy for temperature On block_1 using q1 with lumped_mass diff
   .
End

Begin Aria Region second_region
   .
   eq energy for temperature On block_1 using q1 with xfer
   .
End Sierra
A skeleton outline of two-way transfer between Region_1 to Region_2 in a steady-state problem would be:

Begin Sierra
  .
  Begin Transfer my_first_transfer
    .
    transfer commands for first_region to second_region
    .
    End
  .
  Begin Transfer my_second_transfer
    .
    transfer commands for second_region to first_region
    .
    End
  .
  Begin Procedure My_Aria_Procedure
    .
    Begin Solution Control Description
      .
      Use System Main
      .
      Begin System Main
        .
        Begin Sequential MySolveBlock
          .
          Advance first_Region
          .
          transfer my_first_transfer
          .
          Advance second_Region
          .
          transfer my_second_transfer
          End
        End
      End
    End
  Begin Aria Region first_region
    .
    eq energy for temperature On block_1 using q1 with diff
    eq species_3 for temperature On block_1 using q1 with xfer
    .
    End
  Begin Aria Region second_region
    .
    eq energy for temperature On block_1 using q1 with xfer
    eq species_3 for species_3 On block_1 using q1 with diff
    .
    End
  End
End Sierra

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Assume an input mesh for an Input_Output Region 26.1 contains a nodal variable ConvCoeff. In this case a skeleton outline for one-way transfer of ConvCoeff to to Region_2 in a steady-state problem would be:

```
Begin Sierra
  .
  Begin Transfer my_first_transfer
    .
    transfer commands for input_output_region to second_region
    .
    SEND field hNd state none TO ConvCoeff state none
    .
  End
  .
  Begin Procedure My_Aria_Procedure
    .
    Begin Solution Control Description
      Use System Main
      Begin System Main
        Begin Sequential MySolveBlock
          Advance first_Region
          transfer my_first_transfer
          Advance second_Region
        End
      End
    End
  .
  Begin Input_Output io_region
    USE FINITE ELEMENT MODEL my_input_transfer
  End
  .
  Begin Aria Region second_region
    .
    USER FIELD REAL NODE SCALAR ConvCoeff on surface_1
    .
  End
  .
End Sierra
```

### 15.2 Transfer

**Scope:** Procedure

```
Begin Transfer Transfer_name
  Abort If Field Not Defined On Copy Transfer Send Or Receive Object All Fields
  Copy Option1 Option2 From From_region_name To To_region_name
  Distance Function Is Closest Receive Node To Send Centroid
```

596
Exclude Ghosted
Experimental Option To Use Sending Mesh In Place
From Option1 To Option2
Gauss Point Integration Order {=|are|is} Order
Interpolate Option1 Option2 From From_region_name To To_region_name
Interpolation Function User_Subroutine
Nodes Outside Region {=|are|is} Option
Search Coordinate Field Source_field_name State Option1 To Destination_field_name State Option2
Search Geometric Tolerance {=|are|is} Geometric_tolerance
Search Surface Gap Tolerance {=|are|is} Surface_gap_tolerance [ Or Less ]
Search Type {=|are|is} [ Option1 Option2 Option3 ]
Select One Receiver For Each Send Object
Select One Unique Receiver For Each Send Object
Send Predefined-transfer Fields
Send Block From_blocks... To To_blocks...
Send Field Source_field_name State Option1 To Destination_field_name State Option2
[ Lower Bound Lower_bound Upper Bound Upper_bound ]
Begin Receive Blocks
End
Begin Send Blocks
End

Summary transfer region/mesh information. the mechanics/variables information will get sorted out by the calling procedure.

15.2.1 Abort If Field Not Defined On Copy Transfer Send Or Receive Object
Scope: Transfer

Summary For testing purposes only. Normally mesh objects in the send or receive mesh which do not have the specified field defined on them are just ignored. This line command allows the construction of tests in which it is known that every mesh object should have the specified field defined on it and to abort if that field is not found.

15.2.2 All Fields
Scope: Transfer

Summary Select all fields for transfer that have same name and state for source and destination regions.
15.2.3 Copy
Scope: Transfer

Copy Option1 Option2 From From_region_name To To_region_name

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>From_region_name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>To_region_name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Copy transfer elements, nodes or constraints from one region to another. The copy transfer is very specific in that the sending and receiving mesh parts must have identical global ids for every element to be copied. The copy transfer works by iterating over all the mesh objects in the receiving mesh and using the global id of the receiving mesh object to find a mesh object in the sending mesh with the same global id. The field to transfer is then copied from the sending to receiving objects. There is no interpolation and the actual coordinates of the sending and receiving objects are not used and could be very different. The copy transfer is used in very special cases where the same mesh was read into both the sending and receiving meshes, there was no element death and there was no adaptivity. In this special case, a copy transfer can be much faster than an interpolation transfer.

15.2.4 Distance Function Is Closest Receive Node To Send Centroid
Scope: Transfer

Summary
To be used in conjunction with "SELECT ONE UNIQUE RECEIVER FOR EACH SEND OBJECT". This helped in the case where the sending and receiving element blocks did not overlap and an element transfer was using element centroids for the distance computation. The elements were very distorted so that a centroid of a surface element could be far from the surface. It was wanted that the receiving element be the one close to the surface of the block and close to the sending element in the adjacent block. Using the corner nodes was enough since it was a tet mesh with plane faces. In this particular and unusual case this alternative method of matching sending and receiving elements was useful, but it is not expected to be used often or maybe never again.

15.2.5 Exclude Ghosted
Scope: Transfer

Summary
exclude ghosted nodes from a copy transfer

15.2.6 Experimental Option To Use Sending Mesh In Place
Scope: Transfer

Summary
For testing purposes only. Normally the sending primary region’s objects are copied to a secondary region. This can require a large amount of memory. If specified this line command will attempt to use the sending region objects without copying them. This does not always
get the exact same answer as the standard method and I’m not sure why. It could be that just processing the mesh objects in a slightly different order causes slight differences or it could be that there is still a bug with this option.

15.2.7 From
Scope: Transfer

Summary Allows the send/receive mesh objects to be different.

15.2.8 Gauss Point Integration Order
Scope: Transfer

Gauss Point Integration Order \{=|are|is\} Order

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Order</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Integration order to use when transferring to Gauss points.

15.2.9 Interpolate
Scope: Transfer

Interpolate Option1 Option2 From From_region_name To To_region_name

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>From_region_name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>To_region_name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Interpolate will transfer elements, nodes or constraints from one mesh to another. The interpolation transfer is very general in that the field values to transfer will be interpolated from the sending to receiving mesh based on the coordinates of the sending and receiving mesh objects.

Many line commands can be used to modify the behavior of the interpolation transfer but the basic algorithm is straightforward. Every mesh object in the receiving mesh is converted into a point. For elements this is the average of the nodal coordinates. An element in the sending mesh containing this point is found. If the field to transfer is nodal, the element shape functions are used to interpolate the nodal field to the receiving point. If the field to transfer is elemental, a bi-linear least squares fit based upon neighboring elements is first performed and then used to define the interpolation of the element field at the receiving point.

15.2.10 Interpolation Function
Scope: Transfer

Interpolation Function User_Subroutine
### 15.2.11 Nodes Outside Region

**Scope:** Transfer

**Summary**
This line command defines what to do when a receiving point is outside the scope of the sending mesh.

- **IGNORE** - The receiving mesh object can be ignored and will receive no value. This is almost never a good idea as it can cause mesh objects just outside to have a zero value when the nodes just inside the mesh might have very large values. This can result in a discontinuous receiving field.

- **EXTRAPOLATE** - This is the default behavior. The sending field is extrapolated beyond the bounds of the sending mesh. This can lead to extrapolation error, such as when a large gradient at the surface causes a negative values when only positive values are acceptable. If this happens to the upper and lower bounds that can be placed on the fields to be transferred with the SEND FIELD command.

- **TRUNCATE** - The receiving coordinate is projected back to the surface of the sending mesh to determine a value. This ensures that the receiving value is outside of the field values in the sending mesh.

- **PROJECT** - This option is similar to TRUNCATE in which the receiving coordinate is projected back to the surface of the sending mesh to determine a value. In this case more effort is made to make sure that the projection is normal to the surface in the sending mesh. Sometimes gives a better result than Truncate but is a little more expensive to compute.

If the PROJECT option is used in transferring of surface values, the sending mesh should envelop the receiving mesh. Failure to satisfy this condition will generally result in failure of the transfer.

### 15.2.12 Search Coordinate Field

**Scope:** Transfer

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Source_field_name</strong></td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td><strong>Destination_field_name</strong></td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Normally the interpolation transfers use the default coordinate field to determine geometry information. This line command can be used to specify an alternate field.
15.2.13 Search Geometric Tolerance
Scope: Transfer

Search Geometric Tolerance \texttt{=} \texttt{are} \texttt{is} \texttt{Geometric
tolerance}

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
</table>
| Geometric
tolerance   | real  | undefined |

15.2.14 Search Surface Gap Tolerance
Scope: Transfer

Search Surface Gap Tolerance \texttt{=} \texttt{are} \texttt{is} \texttt{Surface
gap
tolerance} \texttt{Or Less}

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
</table>
| Surface_gap
tolerance | real  | undefined |

Summary
This is a tricky parameter best ignored, let it default to some small number. During the interpolation transfer there is a geometric search based on the coordinates of the send and receive objects. As part of this search, an axis aligned bounding box is contracted for each sending object and SEARCH GAP TOLERANCE is used to make this box bigger than just a tight bounding box. Lists of receiving points are then quickly found within these axis aligned boxes.

If all points in the receiving mesh are within at least one box, no additional searching needs to be done and the search algorithm is fast. If there are still points in the receiving mesh that were outside of EVERY box, then a warning message will be issued about an "expensive search for extrapolation" for these points. This "expensive search" can be very costly if a large number of receiving objects fall into this category and this line command is provided for those special cases.

The OR LESS optional parameter is used when the tolerance must be set to large value for one part of the mesh but much of the mesh needs a much smaller value. In some cases it is necessary for the tolerance to be set to the actual largest surface gap tolerance which may be far too large a gap for the rest of the mesh. Setting OR LESS allows the search tolerance to be reduced in areas of the mesh thus resulting in a faster search.

15.2.15 Search Type
Scope: Transfer

15.2.16 Select One Receiver For Each Send Object
Scope: Transfer

Summary
This option will cause each sending object to be used once and only once. This will have the side effect of some receiving objects not getting any value at all. If you use this option, you will also want to set NODES OUTSIDE REGION IGNORE The example which necessitated this option was a case in which there was a delta function defined on an element in the sending mesh. It was desirable that the delta functions be summed into the receiving mesh such that the total value of the sending was conserved. It was better to have only a single element on
the receiving side have a non-zero value that was the sum of sending values and not worry about how close the receiving element was to the sending element. A check that this option is working is to use Encore to computer the sum of the values of the sending and receiving fields to make sure the total sum is the same.

15.2.17 Select One Unique Receiver For Each Send Object

**Scope:** Transfer

**Summary**
An unusual flag to get around an odd problem. Normally each receive object transfers from the nearest sending object so it is almost always the case that a send object will be used multiple times to define a receiving value. This option will cause each sending object to be used only once. This will have the side effect of some receiving objects not getting any value at all. If you use this option, you will also want to set NODES OUTSIDE REGION IGNORE or else the uniqueness will be lost for nodes outside the sending region. The example which necessitated this option was a case in which there was a delta function defined on an element in the sending mesh. It was desirable that the delta function be defined on the receiving mesh for only a single element in the neighborhood of the sending element. The analysis was more sensitive to the number of delta functions on the receiving side than the location. So it was better to have only a single element on the receiving side have a non-zero value and not worry about how close the receiving element was to the sending element.

15.2.18 Send

**Scope:** Transfer

**Send Predefined-transfer Fields**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Predefined-transfer</td>
<td>{}</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Use predefined transfer semantics provided by the specified name.

15.2.19 Send Block

**Scope:** Transfer

**Send Block From_blocks... To To_blocks...**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>From_blocks</td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td>To_blocks</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Add element blocks to a particular same mesh element copy transfer operator.

The copy transfer can have multiple of these lines to define many blocks, but each line sends a single block to a single block: SEND BLOCK block_1 TO block_1 SEND BLOCK block_101 TO block_101

The interpolation transfer can have only a single SEND BLOCK line, but can define many from/to blocks: SEND BLOCK block_3 block_5 block_6 TO block_3 block_5
15.2.20 Send Field

Scope: Transfer

Send Field Source_field_name State Option1 To Destination_field_name State Option2 [ Lower Bound Lower_bound Upper Bound Upper_bound ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Source_field_name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Destination_field_name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary

Specifies the mapping between source and destination field names. Vector and tensor fields can be subscripted using parenthesis and 1’s based or brackets and 0 based. Notes on subscripting:

(0) Does not work for COPY transfers, only INTERPOLATION type transfers. (1) If the field name itself actually contains either parenthesis or brackets then we are in trouble and an error is going to be thrown due to a syntax error in index specification. (2) Only a single subscript is allowed so vectors of vectors or higher order tensors can not use double subscripts. But it should be possible to determine the correct offset within the field and pick out the correct value with a little effort. (3) Once subscripted, only a single value will be transferred. It is not possible to transfer multiple values starting at a certain index, instead multiple line commands must be used, as shown above. (4) The indexes can be 0 based with brackets or 1 based when using parenthesis. Although this could be very confusing if mixed within a single line command. (5) Both the from and to fields can be subscripted independently on the same line.

example SEND FIELD velocity TO velocity SEND FIELD temp TO temperature lower bound 0 SEND FIELD x TO y lower bound 10 upper bound 100 SEND FIELD A(2) TO B(3) lower bound 10 upper bound 100 SEND FIELD A[1] TO B[2] lower bound 10 upper bound 100
Chapter 16

Time Integration Commands

16.1 Setting Up a Transient Problem

Aria uses the Solution Control library from the SIERRA Mechanics Framework to orchestrate the solution sequence for a simulation. Thus all Aria input files must include a Solution Control Description block in the Procedure section of the input file. The following example illustrates use of the solution control command block. This example demonstrates that time stepping for transient analysis can be carried out with either fixed 16.3 or adaptive time stepping 16.4.

```
Begin Procedure My_Aria_Procedure

Begin Solution Control Description

Use System Main

Begin System Main

Simulation Start Time = 0.0
Simulation Termination Time = 10.0
Simulation Max Global Iterations = 1000

Begin Transient Time_Block_1
  Advance My_Aria_Region
End

Begin Transient Time_Block_2
  Advance My_Aria_Region
End

End

Begin Parameters For Transient Time_Block_1

Start Time = 0.0
Number of steps = 8

Begin Parameters For Aria Region My_Aria_Region
  Time Step Variation = Fixed
  Initial Time Step Size = 0.001
End

End
```
It is worth pointing out that for each named "Begin Transient" command block there should be a corresponding named "Begin Parameters for Transient" command block. This correlation establishes the mapping between a time block and its associated parameters. The intent of time block parameters is often misunderstood thus it is worth describing some of the underlying detail how the parameters are used.

16.2 Initial Time Step Estimation for Thermal Problems

The analysis of a transient diffusion problem requires the selection of a suitable time step for the integration procedure. This process can be automated with the time step being adaptively selected to preserve a specified time truncation error. Aria provides an adaptive time-stepping functionality, as described in Section 16.4. However, the time step selection process is not self-starting and some initial estimate of an appropriate time step is still required. In the following the discussion is primarily limited to implicit integration methods, since these methods are usually the methods of choice for most conduction models.

The selection of too large an initial time step can result in the loss of temporal accuracy in the solution and produce a nonphysical oscillatory response. Likewise, an inappropriately small initial time step may produce nonphysical, spatial oscillations in the early time temperature field due to the limited resolution ability (of temperature gradients) of the finite element mesh. Either of these difficulties may lead to stability problems if the boundary value problem is nonlinear. In any event, the solution during the oscillatory period is not accurate and these occurrences are to be avoided. A method for estimating an appropriate initial time step is outlined below. This procedure is originally due to Levi ([25]) though it has been discussed by several other authors ([26, 27, 28]).

In the following development, it is assumed that a finite element mesh has been constructed that will adequately model the thermal phenomena of interest (e.g., thermal shock problems will require a fine mesh near a boundary while slower thermal transient will be less demanding on mesh refinement). For a given spatial discretization, a local characteristic length, $\Delta x$, is chosen based on element size. Typically, this characteristic length is measured normal to a boundary on which a temperature or heat flux (source) disturbance occurs. Based on the characteristic length, the local heat transfer coefficient, $h$, and the local thermal conductivity, $k$, a local element Biot number ($Bi = h\Delta x/k$) can be computed. Note that for a prescribed temperature, heat flux or heat source boundary condition, the heat transfer coefficient, $h$, is assumed to be large leading to a large Biot number.

In order to bound the thermal gradient that will occur at the boundary in the first time step, the ratio of the temperature at a distance $\Delta x$ (characteristic length) from the boundary to the temperature on the boundary is selected. Let this ratio be defined by $\Theta = T(\Delta x)/T_{surface}$. Typical values of $\Theta$ will range from 0.10 to 0.25. With the estimated values for local Biot number and temperature ratio $\Theta$, a local Fourier number ($Fo = \alpha \Delta t/\Delta x^2$) may be found from the charts in Figures 16.2 and 16.3. These graphs are based
on an analytic solution for one-dimensional conduction with a convective boundary condition ([29]). A value of the local Fourier number and values for the characteristic length and thermal diffusivity, $\alpha$, then allow a time step to be computed.

As an example of the procedure outlined above consider the transient problem described in the following example:

Consider the transient thermal response of a finned tube radiator, as shown in Figure 16.1(a). Thermal properties for the radiator are as follows: $\rho = 2.7071 \times 10^3$ kg/m$^3$, $C = 870.854$ J/kg-K, $k = 207.566$ W/m-K, and $\alpha = 8.806 \times 10^{-5}$ m$^2$/s. The characteristic size is based on the average size of the elements along the radiator fin, as shown in Figure 16.1(b), and is equal to $\Delta x = 3.175 \times 10^{-3}$ m. The boundary condition is an applied heat flux, so the Biot number is assumed large (infinite). Using a temperature ratio of $\Theta = 0.10$, then Figure 16.2 yields a Fourier number of $Fo \approx 0.20$. Thus,

$$\Delta t = \frac{(Fo)(\Delta x^2)}{\alpha} = \frac{(0.20)(3.175 \times 10^{-3})^2}{(8.806 \times 10^{-5})} = 0.023 \text{ seconds.}$$

(a) Schematic

(b) Computational mesh

Figure 16.1. Geometry and mesh of finned radiator problem. Dimensions are in inches.

The above procedure can also be used to estimate the overall response time for a region. In this case the length scale is a characteristic length for the entire region and the temperature ratio should be order unity. The Fourier number will then produce the approximate time interval required to reach equilibrium.

16.3 Fixed Time Step Selection

Since Aria supports both adaptive and fixed time stepping it follows that this behavior be explicitly called out in the input file. For any given time block fixed time stepping is invoked via the combination of:

**TIME STEP VARIATION = FIXED**
**INITIAL TIME STEP SIZE = value**
**PREDICTOR ORDER = 0**

command lines within the Parameters for Transient command block. These command lines are further described in the adaptive time step selection section 16.4. Note that omission of the PREDICTOR ORDER = 0 command line for fixed time stepping will result in a default predictor error criteria to be considered in time step selection and may lead to unexpected results for presumed fixed time stepping scheme.
16.4 Adaptive Time Step Selection

Time step failure in Aria is defined as a Newton solution step that has not converged when the solution time was advanced. Aria time step selection is carried out by first addressing possible time step failures and then applying the type of time stepping requested, fixed or adaptive. If adaptive time stepping is requested then a number of criteria can be considered when selecting the next time step with a primary objective of controlling solution error. Optimistically, use of the adaptive time step selection will result in a converged Newton step solution satisfying a user specified solution error tolerance and the solution time can again be advanced. Additional adaptive time stepping criteria are considered secondary relative to the error based criteria.

Foundationally, error based adaptive time stepping is based upon a solution error obtained using a predicted solution obtained from successful solves. Thus the time integration method often dictates how many solution steps into a simulation adaptive time stepping can begin. In most cases one must be able to obtain at least two successive solves using a fixed initial time step before adaptive time stepping begins. In many fluid simulations one routinely increases this number of constant steps to three or four using the PREDICTOR-CORRECTOR BEGIN AFTER STEP command line.

Certain model features like chemistry 36.1 will require that the phenomenological behavior within the current time step be considered when advancing the time step. We note that when adaptive time step selection...
is used, the subcycled chemistry timestep cannot be included directly in the adaptive time step selection as the chemistry time step will seldom satisfy the MINIMUM TIME STEP SIZE RATIO criteria 16.4.2. Here the minimum subcycled chemistry time step is modified to be comparable with other adaptive time step criteria 36.4. Additionally, the adaptive time step selection can also be overridden by user requests for output to occur at specific times.

One of the quantities used in the selection of time steps is the time step size ratio $r = \frac{\Delta t_{\text{new}}}{\Delta t_{\text{old}}}$, the ratio of a proposed (new) time step to the current (old) time step. Experience tells us that for a first and second order integrator bounding the time step size ratio, $0.5 \leq r \leq 2.0$. While maintaining these bounds usually minimizes the time truncation error, a user is permitted to manually alter these bounds. For the BDF2 integrator the suggested theoretical bounds are $0.5 \leq r \leq 2.14$, where the upper bound is strictly enforced to ensure stability of the integrator. The approach used in Aria to control time stepping strictly relevant to physical phenomena such as phase change is to artificially modify $r$ via a Phase Change Relaxation Factor.

### 16.4.1 Time Step Failure

Aria transient simulations are monitored for time step failure. For each solution time step we expect that the specified nonlinear tolerance will be achieved in the specified number of nonlinear iterations within a Newton step. If the convergence condition is not met the time step is said to have "Failed" and solution time will not be allowed to advance unless the user has explicitly stated that failed time steps will be accepted.
with the (ACCEPT SOLUTION AFTER MAXIMUM NONLINEAR ITERATIONS = TRUE) command line. If the non-converged solution is not accepted, a failed time step counter is incremented, the time step is reduced by the factor defined as FAILED TIME STEP SIZE RATIO (default 0.5) and the Newton step will be re-tryed with the revised time step. This process will be repeated until either a convergence condition is reached (convergence is achieved) or when the trial time step satisfies the FAIL TIME STEP WHEN TIME STEP SIZE RATIO IS BELOW criteria (no convergence). In the first case the time step is advanced and in the second case the simulation is terminated.

During phase change, the model using the energy equation for temperature employs a mushy zone formulation hence it is important that the temperature field values not bypass the range of the mushy zone so that the heat of fusion be accounted for. In this case an additional algorithm is used to detect when a nodal temperature has bypassed the mushy zone. When the mushy zone has been bypassed within a timestep, the Newton step will be marked as failed and the current time step will be scaled by a user specified PHASE CHANGE RELAXATION FACTOR in the same way that the failed time step size ratio is applied.

In some simulations a user may wish to interrupt the Newton step (fail the step) based upon the residual behavior. For many problems an aberrant behavior is recognized early on by noting the ratio of the current residual to the first residual. In cases where this ratio continues to grow it may be appropriate to terminate the Newton step and re-try with a different time step. This ratio can be specified by the user with the MAXIMUM LINEAR RESIDUAL RATIO.
16.4.2 Adaptive Time Step Selection

Adaptive time step selection consists of applying different criteria in the selection process. Each of the criteria are first described before linking them to the final selection procedure 16.4.3

Predictor Corrector

Let the predictor corrector tolerance, $\epsilon$, be specified from the input file.

$$\frac{\Delta t_{n+1}}{\Delta t_n} = \left( b \frac{\epsilon}{d_{n+1}} \right)^m$$

For 1st order time integrator $m = 1/2$, $b = 2.0$. For 2nd order time integrator $m = 1/3$

$$b = 3 \left( 1 + \frac{\Delta t_{n-1}}{\Delta t_n} \right)$$

$$d_{n+1} = \frac{1}{U_{max}} \left[ \sum_{i=1}^{N} \left( U_{i(n+1)} - U_{i(n+1)}^p \right)^2 \right]^{1/2}$$

so that the timestep based on predictor criteria is

$$\Delta t_{P_{red}} = \Delta t_{n+1} = \Delta t_n \left( b \frac{\epsilon}{d_{n+1}} \right)^m .$$

Max _CFL

In some instances the stability of a flow simulation can be governed by a Courant-Freidrichs-Levy (CFL) number, $CFL_{max}$ for the flow. If a user specifies that a CFL number be be carried out over the model then a candidate time step can be computed as $\Delta t_{maxCFL} = CFL_{lim}\Delta t_n/CFL_{max}$, where $CFL_{lim}$ is provided using the COURANT LIMIT command line. The CFL criteria possibly considered for $\Delta t$ are

- $\Delta t_{maxCFL}$ standard fluid
- $\Delta t_{maxCFL}$ standard fluid phase A
- $\Delta t_{maxCFL}$ standard fluid phase B
- $\Delta t_{maxCFL}$ standard fluid phase C
- $\Delta t_{maxCFL}$ level set.

Minimum Time Step Based Upon MAXIMUM TIME STEP SIZE RATIO

A time step size ratio involving the future time step and the current time step is $r = \Delta t_{n+1}/\Delta t_n$. Letting $r_{max}$ be the user specified maximum time step size ratio then the largest timestep achievable with $r_{max}$ is

$$\Delta t = \Delta t_n * r_{max} .$$

Minimum Time Step Based Upon MINIMUM ACCEPTABLE TIME STEP SIZE RATIO

A time step size ratio involving the adaptive time step and the current time step is $r = \Delta t_a/\Delta t_n$. Letting $r_{min}$ be the user specified minimum acceptable time step size ratio then we require that $r > r_{min}$ else the time step is considered to be failed, the proposed time step is too small.
Maximum Solution Increment

Let the user specified maximum increment in any solution DOF be \( (\Delta U)_{\text{Max}} \). From a previous successful solution we compute the maximum change in any DOF \( (\Delta U)_{\text{MaxSoln}} \) and use it to compute an estimate of its corresponding maximum derivative in the previous time step as \( U_{\text{max}} = (\Delta U)_{\text{MaxSoln}} / \Delta t_n \). For a new time step \( \hat{t}_n \) we might expect that the maximum value of \( U \) in the next step might reach \( (\Delta \hat{U})_{\text{max}} = \hat{t}_n U_{\text{max}} \). Now if \( (\Delta \hat{U})_{\text{max}} < (\Delta U)_{\text{Max}} \) we retain \( t_n \) otherwise we propose a new time step \( \Delta t_n = 0.95 * (\Delta U)_{\text{Max}} / U_{\text{max}} \) with the hope that the solution increment \( (\Delta U)_{\text{Max}} \) will not be exceeded in the next time plane. The same process is repeated for all solution DOF to produce a representative \( \Delta t_n \) for the given problem. Although adaptive time stepping with Solution Increment requires at least one successful solve here we require at least two successful solves since control of solution error is the primary objective.

16.4.3 Calculate Adaptive Timestep

We now proceed to select an adaptive time step \( t_a \) based upon the minimum value associated with:

- \( \Delta t_{\text{Pred}} \) \( 16.4.2 \)
- Courant Limit, \( \Delta t_{\text{cfl}} \) \( 16.4.2 \)
- Maximum Time Step Size, \( \Delta t_{\text{max}} \)
- Maximum Time Step Size Ratio, \( r_{\text{max}} \) \( 16.4.2 \)
- Maximum Solution Increment \( 16.4.2 \)
- Minimum Time Step Size

- Note: If chemistry is present, it will take precedence over all other candidates above

Let \( n_{tb} \) be the timeblock counter (step within the current timeblock).

Let \( n_{P\text{red}0} \) be the user specified number of constant steps that begin any timeblock (default minimum is 2).

Let the current timestep be \( \Delta t_n \) (INITIAL TIME STEP SIZE when starting a simulation), the adaptive timestep be \( \Delta t_a = \text{Real}_\text{Max} \), and predictor corrector timestep be \( \Delta t_{\text{Pred}} = \text{Real}_\text{Max} \).

Let \( \Delta t_{\text{max}} \) be the user specified MAXIMUM TIME STEP SIZE and \( \Delta t_{\text{min}} \) the MINIMUM RESOLVED TIMESTEP SIZE.

To begin the timestep selection we first compute the desired predictor corrector timestep \( \Delta t_{\text{Pred}} \) \( 16.4.2 \) with the understanding that we wish to obtain a nominal value \( \Delta t_{\text{Pred}} \) which may not be the same as \( \Delta t_{\text{Pred}} \).

if \( n_{tb} < n_{P\text{red}0} \)
compute predictor error
\( \Delta t_{\text{Pred}} = \Delta t_n \)
else
if \( \Delta t_{\text{Pred}} < \Delta t_{\text{min}} \) then
if \( \Delta t_n > \Delta t_{\text{min}} \) then
\( \Delta t_{\text{Pred}} = \Delta t_{\text{min}} \)
else
if \( \Delta t_{\text{Pred}} < \Delta t_n \) then
\( \Delta t_{\text{Pred}} = \Delta t_n \)
else
\[ \Delta t^{Predict} = \Delta t^{Pred}. \]

else
\[ \Delta t^{Predict} = \Delta t^{Pred}. \]

Finally, the candidate \( t_a \) value must then satisfy the MINIMUM ACCEPTABLE TIME STEP SIZE RATIO, \( r_{\text{min}} \), 16.4.2 to be accepted as a viable time step.

The overall time stepping procedure is demonstrated in the figure below.

![Time Step Selection Schematic](image)

**Figure 16.4.** Time Step Selection Schematic.

Time integration is currently defined in the same way for the various physics contained within Aria. The command lines used to specify parameters used in that definition are described in what follows.

### 16.5 Parameters For Aria Region

**Scope:** Parameters For

Begin Parameters For Aria Region **RegionName**

  - Allow Order Switch For HowMany Steps
  - Courant Limit \( =|\text{are}|\text{is} \) Courant_limit
  - Fail Time Step On Bad Aztec Solver Status
  - Fail Time Step When Time Step Size Ratio Is Below Ratio
  - Failed Time Step Size Ratio \( =|\text{are}|\text{is} \) Ratio
  - Initial Nonlinear Residual Tolerance For Time Step Control \( =|\text{are}|\text{is} \) tolerance
  - Initial Time Step Size \( =|\text{are}|\text{is} \) \( Dt \)
  - Interface Courant Limit \( =|\text{are}|\text{is} \) Interface Courant_limit

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Known Time Discontinuities \(\text{times}\)...

Limit Solution Increment Field name \(\text{Value}\ [\text{Action}]\)

Maximum Acceptable Linear Residual Ratio \(\text{Ratio}\)

Maximum Global Var Limit Encore name \(\text{Value}\ [\text{Action}]\)

Maximum Linear Residual Ratio \(\text{Ratio}\)

Maximum Solution Limit Field name \(\text{Value}\ [\text{Action}]\)

Maximum Time Step Size \(\text{Dt}\)

Minimum Global Var Limit Encore name \(\text{Value}\ [\text{Action}]\)

Minimum Resolved Time Step Size \(\text{Dt}\)

Minimum Solution Limit Field name \(\text{Value}\ [\text{Action}]\)

Minimum Time Step Size \(\text{Dt}\)

Predictor Order \(\text{order}\)

Predictor-Corrector Begin After Step \(\text{Step}\)

Predictor-Corrector Field Normalization Field name \(\text{Norm_type}\ [\text{Scaling}]\)

Predictor-Corrector Normalization \(\text{Normalization}\ [\text{Scaling}]\)

Predictor-Corrector Tolerance \(\text{Predictor_corrector_tolerance}\)

Slope Of Time Step Size \(\text{slope}\)

Stop When Initial Nonlinear Residual Is Below Tol

Time Integration Method \(\text{Time_integration_method}\)

Time Step Variation \(\text{Time_step_variation}\)

Use Initial Nonlinear Residual For Time Step Control \(\text{enable}\)

End

---

Summary  Defines region specific time stepping data

16.5.1 Allow Order Switch For

Scope: Parameters For Aria Region

<table>
<thead>
<tr>
<th>Allow Order Switch For HowMany Steps</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter</td>
</tr>
<tr>
<td>HowMany</td>
</tr>
</tbody>
</table>

Summary  Allow the switching of time integration order from a higher order to first order.

Description  This command allows for the switching of a higher order method to a first order method for a specified number of steps in order to improve the robustness of the simulation especially in the region of discontinuities such as chemistry shutoff. This command is mostly useful in the context of a SECOND_ORDER time integration method.

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16.5.2 Courant Limit
Scope: Parameters For Aria Region

\[ \text{Courant Limit} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>[ \text{Courant_limit} ]</td>
<td>real</td>
<td>0.0</td>
</tr>
</tbody>
</table>

Summary The Courant Number limit. A value of 0.0 denotes INACTIVE.

16.5.3 Fail Time Step On Bad Aztec Solver Status
Scope: Parameters For Aria Region

Summary OPT to fail time the time step when the AZTEC solver status is -2 (Numerical Breakdown) or -3 (Loss of Precision).

16.5.4 Fail Time Step When Time Step Size Ratio Is Below
Scope: Parameters For Aria Region

\[ \text{Fail Time Step When Time Step Size Ratio Is Below} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>[ \text{Ratio} ]</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specify lower bound for adaptive time step size ratio failure criteria.

If the adaptive time step size ratio falls below this value, Aria will fail the time step. This criteria is specified in terms of the size of the time step size ratio, defined as \( dt_{n+1}/dt_n \). When the adaptive time stepper senses the time step being cut back too much, the time step fails. The default is equal to the FAILED TIME STEP SIZE RATIO.

In simulations where the predicted time step is reduced substantially, a time step may fail repeatedly on this criterion. In that event the user may want to set this Ratio to a small positive value, possibly even zero.

16.5.5 Failed Time Step Size Ratio
Scope: Parameters For Aria Region

\[ \text{Failed Time Step Size Ratio} \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>[ \text{Ratio} ]</td>
<td>real</td>
<td>0.5</td>
</tr>
</tbody>
</table>

Summary Specifies the factor used to cut the time step size when a step fails.

When a time step fails, the time step size is reduced by this factor.
16.5.6 Initial Nonlinear Residual Tolerance For Time Step Control

**Scope:** Parameters For Aria Region

**Initial Nonlinear Residual Tolerance For Time Step Control {=|are|is} tolerance**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>tolerance</td>
<td>real</td>
<td>1.0</td>
</tr>
</tbody>
</table>

**Summary**
Set initial nonlinear residual tolerance for time step control.

**Description**
Sets the tolerance on the initial nonlinear residual to be used for time step control.

16.5.7 Initial Time Step Size

**Scope:** Parameters For Aria Region

**Initial Time Step Size {=|are|is} Dt**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dt</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specifies the initial time step size for the given parameter block. The initial time step size should always be specified for FIXED time stepping and for the starting time block of ADAPTIVE time stepping. When no other command lines affecting timestep are provided in the time step parameter block the time step will remain constant over the time extent of the parameter block.

16.5.8 Interface Courant Limit

**Scope:** Parameters For Aria Region

**Interface Courant Limit {=|are|is} Interface Courant_limit**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Interface Courant_limit</td>
<td>real</td>
<td>0.0</td>
</tr>
</tbody>
</table>

**Summary**
The Interface Courant Number limit. Applies to both diffuse level set and CDFEM problems. A value of 0.0 denotes INACTIVE.

16.5.9 Known Time Discontinuities

**Scope:** Parameters For Aria Region

**Known Time Discontinuities {=|are|is} times...**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>times</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specifies known time discontinuities in the solution that the adaptive time stepping procedure must capture.
Description | Allows user to specify certain times that the adaptive time stepper must solve at because the solution is known to have a discontinuity there. This is useful for problems where the boundary conditions change suddenly for example.
---|---

16.5.10 Limit Solution Increment

**Scope:** Parameters For Aria Region

Limit Solution Increment $Field_name \{=|are|is\} Value [ Action ]$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$Field_name$</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Value</td>
<td>real</td>
<td>REAL_MAX</td>
</tr>
</tbody>
</table>

**Summary** Provides user control for limiting maximum change in a chosen solution Field over a timestep. When action is PREDICT a time step estimation is computed based upon the predicted solution change. When action is FAIL_STEP the time step is still predicted but the the time step will also fail if the suggested solution increment is exceeded.

**Description**

This command allows one to roughly specify the maximum magnitude that a solution Field can change within a step. A solution increment is estimated based upon the maximum change in the Field in the previous timestep and the proposed adaptive timestep. When the specified value of maximum solution increment is exceeded, the proposed adaptive timestep is reduced in an attempt to limit the solution increment.

One command line should be supplied for each solution Field one wishes to restrict. The restriction is applied directly to the solution Field values, no scaling is invoked.

16.5.11 Maximum Acceptable Linear Residual Ratio

**Scope:** Parameters For Aria Region

Maximum Acceptable Linear Residual Ratio $\{=|are|is\} Ratio$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ratio</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary** Specifies a failure criterion for a nonlinear step based on the ratio of the final linear residual to the initial nonlinear residual.

The ratio must be greater than zero and less than one.

16.5.12 Maximum Global Var Limit

**Scope:** Parameters For Aria Region

Maximum Global Var Limit $Encore_name \{=|are|is\} Value [ Action ]$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$Encore_name$</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Summary  
This command allows the simulation to continue only when the given Encore variable is lower than a certain magnitude. Inclusion of the optional argument RETRY will cause the solution step to fail and the time step to be reduced. This sequence will be carried out a maximum of three steps, after which the simulation will be terminated.

Description  
This command allows the simulation to continue only when the given Encore variable is lower than a certain magnitude.

16.5.13 Maximum Linear Residual Ratio
Scope: Parameters For Aria Region

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ratio</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  
Specifies a failure criterion within a nonlinear step based on the ratio of a linear residual to the initial nonlinear residual. This criteria is not be used unless requested by the user.

16.5.14 Maximum Solution Limit
Scope: Parameters For Aria Region

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Field_name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Value</td>
<td>real</td>
<td>REAL_MAX</td>
</tr>
</tbody>
</table>

Summary  
Provides user control for limiting maximum value of a Field solution over a time step. Optional parameters are continue or terminate. When a step fails the step will be cut back and the optional behavior is invoked when three consecutive fails have occurred. Default behavior is to terminate.

Description  
This command allows one to restrict the maximum magnitude that a solution Field can achieve within a simulation by causing the time step to fail when the Field value exceeds a user specified magnitude. Often times the maximum value will be tied to an event that occurs at a particular magnitude of the Field. This command can be used to impose a different solution behavior when this magnitude of the Field is realized by forcing a failure of the solution step. When the step is failed due to solution limit then cutback of the time step is allowed to occur for two consecutive steps. If the time step is cut back a third time in succession, the specified action of the command will be invoked.

By default the simulation will terminate when the maximum value is reached. If the optional argument is "continue" then time step failure will be over-ridden and the solution sequence will continue, presumably evolving toward lower magnitude values of the Field. This behavior may enable one to overcome large reductions in time step at discontinuous events which may occur for maximum values of the Field.
16.5.15 Maximum Time Step Size
Scope: Parameters For Aria Region

Maximum Time Step Size \( = \) are is \( Dt \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( Dt )</td>
<td>real</td>
<td>REAL_MAX</td>
</tr>
</tbody>
</table>

Summary
Specifies the maximum time step size for this parameter block.
The time step will not be allowed to exceed this value, regardless of adaptive time step selection and regardless of time step requests for other regions (for loosely coupled simulations).

16.5.16 Maximum Time Step Size Ratio
Scope: Parameters For Aria Region

Maximum Time Step Size Ratio \( = \) are is \( Ratio \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( Ratio )</td>
<td>real</td>
<td>2.0/2.4</td>
</tr>
</tbody>
</table>

Summary
Specifies the maximum allowable growth rate of the adaptive time step size.
If the time step selection computes a new time step size that exceeds the previous step size by more than this amount, then the new step size is limited to the product of this factor and the previous step size.
The default is 2.0 for FIRST_ORDER and SECOND_ORDER integration methods.
For BDF2 a value of 2.4 is imposed to enforce stability of the integrator.

16.5.17 Minimum Global Var Limit
Scope: Parameters For Aria Region

Minimum Global Var Limit \( Encore \_ name \) \( = \) are is \( Value \) \[ Action \]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( Encore _ name )</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>( Value )</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
This command allows the simulation to continue only when the given Encore variable is higher than a certain magnitude. Inclusion of the optional argument, RETRY, will cause the solution step to fail and the time step to be reduced. This sequence will be carried out a maximum of three steps, after which the simulation will be terminated.

Description
This command allows the simulation to continue only when the given Encore variable is higher than a certain magnitude.

16.5.18 Minimum Resolved Time Step Size
Scope: Parameters For Aria Region
Minimum Resolved Time Step Size \(\text{=}|\text{are}|\text{is} \ D_t\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(D_t)</td>
<td>real</td>
<td>0.0</td>
</tr>
</tbody>
</table>

Summary

Specifies the minimum resolved time step size. If the time step size lies below this value, the step will not fail when the time step size ratio falls too low.

This setting serves as a floor value since the adaptive time stepper will not select a time step size below this value, and any predictor-corrector errors are neglected when the time step size falls below this value.

16.5.19 Minimum Solution Limit

Scope: Parameters For Aria Region

Minimum Solution Limit Field_name \(\text{=}|\text{are}|\text{is} \ Value \ [ \text{Action} \ ]\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Field_name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary

Provides user control for limiting minimum value of a Field solution over a time step. Optional parameters are continue or terminate. When a step fails the step will be cut back and and the optional behavior is invoked when three consecutive fails have occurred. Default behavior is to terminate.

Description

By default the solution minimum limit for all Fields is not active. This command allows one to restrict the minimum magnitude that a solution Field can achieve within a simulation by causing the time step to fail when the Field value goes below a user specified value.

See MAXIMUM SOLUTION LIMIT command for more info.

16.5.20 Minimum Time Step Size

Scope: Parameters For Aria Region

Minimum Time Step Size \(\text{=}|\text{are}|\text{is} \ D_t\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(D_t)</td>
<td>real</td>
<td>0.0</td>
</tr>
</tbody>
</table>

Summary

Specifies the minimum time step size for the given parameter block. The time step will not be allowed to fall below this value, regardless of adaptive time step selection and regardless of time step requests from other regions (for loosely coupled simulations).

16.5.21 Predictor Order

Scope: Parameters For Aria Region

Predictor Order \(\text{=}|\text{are}|\text{is} \ order\)
### Parameter: Predictor Order

**Summary:** Specifies the order of the extrapolation used to predict the solution in transient simulations. An order of 0 corresponds to no extrapolation, instead copying the previous solution into the predicted solution.

**Description:** This command sets the order of extrapolation in the predictor for transient problems. A value of 0 basically turns off the extrapolation, and instead copies to old solution to the predicted solution.

The `PREDICTOR ORDER` defaults to 0 for problems if Gas Dynamics equations are defined. Otherwise, it is set to the order of the time integrator for the current time block i.e 1 for first order and 2 for second order.

### 16.5.22 Predictor-Corrector Begin After Step

**Scope:** Parameters For Aria Region

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Step</td>
<td>integer</td>
<td>2 / 3</td>
</tr>
</tbody>
</table>

**Summary:** Specifies after which solution step the Predictor-Corrector timestep calculation will begin to be used in computing an adaptive timestep. Predictor error will always be computed but will not actually be used until the prescribed step is reached.

Default value is 2 for first-order time integration and 3 for second-order time integration. Thus for first-order time integration, a candidate predictor-corrector timestep could appear in step 3.

### 16.5.23 Predictor-Corrector Field Normalization

**Scope:** Parameters For Aria Region

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Field_name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** Allows user to change the predictor-corrector normalization type for a given field variable separately from the global option.

**Description:** Allows user to change the predictor-corrector normalization type for a given field variable separately from the global option.

Normalization types are:

- **NONE:** Scale by unity
- **MAX** (default): Scale by the maximum field value
- **USER:** Scale by a constant (user-supplied) value
16.5.24 Predictor-Corrector Normalization

Scope: Parameters For Aria Region

Summary
Provides the ability to specify how the predictor-corrector error is normalized. Currently supported values are MAX, NONE, and USER. If desired, L2 could be added easily. A possibly better method would be to allow user-specified scales for the variables but that’s not yet supported.

The reason for allowing NONE (or even better, a user-specified scale) is to better handle problems that involve something that is zero-based with finite changes. This can happen in ALE problems where the adaptive time stepper is overly restrictive initially when the displacements are all nearly zero, but growing to some finite value.

Default is MAX.

16.5.25 Predictor-Corrector Tolerance

Scope: Parameters For Aria Region

<table>
<thead>
<tr>
<th>Predictor-Corrector Tolerance {=</th>
<th>are</th>
<th>is} Predictor_corrector_tolerance</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Parameter</strong></td>
<td><strong>Value</strong></td>
<td><strong>Default</strong></td>
</tr>
<tr>
<td>Predictor_corrector_tolerance</td>
<td>real</td>
<td>0.001</td>
</tr>
</tbody>
</table>

Summary
Specifies the tolerance for the difference between the predicted solution and the implicitly solved corrector solution used sed in adaptive time step selection.

The adaptive time step selection formula is

\[
\Delta t_{n+1} = \Delta t_n \left( b \frac{\epsilon}{d_{n+1}} \right)^m
\]  

(16.2)

where \(\Delta t_n \equiv t_{n+1}-t_n\) is the time step size from the most recent solution, \(\Delta t_{n+1} \equiv t_{n+2}-t_{n+1}\) is the new time step size, \(\epsilon\) is the predictor-corrector tolerance and \(d_{n+1}\) is the norm of the difference between the predicted and actual solutions at time \(t_{n+1}\). For first order time integration \(m = 1/2\) and \(b = 2\). For second order time integration \(m = 1/3\) and \(b = 3(1 + \Delta t_{n-1}/\Delta t_n)\). See [30].

16.5.26 Slope Of Time Step Size

Scope: Parameters For Aria Region

| Slope Of Time Step Size {=|are|is} slope |
|----------------------------------------|
| **Parameter** | **Value** | **Default** |
| slope            | real                  | undefined |

Summary
Specifies the linear growth rate of the time step size for the given parameter block. The time step size will grow by this amount at each step unless limited by other factors.
16.5.27  Stop When Initial Nonlinear Residual Is Below
Scope: Parameters For Aria Region

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stop When Initial Nonlinear Residual Is Below <em>Tol</em></td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specify time block termination criteria based upon nonlinear residual value. If the initial nonlinear residual falls below this value, Aria will end the time block. This is useful for obtaining pseudo-transient solutions of steady-state problems.

16.5.28  Time Integration Method
Scope: Parameters For Aria Region

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time Integration Method {=</td>
<td>are</td>
<td>is} <em>Time_integration_method</em></td>
</tr>
</tbody>
</table>

Summary Specifies the order of time integration. Valid options are FIRST_ORDER, SECOND_ORDER and BDF2. Note that the stability criteria of BDF2 will restrict the timestep growth by imposing a MAXIMUM TIME STEP SIZE RATIO of 2.14.

16.5.29  Time Step Variation
Scope: Parameters For Aria Region

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time Step Variation {=</td>
<td>are</td>
<td>is} <em>Time_step_variation</em></td>
</tr>
</tbody>
</table>

Summary Specifies how the time step sizes are to be derived. Choose between FIXED and ADAPTIVE time step selection methods. When the time step is ADAPTIVE the time step is determined using a predictor-corrector error criteria.

16.5.30  Use Initial Nonlinear Residual For Time Step Control
Scope: Parameters For Aria Region

Summary Enable use of initial nonlinear residual as an error estimate to control timestep for loosely coupled problems.
When enabled this option will treat the initial non-linear residual as a proxy for the error in region-region convergence for loosely coupled problems. The time step will be controlled using a predictor-corrector like algorithm.

16.6 Other Timestep Related Commands

Because Aria can solve for many solution Fields it is inappropriate to include Field specific commands within the time block parameters. Because interaction of time stepping with particular simulation physics is often required, these interactions are handled via command lines appearing at the Region scope as illustrated below.

Begin Procedure My_Aria_Procedure

Begin Solution Control Description

  time stepping related commands

End

Begin Aria Region My_Region

  other physics timestep related commands

End

End

16.6.1 Phase Change Relaxation Factor

Scope:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prf</td>
<td>real</td>
<td>0.5</td>
</tr>
</tbody>
</table>

Summary: Allows a user to specify the phase change relaxation factor for ensuring that the current computation step does not miss a freeze or melt event. This factor must lie between 0.0 and 1.0. Over-relaxation or under-relaxation time step reduction factor when phase change occurs within a time step. Used in conjunction with a phase change specific heat material model.
16.7 Predictor Related Commands

Because Aria can solve for many solution Fields it is inappropriate to include Field specific commands within the time block parameters. Nevertheless, since interaction of time stepping with solution prediction is required, these interactions are handled via command lines appearing at the Region scope as illustrated below.

```plaintext
Begin Procedure My_Aria_Procedure

Begin Solution Control Description
  .
  time stepping related commands
  .
End
.

Begin Aria Region My_Region
  .
  predictor related commands
  .
End

End
.
```

16.7.1 Display Truncation Error For

Scope:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Solution_dof</em> [ With <em>HowMany</em> Entries ]</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Provides ability to request log file output of the maximum computed truncation error values.

**Description**

This command allows one to request the maximum magnitudes of predictor error at a small number of DOF to the log file. For vector Fields only the magnitude over all components is considered. Note that additional log file output can be IO intensive so use of this command line should be restricted to debugging of a model.

Default is 5 values.

16.7.2 Maximum Number Of Rebalances

Scope:
## Maximum Number Of Rebalances

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Count</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Specify the maximum number of times to perform a mesh rebalance.

### 16.7.3 Predictor Fields

**Scope:**

**Predictor Fields**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Field_names</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Specification of active fields to be used in predictor algorithm.

**Description**

Specifies which fields to examine or ignore in the algorithm for adaptive time step selection. Fields that are *not* predictor fields will not be predicted solutions for the first two time steps. This is important for fields like pressure and electrostatic potential which may exhibit large jumps in their solutions between the initial conditions and the first solutions owing to the absence of time dependent terms in their governing equations during startup.

The selected fields will contribute to the $d_{n+1}$ norm discussed in the **TIME TRUNCATION ERROR** time block command. This command line can be provided multiple times with cumulative results.

By default all fields are included. All field names following the optional "NOT" keyword will be excluded from the selection algorithm.
Chapter 17

Nonlinear Solution Strategy

17.1 Equation System Overview

In coupled physics problems one is tasked with solving a set of nonlinear coupled equations and ideally one would solve these equations as a fully-coupled monolithic system. In practice solving the set of equations monolithically may prove impractical owing to large linear system memory requirements and difficulty in locating a small radius of convergence. Thus in some cases one may wish to perform a segregated solve of the equations where each set of equations can be solved in a slightly different manner. This ability to solve subsets of the entire equation set is supported through a feature called Equation System by which criteria for solving a set of nonlinear equations can be applied to a select group of equations.

By default a single set of coupled equations is treated internally as an Equation System without need of additional input syntax. Here the solution strategy for the Equation System is solely determined by a collection of Nonlinear Solution Specifications 17.3. A typical setup for a single equation system might be

```
Begin Aria Region myRegion
  .
  Use Linear Solver solve_temperature
  .
  eq energy for temperature On block_1 using q1 with lumped_mass diff
  .
  nonlinear solution strategy = newton
  use dof averaged nonlinear residual
  maximum nonlinear iterations = 20
  nonlinear residual tolerance = 1.e-6
  .
End Aria Region myRegion
```

Simulations in which different subsets of the coupled equations are solved in a segregated manner require that each subset be explicitly defined as an Equation System with a corresponding solution strategy which may differ for each equation system. When multiple Equation Systems are present, the overall segregated solution is governed by iteration and convergence criteria at a global/outer level 17.2.
A typical setup for segregated solution of multiple equation systems might be

```
Begin Aria Region myRegion
  .
  Segregated global nonlinear residual tolerance = 6.0e-9
  .
  Begin Equation System temperature
    .
    eq energy for temperature On block_1 using q1 with lumped_mass diff
    Use Linear Solver solve_temperature
    .
    nonlinear solution strategy = newton
    use dof averaged nonlinear residual
    maximum nonlinear iterations = 20
    nonlinear residual tolerance = 1.e-6
    .
  End Equation System temperature
  .
  Begin Equation System species
    .
    eq species for species_0 On block_1 using q1 with diff
    Use Linear Solver solve_species
    .
    nonlinear solution strategy = newton
    use dof averaged nonlinear residual
    maximum nonlinear iterations = 10
    nonlinear residual tolerance = 1.e-8
    .
  End Equation System temperature
  .
End Aria Region myRegion
```

17.2 Global Solution For Equation Systems

The global solution for a set of Equation Systems is controlled by a set of segregated solve criteria. A simulation will advance to the next time step when these criteria are satisfied.
17.2.1 Segregated Global Minimum Convergence Rate

Scope:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>MinRate</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary

Set a minimum convergence rate based on the initial nonlinear residual of each equation system in the problem. At each segregated iteration the equation system with the largest initial residual is tested using \( r_N / r_{N-MinSteps} < MinRate^{MinSteps} \). If that condition is not met then the time step fails and is retried with a shorter dt.

17.2.2 Segregated Global Nonlinear Initial Residual Tolerance

Scope:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pt</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary

Convergence tolerance for global segregated nonlinear residual for the iteration. The iteration is considered finished when all of the individual equation systems have an initial nonlinear residual less than the given tolerance. This is cheaper than other tests that require recomputing the nonlinear residual. However, care should be taken since the initial nonlinear residual of the individual equation systems might not be a good measure of the final nonlinear residual of those systems due to deltas produced by other equation systems.

17.2.3 Segregated Global Nonlinear Number Of Steps

Scope:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number_of_steps</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary

Maximum number of segregated solution steps within a time step. Depending on the failValue the solution will either be accepted, or the time step will fail when the maximum number of steps is reached.

17.2.4 Segregated Global Nonlinear Relative Residual Tolerance

Scope:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pt</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Parameter | Value | Default
---|---|---
Pt | real | undefined

**Summary**

Convergence tolerance for global segregated nonlinear residual for the iteration. Satisfaction of this criterion is sufficient for the iteration to be considered finished. This line command can be used when the user desires to enforce a global relative residual tolerance for the entire segregated system that may be different from the individual equation systems value.

### 17.2.5 Segregated Global Nonlinear Residual Tolerance

**Scope:**

```
Segregated Global Nonlinear Residual Tolerance {=|are|is} Pt [ Minimum Steps {=|are|is} minStepsValue ]
```

**Parameter** | **Value** | **Default**
---|---|---
Pt | real | undefined

**Summary**

Convergence tolerance for global segregated nonlinear residual for the iteration. Satisfaction of this criterion is sufficient for the iteration to be considered finished. This line command can be used when the user desires to enforce a global residual enforcement for the entire segregated system that may be different from the individual equation systems value.

### 17.2.6 Segregated Global Nonlinear Target Number Of Steps Per Time Step

**Scope:**

```
Segregated Global Nonlinear Target Number Of Steps Per Time Step {=|are|is} target_iterations
```

**Parameter** | **Value** | **Default**
---|---|---
target_iterations | integer | undefined

**Summary**

Set a target number of segregated iterations per time step to use for time step control. If more than the specified number of iterations are required to converge a time step the next time step size will be decreased proportional to the ratio of the target number and the number taken. If a maximum number of allowed segregated iterations is set it is recommended to set the target to 1-2 fewer iterations to avoid an excessive number of failed time steps.
17.3 Nonlinear Solution Specifications

The general solution strategy used in Aria is to solve the requested system of equations for solution variables \( \phi \) incrementally. In the context of a Newton formulation then within the Newton step one will solve

\[
J \Delta \phi = -R(\phi)
\]

where \( J \) is the Jacobian matrix, \( \Delta \phi \) is the solution increment and \( R(\phi) \) is the system residual for the governing equations described in Chapter 5. This chapter describes basic nonlinear solution command line settings that influence how a converged solution will be obtained. For transient solutions these settings are used in combination with time integration settings described in Chapter 16 for advancing a solution to the next time step.

It is instructive here to review a typical log file output of the Aria solution summary in order to establish how a user might influence the solution procedure.

---

**Equation System AriaRegion->main:**

* Step : Transient, Strategy: NEWTON, Time: 2.20e+03, Step: 5.29e+00
* Mesh : Processor 0 of 1: 1253 of 1253 elems, 1363 of 1363 nodes
* Computing View Factors for enclosure space

<table>
<thead>
<tr>
<th>Step</th>
<th>Resid</th>
<th>Delta</th>
<th>Itns</th>
<th>Status</th>
<th>Resid</th>
<th>Asm/Slv Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>6.37e-02</td>
<td>1.10e+01</td>
<td>59</td>
<td>ok</td>
<td>8.23e-07</td>
<td>7.0e-03/2.0e-03</td>
</tr>
<tr>
<td>2</td>
<td>1.10e-04</td>
<td>8.18e-03</td>
<td>58</td>
<td>ok</td>
<td>1.19e-09</td>
<td>7.0e-03/2.0e-03</td>
</tr>
<tr>
<td>3</td>
<td>3.06e-07</td>
<td>8.81e-05</td>
<td>61</td>
<td>ok</td>
<td>3.68e-12</td>
<td>6.0e-03/2.0e-03</td>
</tr>
</tbody>
</table>

Termination reason: 8.80515e-05 < nonlinear_correction_tolerance(0.001)

---

In the log file output shown above, columns under the NONLINEAR heading are related to the command lines described in this chapter. The columns **Resid** and **Delta** represent L2 norms of an aggregate residual \( R^*(\Delta \phi) \) and solution increment vector \( \Delta \phi \) while the column denoted **Step** denotes the iteration within this Newton step. Columns under the LINEAR heading represent the number of linear solver iterations (**Itns**), the **Resid** column represents the linear system residual \( R^*(\Delta \phi) \). Here \( R^*(\Delta \phi) = J\Delta \phi + R(\phi) \), the residual for the assembled system of equations, is solved for \( \Delta \phi \) to the tolerance requested in the Linear Solver specifications 20. The column labeled **Status** indicates whether the linear solve completed successfully, **ok** or **fail**, within the allowable number of iterations defined in the Linear Solver specifications. The column labeled **Asm/Slv Time** contain assembly and solution times for the system of equations.

Given a current value of \( \phi \) then within each Newton step the residual \( R^*(\Delta \phi) \) is first evaluated using \( \Delta \phi = 0 \) and tested for convergence. Here the convergence conditions are satisfied when either the **NONLINEAR RESIDUAL TOLERANCE** or **NONLINEAR CORRECTION TOLERANCE** are achieved. If the solution is not yet converged, the system of equations is turned over to the equation solver where it is solved for a new \( \Delta \phi \) subject to the Linear Solver specifications. Then the Solver solution is used to update the nonlinear solution \( \phi = \phi_{old} + \Delta \phi \) and the process begins anew subject to a user supplied **MAXIMUM NONLINEAR ITERATIONS**.

Within the Newton step a solution is said to **FAIL** if it is not converged within **MAXIMUM NONLINEAR ITERATIONS** and this occurrence will be noted in the log file. For transient simulations one will normally repeat the solution procedure with a smaller time step as described in Chapter 16. In some cases the user may chose to override the **FAIL** condition using the **ACCEPT SOLUTION AFTER MAXIMUM NONLINEAR ITERATIONS** command line but one is cautioned that this option should be used with discretion as it may discredit any expectation of accuracy in transient results. For completeness, a summary of pass, fail, and fail but accepted steps are included at the end of the log file.
17.3.1 Accept Solution After Maximum Nonlinear Iterations

Scope:

Accept Solution After Maximum Nonlinear Iterations \(=|\text{are}|\text{is}\) \(\text{Bool}\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\text{Bool})</td>
<td>{false</td>
<td>true}</td>
</tr>
</tbody>
</table>

Summary: Specifies whether solution is considered to be converged after MAXIMUM NONLINEAR ITERATIONS even if none of the convergence criteria are satisfied.

17.3.2 Filter Nonlinear Solution

Scope:

Filter Nonlinear Solution For \(\text{AssociatedDoF FilterType}\) \(=|\text{are}|\text{is}\) \(\text{Values...}\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\text{AssociatedDoF})</td>
<td>(\text{string})</td>
<td>undefined</td>
</tr>
<tr>
<td>(\text{FilterType})</td>
<td>(\text{string})</td>
<td>undefined</td>
</tr>
<tr>
<td>(\text{Values})</td>
<td>(\text{string...})</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Restrict the value of a solution variable used in the nonlinear solver iterations of a solution step.

Description: The nonlinear solution can be restricted to a range with upper and lower limits using the FILTER_TYPE = RANGE while supplying two bounding values. To set an upper or lower bound use FILTER_TYPE = MAXIMUM or FILTER_TYPE = MINIMUM while supplying a single bounding value.

17.3.3 Maximum Nonlinear Iterations

Scope:

Maximum Nonlinear Iterations \(=|\text{are}|\text{is}\) \(N_p\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(N_p)</td>
<td>(\text{integer})</td>
<td>20</td>
</tr>
</tbody>
</table>

Summary: Number of allowable nonlinear iterations in one linear solution step.

Description: The solution step will terminate when the number of nonlinear iterations are exceeded. Default value = 20.

17.3.4 Minimum Nonlinear Solves

Scope:

Minimum Nonlinear Solves \(=|\text{are}|\text{is}\) \(N_p\)
Parameter Value Default
---
\(N_p\) integer 1

Summary  Minimum number of actual nonlinear solves that are required.

Description  Each nonlinear iteration involves first assembling the nonlinear residual and Jacobian matrix and second solving the linear system for a nonlinear correction that is applied to the estimate of the nonlinear solution.

The nonlinear solver will continue iterating until this minimum number of solves are performed. The default value is one (1) but setting this to zero can be useful in some situations.

### 17.3.5 Nonlinear Correction Tolerance

Scope:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nonlinear Correction Tolerance (=</td>
<td>are</td>
<td>is) (Pt)</td>
</tr>
</tbody>
</table>

Summary  Convergence tolerance of nonlinear correction norm for the iteration. Satisfaction of this criterion is sufficient for the iteration to be considered finished.

### 17.3.6 Nonlinear Relaxation Factor

Scope:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nonlinear Relaxation Factor (=</td>
<td>are</td>
<td>is) (Pr_f)</td>
</tr>
</tbody>
</table>

Summary  Weighting factor for fraction of a new nonlinear solution that will be applied to the linear solution update. Weighting factor must lie in the range 0.0 to 1.0.

### 17.3.7 Nonlinear Residual Ratio Tolerance

Scope:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nonlinear Residual Ratio Tolerance (=</td>
<td>are</td>
<td>is) (Pt)</td>
</tr>
</tbody>
</table>

Summary  Convergence tolerance for the ratio of the nonlinear residual to the initial nonlinear residual for the solution iteration. Convergence is satisfied if the ratio is less than this specified multiple of the initial nonlinear residual.
17.3.8 Nonlinear Residual Tolerance

Scope:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Pt)</td>
<td>real</td>
<td>(1.0\times10^{-6})</td>
</tr>
</tbody>
</table>

Summary: Convergence tolerance of nonlinear residual for the iteration. Satisfaction of this criterion is sufficient for the iteration to be considered finished.

17.3.9 Nonlinear Solution Strategy

Scope:

Summary: Specifies the nonlinear solution strategy.

Description: Nonlinear solution strategy must be NEWTON, NEWTON\_FINITE\_DIFFERENCE or POINT\_IMPLICIT. There is no default value so this line command is required.

For POINT\_IMPLICIT one must also set: ACCEPT SOLUTION AFTER MAXIMUM NONLINEAR ITERATIONS = TRUE and MAXIMUM NONLINEAR ITERATIONS = 1

17.3.10 Use Dof Averaged Nonlinear Residual

Scope:

Summary: Select whether to use a nonlinear residual normalized by the number of degrees of freedom instead of the raw global residual reported by the linear system.

Description: Specifies that the reported nonlinear residual is scaled (divided by) the number of DOF in the problem. In selecting appropriate values for the NONLINEAR RESIDUAL TOLERANCE users are cautioned that the average nonlinear residual reported values may be several orders of magnitude smaller than when using the global nonlinear residual. This option is provided to allow comparison with residual values from other applications.
Chapter 18

Writing User Plugins

18.1 About Plugins

Users are free to extend Aria’s library of material models, constitutive equations, boundary conditions, distinguishing conditions and source terms through the use of plugins. In order to do this, a user writes the C++ code to implement an Expression class. Thus plugins should not be confused with user_subroutines 33.19. Before delving into Aria plugins it’s worth reading section 42.2 and scanning section 42.5 for an introduction to Aria’s Expression system.

When a user supplies their own plugin it becomes a legitimate piece of Aria. Plugins have no performance penalty over Aria’s built-in functionality and plugins have no added restrictions over built-in Expression regarding what can and can’t be done. In fact, taking a user plugin and adding it to Aria proper (so that it becomes “owned” by Aria) is a fairly straightforward process.

It’s worth re-stating here that since Aria employs a Newton formulation, many of the algorithms make use of DOF sensitivities, i.e., Aria often needs to know the derivative of your model with respect to all of the unknowns in the problem. There are three ways to supply the sensitivities in Aria: manually code them, use a numerical finite-difference function, or use (forward) automatic differentiation (FAD). The FAD method is ideal for plugins because the sensitivities are are performed analytically and exact, though there may be a small performance degradation. However, if only a few models use FAD there may be no measurable performance penalty. For that reason, we’ve designed our plugin system to use FAD by default. If one has different needs, please contact the user support group sierra-helpsandia.gov for help.

18.2 Compiling and Using Plugins with Dynamically Loaded Libraries

The easiest and most efficient way to compile and use plugins with Aria is to compile them into a dynamically loaded library that a pre-built aria executable is able to load at runtime.

18.2.1 Compiling a Plugin into a Shared Library

The easiest way to compile your plugin into a share library (.so file) is to use the jamsub tool. Within SNL, this tool is normally located in the /sierra/Dev/tools/ directory. It’s pretty simple to use:

$ jamsub -P aria -sierradir /path/to/sierra/project

If you happen to be working in a Sierra project that contains the Aria you’d like to build against then one can skip the -sierradir option. You can use the -s option to override the system for which you want
to build; see the help (-h) for more options.

This command will find all of the .C files in the current directory and compile them into a file of the same name but with the .so extension. Note that you do not have to put your .C files in the Aria source directories – they can live with your problem input files in any working directory.

In the event one is not building on a standard Linux machine, the command line must also reference the target platform as follows:

For code versions before 4.26,

$ module load sierra
$ jamsub -P aria -sierradir /path/to/sierra/project/target_code_version

this will compile all plugins in the current directory.

For code versions 4.26 and beyond,

$ module load sierra-devel
$ module load sierra-apps/target_code_version
$ sierra -make aria -i input_filename.i

here the input file is first scanned and only the needed plugins will be compiled.

Running the above command line with an invalid platform name will provide the available platforms for which one can build an Aria executable.

18.2.2 Pointing to a Plugin from your Aria Input File

Begin Sierra My_Sierra_Job
  Load User Plugin file ./My_Density_Plugin.so
  Load User Plugin file ./My_Viscosity_Plugin.so
  
18.2.3 Running Aria with Dynamically Loaded Plugins

The Aria executable is built to accommodate dynamically loaded libraries. Execution when using plugins is handled in the same way as when plugins are not being used:

$ sierra aria -i use_my_density.i -x /path/to/sierra/project

18.3 An Important Note About Model Names

In order to avoid name clashes, model names always end with the generic name of the quantity they provide. So, density model names always end in _DENSITY, viscosity model names end in _VISCOSITY, source model
names end in \_SOURCE, etc. For material models, it’s easy to know what that ending is because it’s the same as the left-hand side of the “\_” sign in the material input block (with spaces replaced by an underscore “\_”). Clearly, it’s important to keep this in mind when naming and referring to your plugin model or else Aria won’t be able to find it properly.

In this example, we’ll name our density \_MODEL\_DENSITY (recall, it must end in \_DENSITY) so in the input file we’ll have to refer to it as \_MODEL since the ending is automatically added.

### 18.4 The Input File

Plugins can be referenced in the input file similar to the way Aria’s built-in Expressions are referenced. The name of the model is just whatever your plugin is named without the file extension, e.g., \_MODEL,

```
Begin Aria Material Kryptonite
  Density = My_Model a=1.0 b=-0.01
  ...
End
```

It’s important to note that the plugin name must not conflict with model names used internally by Aria (the outcome would be, at least to users, ambiguous). Aria will verify this and produce an error if there’s a name clash.

### 18.5 Example Plugin Code: \_Density.C

In this section we’ll write the code to supply a density function which is a cubic polynomial in temperature ($T$),

$$
\rho = a + bT + cT^2 + dT^3.
$$

Normally, writing C++ code requires using a header (.h) file and an implementation (.C) file but since no other code needs to see our class definition, we can skip the header file and place the definition right in the .C file.

The first thing we need to do is include a header file which will give us all we need to write an Expression. Since our plugin will live in isolation, we’ll also add a using declaration to make life easier on ourselves; without it we’d have to declare the namespaces or prepend \_Aria:: to lots of data types. So far, we have this:

```c
#include <Aria_Plugin_Expression.h>
using namespace sierra::Aria;
```

Next we need the basic declaration of our class. This tells the compiler which methods and data members we want to have. Here’s ours:

```c
// Class definition -- could go in a header file.
class My_Density : public FAD_Expression
{
 public:
  My_Density(Expression_Manager * const manager,
```
const sierra::Identifier & expr_model_name,
const Subdomain_Tag & subdomain_tag,
const Int & subindex,
const Phase_Label & phase_label,
const sierra::String & params);
virtual ~My_Density() {}
virtual void compute_FAD_values();
private:
  Real a;
  Real b;
  Real c;
  Real d;

  FAD_Expression_Handle temperature;
};

This is mostly boiler-plate code. Here we declare a constructor and an empty destructor and the method
needed for computing our density’s values. We also add some private data to store our polynomial coefficients.

Now it’s time to write some code. First, we write the constructor which tells Aria, in a generic sense,
what we provide (density), what we depend on (temperature) and what parameters we require from the user
\((a, b, c \text{ and } d)\).

My_Density::My_Density(Expression_Manager * const manager,
  const sierra::Identifier & expr_model_name,
  const Subdomain_Tag & subdomain_tag,
  const Int & subindex,
  const Phase_Label & phase_label,
  const sierra::String & params) :
  FAD_Expression(manager,subdomain_tag,subindex,phase_label,params)
{
  my_tensor_order = 0;
  my_expression_tag = Expression_Tag(EXPR::DENSITY,NO_OP,subindex,phase_label);
  my_expression_model_name = expr_model_name;

  // List the expressions that are required for this model.
  const Expression_Tag temperature_tag(EXPR::TEMPERATURE,NO_OP,subindex,phase_label);
  add_prereq(temperature_tag,temperature);

  // Get my model parameters.
  a = b = c = d = 0.0;
  get_optional_param("A",a);
  get_optional_param("B",b);
  get_optional_param("C",c);
  get_optional_param("D",d);

  if(a == 0.0 && b == 0.0 && c == 0.0 && d == 0.0)
  {
    throw sierra::RuntimeUserError() << "ERROR: All MY_MODEL_DENSITY parameters are zero.";
  }

  // Make myself known to the manager.
  register_myself();

  // ...
The `my_tensor_order` tells Aria what kind of field your Expression creates, viz. scalar, vector or tensor, tensor order 0, 1 or 2. If left unspecified, the default type is scalar (`my_tensor_order == 0`). Your expression also has a variable called `my_tensor_dimension` which is dimensionality of each tensor order. By default, `my_tensor_dimension` is set to the physical dimension of the problem, i.e., 2 for 2D and 3 for 3D. This variable is also set for scalar fields. Combined, these two variables define the number of values required to fully specify your Expression at a point: `pow(my_tensor_dimension, my_tensor_order)`. These also define the expected signature of the `values` data used below.

Next, we write the code that implements our density function. **Important note:** the `FAD_values` arrays are always initialized to zero before this method is called. Here `FAD` type variables are used for non-constant values related to the `FAD_values` calculation.

```cpp
void My_Density::compute_FAD_values()
{
    for(Int point=0; point < num_points; ++point)
    {
        const FAD_Type & T = temperature(point);
        FAD_values(point) = a + T*(b + T*(c + T*(d))));
    }
}
```

Note that since our Expression is a scalar in this example, the signature of `FAD_values` is `FAD_values(point)`. If our result was a vector or tensor, the signature would be `FAD_values(point,r)` and `FAD_values(point,r,s)`, respectively, where `0 <= r,s < my_tensor_dimension`.

The last thing we need to do is the actual plugin step. This one line of code,

```cpp
ExprPluginFactory<My_Density> my_density_creator("MY_MODEL_DENSITY");
```

takes care of making your Expression known to Aria. The quoted string is the string that is used in your input file (except for the ending part, see section 18.3. If this string has any spaces in it, Aria will automatically replace them with an underscore.

For completeness the whole plugin code is given here.

```cpp
#include <Aria_Plugin_Expression.h>

using namespace sierra::Aria;

// Class definition -- could go in a header file.
class My_Density : public FAD_Expression
{
public:
    My_Density(Expression_Manager * const manager,
                  const sierra::Identifier & expr_model_name,
                  const Subdomain_Tag & subdomain_tag,
                  const Int & subindex,
                  const Phase_Label & phase_label,
                  const sierra::String & params);
    virtual ~My_Density() {}  
    virtual void compute_FAD_values();
```
private:
    Real a;
    Real b;
    Real c;
    Real d;

    FAD_Expression_Handle temperature;
};

My_Density::My_Density(Expression_Manager * const manager,
                        const sierra::Identifier & expr_model_name,
                        const Subdomain_Tag & subdomain_tag,
                        const Int & subindex,
                        const Phase_Label & phase_label,
                        const sierra::String & params) :
    FAD_Expression(manager,subdomain_tag,subindex,phase_label,params)
{
    my_tensor_order = 0;
    my_expression_tag = Expression_Tag(EXPR::DENSITY,NO_OP,subindex,phase_label);
    my_expression_model_name = expr_model_name;

    // List the expressions that are required for this model.
    const Expression_Tag temperature_tag(EXPR::TEMPERATURE,NO_OP,subindex,phase_label);
    add_prereq(temperature_tag,temperature);

    // Get my model parameters.
    a = b = c = d = 0.0;
    get_optional_param("A",a);
    get_optional_param("B",b);
    get_optional_param("C",c);
    get_optional_param("D",d);

    if(a == 0.0 && b == 0.0 && c == 0.0 && d == 0.0)
    {
        throw sierra::RuntimeUserError() << "ERROR: All MY_MODEL_DENSITY parameters are zero."
    }

    // Make myself known to the manager.
    register_myself();
}

void My_Density::compute_FAD_values()
{
    for(Int point=0; point < num_points; ++point)
    {
        const FAD_Type & T = temperature(point);
        FAD_values(point) = a + T*(b + T*(c + T*(d)));
    }
}

ExprPluginFactory<My_Density> my_density_creator("MY_MODEL_DENSITY");
18.6 Testing Your Plugin

There are two good tests you can perform to test your plugin. The first is to run Aria in debug mode. To do this, add `-o dbg` to your `build` command line to build a debug executable. Then, add `-d` to your `sierra` command line to run the debug executable. In debug mode Aria will, among other things, perform bounds checking on your `FAD_MDArray` objects like `values(...)`, `dself(...)`, and `sens(...)` in this example.

Secondly, if you hand-code your Newton sensitivities (not done in this example), you can test the coding of your sensitivities by adding `-arialog sens_check` to your `sierra` command line. This will cause Aria to compare the computed sensitivity values with numerical approximations. The sensitivity checker will cause your code to run slower but it will work in either debug or optimized mode. Aria’s sensitivity checker is designed to only report discrepancies that have a high probability of being true errors so it’s possible that it may miss some small errors. However, reported errors are most probably real.

Oftentimes the easiest way to debug your code is to print information to the screen. Aria provides a facility to support this. To print information to the log file, use the `arialog` C++ output stream. For example,

```c++
for(Int point=0; point < num_points; ++point)
{
    ...
    FAD_values(point) = ...
    arialog.m(LOG_PLUGIN) << "value(" << point << ") = " << value(point) << endl;
}
```

Then, you can activate this output by adding the option `-arialog plugin` to your `sierra` command line. If you leave off the `.m(LOG_PLUGIN)` part then it will always write your output to the log file. There’s a performance penalty for having this code option present (even if you don’t turn the logging on) so you probably want to remove it once you’re done debugging.
Chapter 19

Dynamic Load Balancing

See, also, the Zoltan homepage [31], the Zoltan User’s Guide [32] and an overview of Zoltan [33].

19.1 ENABLE REBALANCE

ENABLE REBALANCE WITH THRESHOLD = REAL USING ZOLTAN PARAMETERS STRING

Description

Enables load balancing for parallel simulations.

Summary

This command causes Aria to occasionally redistribute the mesh across the processors in a parallel run in order to (hopefully) balance the work. In order to perform this balancing, Aria must supply a “load” for each element on each processor to the Zoltan library. See the REBALANCE LOAD MEASURE command (19.2) for load measuring options.

The loads can be output to the results database by adding the following line to the RESULTS OUTPUT block in the input file:

Element Variables = rebalLoadMeasure as Load

The number supplied as a threshold determines how far out of balance the load can become before load balancing is performed and the STRING argument names the Zoltan parameter block to use. Hence, a ZOLTAN PARAMETERS block must also be supplied in the input file.

Parent Block(s) ARIA_REGION

19.2 REBALANCE LOAD MEASURE

REBALANCE LOAD MEASURE = STRING

Description

Selects the method for measuring element loads for rebalancing.

Summary

Valid options are ELEMENT (default), PROCESSOR and CONSTANT.

ELEMENT : (default) assigns an element weight equal to the total cost of assembling the element for a timestep divided by the number of nonlinear iterations.

PROCESSOR : assigns same weight to all elements on a processor equal to the average cost of assembling an element for a timestep divided by the number of nonlinear iterations.

CONSTANT : assigns the same weight to each element, and is useful for regression testing.
19.3 **MAXIMUM NUMBER OF REBALANCES**

**MAXIMUM NUMBER OF REBALANCES = INT**

**Description**
Disables dynamics mesh rebalancing after the specified number of mesh rebalances have been performed.

**Summary**
The number of rebalances only increments if elements are actually migrated across processors. By default, Aria will rebalance as many times as needed.

19.4 **REBALANCE TIME STEP FREQUENCY**

**REBALANCE TIME STEP FREQUENCY = INT**

**Description**
Causes Aria to only examine the load balance every $N$ steps, where $N$ is the number supplied by this command line.

**Summary**
By default, Aria will check the load balance every time step.

19.5 **Zoltan Parameters**

**Scope:** Sierra

```
Begin Zoltan Parameters Parameter_block_model
   Algorithm Debug Level {=|are|is} Level
   Check Geometry {=|are|is} Bool
   Debug Memory {=|are|is} Level
   Debug Processor Number {=|are|is} Proc
   Deterministic Decomposition {=|are|is} Bool
   Imbalance Tolerance {=|are|is} Tol
   Keep Cuts {=|are|is} Bool
   Load Balancing Method {=|are|is} LoadBalancingMethod
   Lock Rcb Directions {=|are|is} Bool
```

### 19.5.1 Algorithm Debug Level

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Algorithm Debug Level</td>
<td>Level</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
- Default: Algorithm Debug Level = 1 Zoltan Equivalence: Algorithm Debug Level = RCB_OUTPUT_LEVEL = RIB_OUTPUT_LEVEL
- Flag controlling the amount of timing and diagnostic output the specific algorithm produces.
- Valid values are 0 (no output), 1 (print summary), or 2 (print data for each processor).

**Examples**
- Algorithm Debug Level = 0
- Algorithm Debug Level = 1
- Algorithm Debug Level = 2

### 19.5.2 Check Geometry

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Check Geometry</td>
<td>Bool</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
- Default: Check Geometry = true Zoltan Equivalence: Check Geometry = CHECK_GEOM
- Flag controlling the invocation of input and output error checking. Valid values are false (don’t do checking), or true (do checking). Default is true.

**Examples**
- Check Geometry = false
- Check Geometry = true
19.5.3 Debug Memory
Scope: Zoltan Parameters

Debug Memory \{=|are|is\} Level

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Level</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Default: Debug Memory = 1 Zoltan Equivalence: Debug Memory = DEBUG_MEMORY
Integer indicating the amount of low-level debugging information about memory-allocation should be kept by Zoltan’s Memory Management utilities. Valid values are 0, 1, 2, and 3
examples Debug Memory = 0 Debug Memory = 1 Debug Memory = 2 Debug Memory = 3

19.5.4 Debug Processor Number
Scope: Zoltan Parameters

Debug Processor Number \{=|are|is\} Proc

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Proc</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Default: Debug Processor = 0 Zoltan Equivalence: Debug Processor Number = DEBUG_PROCESSOR
Processor number from which trace output should be printed when the zoltan parameter, DEBUG_LEVEL, is set to 5.
example Debug Processor Number = 2

19.5.5 Deterministic Decomposition
Scope: Zoltan Parameters

Deterministic Decomposition \{=|are|is\} Bool

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bool</td>
<td>{false</td>
<td>true}</td>
</tr>
</tbody>
</table>

Summary
Default: Deterministic Decomposition = true Zoltan Equivalence: Deterministic Decomposition = DETERMINISTIC
If this value is set to true, Zoltan’s computation of the new decomposition is deterministic: i.e., executing the same algorithm with the same input on the same number of processors always produces the same results.
When this parameter is false, message order and other factors may cause variations in decompositions even under identical operating conditions.
It is highly recommended not to change the default of true.
examples Deterministic Decomposition = true Deterministic Decomposition = false
19.5.6 Imbalance Tolerance

Scope: Zoltan Parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tol</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Default: Imbalance Tolerance = 1.1 Zoltan Equivalence: Imbalance Tolerance = IMBALANCE_TOL Greater than or equal to 1.0.
The amount of load imbalance the partitioning algorithm should deem acceptable. The load on each processor is computed as the sum of the weights of objects it is assigned. The imbalance is then computed as the maximum load divided by the average load. A value for IMBALANCE_TOL of 1.2 indicates that should not exceed 1.2 example Imbalance Tolerance = 1.2

19.5.7 Keep Cuts

Scope: Zoltan Parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bool</td>
<td>{false</td>
<td>true}</td>
</tr>
</tbody>
</table>

Summary
Default: Keep Cuts = true Zoltan Equivalence: Keep Cuts = KEEP_CUTS Should information about the cuts determining the RCB, RIB, or HSFC type decomposition be retained? It costs a bit of time to do so, but this information is necessary if application wants to add more objects to the decomposition via calls to Zoltan_Point_Assign or to Zoltan_Box_Assign. Valid values are: false (don’t keep cuts), or true (keep cuts).
examples Keep Cuts = false Keep Cuts = true

19.5.8 Load Balancing Method

Scope: Zoltan Parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>LoadBalancingMethod</td>
<td>{hilbert space filling curve</td>
<td>recursive coordinate bisection</td>
</tr>
</tbody>
</table>

Summary
Default: Load Balancing Method = Recursive Coordinate Bisection Zoltan Equivalence: Load Balancing Method = LB_METHOD Dynamic load rebalancing partitioning decomposition method. Valid values are: Recursive Coordinate Bisection (RCB), Recursive Inertial Bisection (RIB), Hilbert Space Filling Curve (HSFC). (Carter Edward’s HFFC algorithm)
examples Load Balancing Method = Hilbert Space Filling Curve
19.5.9 Lock Rcb Directions

Scope: Zoltan Parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lock Rcb Directions</td>
<td>Bool</td>
<td>undefined</td>
</tr>
<tr>
<td></td>
<td>{false</td>
<td>true}</td>
</tr>
</tbody>
</table>

Summary

Default: Lock RCB Directions = false
Zoltan Equivalence: Lock RCB Directions = RCB_LOCK_DIRECTIONS

Flag that determines whether the order of the directions of the cuts is kept constant after they
are determined the first time RCB is called. Valid values are: false (Don’t lock directions),
or true (lock directions).

examples
Lock RCB Directions = false
Lock RCB Directions = true

19.5.10 Over Allocate Memory

Scope: Zoltan Parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Over Allocate Memory</td>
<td>Alloc</td>
<td>undefined</td>
</tr>
<tr>
<td></td>
<td>real</td>
<td></td>
</tr>
</tbody>
</table>

Summary

Default: Over Allocate Memory = 1.1
Zoltan Equivalence: Over Allocate Memory = RCB_OVERALLOC = RIB_OVERALLOC
Greater than or equal to 1.0.

The amount by which to over-allocate temporary storage arrays for objects within the algo-

rithm when additional storage is due to changes in processor assignments. Valid values are:
1.0 (no extra storage allocated), 1.5 (50

examples
Over Allocate Memory = 1.0
Over Allocate Memory = 1.5

19.5.11 Rectilinear Rcb Blocks

Scope: Zoltan Parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rectilinear Rcb Blocks</td>
<td>Bool</td>
<td>undefined</td>
</tr>
<tr>
<td></td>
<td>{false</td>
<td>true}</td>
</tr>
</tbody>
</table>

Summary

Default: Rectilinear RCB Blocks = false
Zoltan Equivalence: Rectilinear RCB Blocks = RCB_RECTILINEAR_BLOCKS

Flag controlling the shape of the resulting regions. If this option is specified, then when a cut
is made, all of the dots located on the cut are moved to the same side of the cut. The resulting
regions are then rectilinear. When these dots are treated as a group, then the resulting load
balance may not be as good as when the group of dots is split by the cut. Valid values are:
false (move dots individually), or true (move dots in groups).

examples
Rectilinear RCB Blocks = false
Rectilinear RCB Blocks = true
19.5.12 Renumber Partitions
Scope: Zoltan Parameters

Renumber Partitions {=|are|is} Bool

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bool</td>
<td>{false</td>
<td>true}</td>
</tr>
</tbody>
</table>

Summary
Default: Renumber Partitions = true Zoltan Equivalence: Renumber Partitions = REMAP
Flag to indicate whether to renumber partitions to maximize overlap between the old decomposition and the new decomposition (to reduce data movement from old to new decompositions). Requests for remapping are ignored when, in the new decomposition, a partition is spread across multiple processors or partition sizes are specified using "Set Partition Sizes".
examples Renumber Partitions = false Renumber Partitions = true

19.5.13 Reuse Cuts
Scope: Zoltan Parameters

Reuse Cuts {=|are|is} Bool

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bool</td>
<td>{false</td>
<td>true}</td>
</tr>
</tbody>
</table>

Summary
Default: Reuse Cuts = true Zoltan Equivalence: Reuse Cuts = RCB_REUSE
Flag to indicate whether to use previous cuts as initial guesses for the current RCB invocation. Valid values are: false (don’t use previous cuts), or true (use previous cuts).
examples Reuse Cuts = false Reuse Cuts = true

19.5.14 Set RCB Directions
Scope: Zoltan Parameters

Set RCB Directions {=|are|is} RCBSetDirections

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>RCBSetDirections</td>
<td>{do not order cuts</td>
<td>xyz</td>
</tr>
</tbody>
</table>

Summary
Default: Set RCB Directions = Do not order cuts Zoltan Equivalence: Set RCB Directions = RCB_SET_DIRECTIONS
Flag permits the order of cuts to be changed so that all of the cuts in any direction are done as a group. The number of cuts in each direction is determined and then the value of the parameter is used to determine the order that those cuts are made in. When 1D and 2D problems are partitioned, the directions corresponding to unused dimensions are ignored. Valid values are: Do not order cuts, xyz, xzy, yxz, yzx, zxy, or zyx.
examples Set RCB Directions = Do not order cuts Set RCB Directions = xyz Set RCB Directions = xzy Set RCB Directions = yxz Set RCB Directions = yzx Set RCB Directions = zyx Set RCB Directions = zxy

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19.5.15 Timer
Scope: Zoltan Parameters

Summary
Default: Timer = wall Zoltan Equivalence: Timer = TIMER
The timer with which you wish to measure time. Valid values are WALL and CPU.
examples Timer = wall Timer = cpu

19.5.16 Zoltan Debug Level
Scope: Zoltan Parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zoltan Debug Level</td>
<td>Level</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Default: Zoltan Debug Level = 1 Zoltan Equivalence: Zoltan Debug Level = DEBUG_LEVEL
0 Quiet mode; no output unless an error or warning is produced. 1 Values of all parameters set by Zoltan _Set_Param and used by the load-balancing method. 2 Timing information for Zoltan’s load-balancing and auto-migration routines. 3 Timing information within load-balancing algorithms (support by algorithms is optional). 4 5 Trace information (enter/exit) for major Zoltan interface routines (printed by the processor specified by the DEBUG_PROCESSOR parameter). 6 Trace information (enter/exit) for major Zoltan interface routines (printed by all processors). 7 More detailed trace information in major Zoltan interface routines. 8 List of objects to be imported to and exported from each processor. 9 10 Maximum debug output; may include algorithm-specific output. example Zoltan Debug Level = 5

19.5.17 Rcb Max Aspect Ratio
Scope: Zoltan Parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rcb Max Aspect Ratio</td>
<td>Ratio</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Default: rcb max aspect ratio = 10 Zoltan Equivalence: RCB_MAX_ASPECT_RATIO = ratio
The maximum aspect ratio for a given part
examples rcb max aspect ratio = 5

19.5.18 Rcb Recompute Box
Scope: Zoltan Parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rcb Recompute Box</td>
<td>Bool</td>
<td>undefined</td>
</tr>
</tbody>
</table>

650
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bool</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Default: `rcb recompute box = 0` Zoltan Equivalence: `RCB_RECOMPUTE_BOX = bool`

0, leave box computations off 1, turn box computations on

eamples `rcb recompute box = 1`
Chapter 20

Linear Solver Reference

20.1 Overview

For most problems Aria makes use of the Trilinos and Aztec equation solvers. Another solver GDSW, geared specifically toward incompressible flows is also available. These solvers provide an object-oriented interface to allow for flexible construction matrix and vector algorithms. Within Aria one provides the solver parameters at the domain scope and then references the solver within the Aria region. See, also, the Trilinos parameters online reference.

Within the input file linear solver command block will appear at the Domain scope as illustrated below. Note that different linear solvers can be used for different equation sets. Furthermore a solver command block can appear in the input but not be used.

Begin Sierra

. Begin Trilinos Equation Solver first_solver
  . Trilinos linear solver commands
  . End

. Begin Aztec Equation Solver second_solver
  . Aztec linear solver commands
  . End

. Begin Aztec Equation Solver third_solver
  . Aztec linear solver commands
  . End

. Begin Procedure My_Aria_Procedure
  . Begin Aria Region My_Region
    . use linear solver first_solver
  . End
20.2 Trilinos Equation Solver

Scope: Sierra

Begin Trilinos Equation Solver SolverName
  Bc Enforcement {=|are|is} BcEnforcement
  Debug Output Level {=|are|is} Level
  Debug Output Path {=|are|is} DebugOutput
  Determine Sharing {=|are|is} FeiDetermineSharing
  Fei Error Behavior {=|are|is} FeiErrorBehavior
  Fei Output Level {=|are|is} FeiOutputLevels
  Ilu Fill {=|are|is} Fill_level
  Ilu Threshold {=|are|is} Threshold
  Matrix Format {=|are|is} MatrixFormat
  Matrix Reduction {=|are|is} AztecReductionType
  Matrix Scaling {=|are|is} MatrixScaling
  Matrix Viewer {=|are|is} Machine:port
  Maximum Iterations {=|are|is} Max_iters
  Num Levels {=|are|is} Num_levels
  Param-Bool Parameter_name Value Bool
  Param-Int Parameter_name Value integer_value
  Param-Real Parameter_name Value Real_value
  Param-String Parameter_name Value String_value
  Polynomial Order {=|are|is} Order
  Preconditioner Subdomain Overlap {=|are|is} Overlap
  Preconditioning Method {=|are|is} TrilinosPrecondMethods
  Preconditioning Steps {=|are|is} Steps
  Residual Norm Scaling {=|are|is} AztecResidualNormScaling
  Residual Norm Tolerance {=|are|is} ResidualNormTolerance
  Restart Iterations {=|are|is} Restart_iters
  Row Ordering {=|are|is} RowOrdering
  Shared Ownership Rule {=|are|is} SharedOwnershipRule
  Solution Method {=|are|is} TrilinosSolverMethods

End
Summary: A set of solver parameters for Trilinos equation solver.

20.2.1 Bc Enforcement

Scope: Trilinos Equation Solver

Bc Enforcement {=|are|is} BcEnforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>BcEnforcement</td>
<td>{exact</td>
<td>exact_no_column_mod</td>
</tr>
</tbody>
</table>

Summary: Controls the way Dirichlet BCs are enforced.

Description: Valid values for this line-command are contained in the BcEnforcement enum.

20.2.2 Debug Output Level

Scope: Trilinos Equation Solver

Debug Output Level {=|are|is} Level

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Level</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Output level for debugging. Generally 0 means no solver screen output, and higher values of this parameter correspond to more screen output.

20.2.3 Debug Output Path

Scope: Trilinos Equation Solver

Debug Output Path {=|are|is} DebugOutput

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>DebugOutput</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify path where debug-logs and other debug output files will be placed.
20.2.4 Determine Sharing

Scope: Trilinos Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Determine Sharing</td>
<td>{fei</td>
<td>sierra}</td>
</tr>
</tbody>
</table>

Summary: Whether FEI should determine sharing internally or receive the info from Sierra.

Description: By default the Eqns::LinearSystem class tells the fei layer which nodes are shared and which processors share them. If this option is set to on, then the fei layer internally determines the sharing info. Requires the fei layer to perform extra communication during the initialize phase.

20.2.5 Fei Error Behavior

Scope: Trilinos Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fei Error Behavior</td>
<td>{abort</td>
<td>returncode}</td>
</tr>
</tbody>
</table>

Summary: Set FEI error behavior to abort (rather than the default which is to simply print a message and return an integer error code).

Description: This is becoming less relevant starting with FEI version 2.11, as the FEI begins to adopt exception handling and abandon the antiquated int-error-code interfaces.

20.2.6 Fei Output Level

Scope: Trilinos Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fei Output Level</td>
<td>{all</td>
<td>brief_logs</td>
</tr>
</tbody>
</table>

Summary: Control the amount of output produced by FEI.

Description: Output level controls the amount of debugging information (screen output, matrix/vector files and log files) produced by FEI. The FeiOutputLevels enum contains the valid values for this line-command. The location of files can be controlled by the ‘debug output path’ line-command.
### 20.2.7 Ilu Fill

**Scope:** Trilinos Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fill_level</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Fill-in parameter for incomplete factorizations.

### 20.2.8 Ilu Threshold

**Scope:** Trilinos Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Threshold</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Threshold parameter for incomplete factorizations.

### 20.2.9 Matrix Format

**Scope:** Trilinos Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>MatrixFormat</td>
<td>{csr</td>
<td>msr</td>
</tr>
</tbody>
</table>

**Summary**
Storage format for the matrix.

**Description**
This parameter only applies to Trilinos and Aztec.

### 20.2.10 Matrix Reduction

**Scope:** Trilinos Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>AztecReductionType</td>
<td>{fei-remove-slaves}</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Remove constraint equations from matrix.

**Description**
Dependent degrees of freedom in constraints are projected out of the equation space, yielding a linear system with smaller dimension, and retaining positive definiteness (whereas the alternative, an augmented matrix arising from the lagrange multiplier formulation, is indefinite).
20.2.11 Matrix Scaling
Scope: Trilinos Equation Solver

Matrix Scaling {=|are|is} MatrixScaling

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>MatrixScaling</td>
<td>{block-jacobi</td>
<td>jacobi</td>
</tr>
</tbody>
</table>

Summary Scaling to be applied to the matrix.

Description Matrix scaling is a pre-solve operation, typically modifying the matrix in place, whereas preconditioning is performed at each iteration during the solve and doesn’t modify the actual matrix.

20.2.12 Matrix Viewer
Scope: Trilinos Equation Solver

Matrix Viewer {=|are|is} Machine:port

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Machine:port</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Host and port-number where matvis is running.

20.2.13 Maximum Iterations
Scope: Trilinos Equation Solver

Maximum Iterations {=|are|is} Max_iters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Max_iters</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Maximum number of solution method iterations.

20.2.14 Num Levels
Scope: Trilinos Equation Solver

Num Levels {=|are|is} Num_levels

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Num_levels</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Number of levels for multi-level/multi-grid solvers.
20.2.15  Param-Bool

Scope:  Trilinos Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter_name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>Bool</td>
<td>{false</td>
<td>true}</td>
</tr>
</tbody>
</table>

Summary  String-Key/Boolean-Value pair to be passed to solver.

Description  Syntax: 'PARAM-BOOL "some-name" VALUE true | false'

20.2.16  Param-Int

Scope:  Trilinos Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter_name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>integer_value</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  String-Key/Integer-Value pair to be passed to solver.

Description  Syntax: 'PARAM-INT "some-name" VALUE 123' Primarily used for passing values to the ML package. ML has a large number of parameters which haven't been individually incorporated into our parameter-parsing system. Can also be used to pass parameters to Aztec, e.g.:

PARAM-REAL "AZ_ilut_fill" VALUE 3

20.2.17  Param-Real

Scope:  Trilinos Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter_name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>Real_value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  String-Key/Real-Value pair to be passed to solver.

Description  Syntax: 'PARAM-REAL "some-name" VALUE 1.23' Primarily used for passing values to the ML package. ML has a large number of parameters which haven't been individually incorporated into our parameter-parsing system. Can also be used to pass parameters to Aztec, e.g.:

PARAM-REAL "AZ_drop" VALUE 1.e-8
20.2.18  Param-String
Scope:  Trilinos Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter_name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>String_value</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Key/Value string-pair to be passed to solver.

Description  Syntax: 'PARAM-STRING "some-name" VALUE "some-value"' (Primarily used for passing values to the ML package. ML has a large number of parameters which haven’t been individually incorporated into our parameter-parsing system.)

20.2.19  Polynomial Order
Scope:  Trilinos Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Order</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Polynomial order of preconditioning method.

20.2.20  Preconditioner Subdomain Overlap
Scope:  Trilinos Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Overlap</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Overlap of Schwarz subdomains (eg, 0,1 or 2).

20.2.21  Preconditioning Method
Scope:  Trilinos Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>TrilinosPrecondMethods</td>
<td>{dd-bilu</td>
<td>dd-icc</td>
</tr>
</tbody>
</table>

Summary  Selection of the preconditioning method.

Description  Valid values for this are contained in the TrilinosPrecondMethods enum.
20.2.22 Preconditioning Steps
Scope: Trilinos Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Steps</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Number of Jacobi, Gauss-Seidel, or other preconditioning methods' applications per iteration.

20.2.23 Residual Norm Scaling
Scope: Trilinos Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>AztecResidualNormScaling</td>
<td>{anorm</td>
<td>none</td>
</tr>
</tbody>
</table>

Summary: Scaling method for the residual norm.
Description: Determines the residual expression used in convergence checks and printing.

20.2.24 Residual Norm Tolerance
Scope: Trilinos Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>ResidualNormTolerance</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Iterative solution method relative residual convergence tolerance.

20.2.25 Restart Iterations
Scope: Trilinos Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Restart_iters</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Number of iterations between GMRES restarts.

20.2.26 Row Ordering
Scope: Trilinos Equation Solver
20.2.27 Shared Ownership Rule

Scope: Trilinos Equation Solver

Summary: Controls the way owning processors are chosen for shared nodes in the FEI.

Description: 'low-numbered-proc' is the default.
'proc-with-local-elem' is another valid value.
'sierra_specifies' is the other valid value.

20.2.28 Solution Method

Scope: Trilinos Equation Solver

Summary: Selection of the linear-system solution algorithm.

Description: The TrilinosSolverMethods enum contains valid values for this line-command. There are several iterative methods (Krylov subspace algorithms), as well as some sparse direct methods available through the Amesos interface package. At this time (Oct’05) not all of the listed direct solvers are available. See the specific entries in the TrilinosSolverMethods enum below for more information.

20.2.29 Solve Transpose

Scope: Trilinos Equation Solver

Summary: Whether to solve for transpose of system matrix.
### 20.2.30 Select Fei

**Scope:** Trilinos Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>WhichFEI</td>
<td>new</td>
<td>old}</td>
</tr>
</tbody>
</table>

**Summary**
Selection of old vs new fei.

**Description**
This parameter will be deprecated, it is used as a transition aid during some FEI refactoring.

### 20.3 Aztec Equation Solver

**Scope:** Sierra

```
Begin Aztec Equation Solver Solver Name
   Bc Enforcement {=|are|is} BcEnforcement
   Debug Output Level {=|are|is} Level
   Debug Output Path {=|are|is} DebugOutput
   Determine Sharing {=|are|is} FeiDetermineSharing
   Fei Error Behavior {=|are|is} FeiErrorBehavior
   Fei Output Level {=|are|is} FeiOutputLevels
   Ilu Fill {=|are|is} Fill_level
   Ilu Omega {=|are|is} Value
   Ilu Threshold {=|are|is} Threshold
   Matrix Format {=|are|is} MatrixFormat
   Matrix Reduction {=|are|is} AztecReductionType
   Matrix Scaling {=|are|is} MatrixScaling
   Matrix Viewer {=|are|is} Machine:port
   Maximum Iterations {=|are|is} Max_iters
   Num Levels {=|are|is} Num_levels
   Orthog Method {=|are|is} OrthogMethod
   Param-Int Parameter_name Value integer_value
   Param-Real Parameter_name Value Real_value
   Param-String Parameter_name Value String_value
   Polynomial Order {=|are|is} Order
   Preconditioner Subdomain Overlap {=|are|is} Overlap
   Preconditioning Method {=|are|is} AztecPrecondMethods
   Preconditioning Steps {=|are|is} Steps
   Residual Norm Scaling {=|are|is} AztecResidualNormScaling
   Residual Norm Tolerance {=|are|is} ResidualNormTolerance
```
Summary A set of solver parameters for Aztec equation solver.

### 20.3.1 Bc Enforcement

**Scope:** Aztec Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>BcEnforcement</td>
<td>{exact</td>
<td>exact_no_column_mod</td>
</tr>
</tbody>
</table>

Summary Controls the way Dirichlet BCs are enforced.

Description Valid values for this line-command are contained in the BcEnforcement enum.

### 20.3.2 Debug Output Level

**Scope:** Aztec Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Level</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Output level for debugging. Generally 0 means no solver screen output, and higher values of this parameter correspond to more screen output.

### 20.3.3 Debug Output Path

**Scope:** Aztec Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>DebugOutput</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specify path where debug-logs and other debug output files will be placed.
20.3.4 Determine Sharing

Scope: Aztec Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>FeiDetermineSharing</td>
<td>fei</td>
<td>undefined</td>
<td></td>
</tr>
</tbody>
</table>

Summary: Whether FEI should determine sharing internally or receive the info from Sierra.

Description: By default the Eqns::LinearSystem class tells the fei layer which nodes are shared and which processors share them. If this option is set to on, then the fei layer internally determines the sharing info. Requires the fei layer to perform extra communication during the initialize phase.

20.3.5 Fei Error Behavior

Scope: Aztec Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>FeiErrorBehavior</td>
<td>abort</td>
<td>undefined</td>
<td></td>
</tr>
</tbody>
</table>

Summary: Set FEI error behavior to abort (rather than the default which is to simply print a message and return an integer error code).

Description: This is becoming less relevant starting with FEI version 2.11, as the FEI begins to adopt exception handling and abandon the antiquated int-error-code interfaces.

20.3.6 Fei Output Level

Scope: Aztec Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>FeiOutputLevels</td>
<td>all</td>
<td>undefined</td>
<td></td>
</tr>
</tbody>
</table>

Summary: Control the amount of output produced by FEI.

Description: Output level controls the amount of debugging information (screen output, matrix/vector files and log files) produced by FEI. The FeiOutputLevels enum contains the valid values for this line-command. The location of files can be controlled by the `debug output path` line-command.
### 20.3.7 Ilu Fill

**Scope:** Aztec Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fill_level</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  
Fill-in parameter for incomplete factorizations.

### 20.3.8 Ilu Omega

**Scope:** Aztec Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  
Omega parameter, dd-rlu uses ILU(k,w), w==omega

### 20.3.9 Ilu Threshold

**Scope:** Aztec Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Threshold</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  
Threshold parameter for incomplete factorizations.

### 20.3.10 Matrix Format

**Scope:** Aztec Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>MatrixFormat</td>
<td>{csr,msr,vbr}</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  
Storage format for the matrix.

**Description**  
This parameter only applies to Trilinos and Aztec.

### 20.3.11 Matrix Reduction

**Scope:** Aztec Equation Solver
Matrix Reduction {=|are|is} AztecReductionType

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>AztecReductionType</td>
<td>{fei-remove-slaves}</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Remove constraint equations from matrix.

Description: Dependent degrees of freedom in constraints are projected out of the equation space, yielding a linear system with smaller dimension, and retaining positive definiteness (whereas the alternative, an augmented matrix arising from the lagrange multiplier formulation, is indefinite).

20.3.12 Matrix Scaling
Scope: Aztec Equation Solver

Matrix Scaling {=|are|is} MatrixScaling

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>MatrixScaling</td>
<td>{block-jacobi</td>
<td>jacobi</td>
</tr>
</tbody>
</table>

Summary: Scaling to be applied to the matrix.

Description: Matrix scaling is a pre-solve operation, typically modifying the matrix in place, whereas preconditioning is performed at each iteration during the solve and doesn’t modify the actual matrix.

20.3.13 Matrix Viewer
Scope: Aztec Equation Solver

Matrix Viewer {=|are|is} Machine:port

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Machine:port</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Host and port-number where matvis is running.

20.3.14 Maximum Iterations
Scope: Aztec Equation Solver

Maximum Iterations {=|are|is} Max_iters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Max_iters</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Maximum number of solution method iterations.
### 20.3.15  Num Levels

**Scope:** Aztec Equation Solver

Num Levels \{=|are|is\} Num_levels

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Num_levels</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  
Number of levels for multi-level/multi-grid solvers.

### 20.3.16  Orthog Method

**Scope:** Aztec Equation Solver

Orthog Method \{=|are|is\} OrthogMethod

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>OrthogMethod</td>
<td>{classical</td>
<td>modified}</td>
</tr>
</tbody>
</table>

**Summary**  
User can choose orthogonalization method used by Aztec GMRES.

**Description**  
Valid values for this line-command are contained in the OrthogMethod enum.

### 20.3.17  Param-Int

**Scope:** Aztec Equation Solver

Param-Int Parameter_name Value integer_value

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter_name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>integer_value</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  
String-Key/Integer-Value pair to be passed to solver.

**Description**  
Syntax: 'PARAM-INT "some-name" VALUE123' Primarily used for passing values to the ML package. ML has a large number of parameters which haven’t been individually incorporated into our parameter-parsing system. Can also be used to pass parameters to Aztec, e.g.: PARAM-REAL "AZ_ilut_fill" VALUE 3

### 20.3.18  Param-Real

**Scope:** Aztec Equation Solver

Param-Real Parameter_name Value Real_value

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter_name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>Real_value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Summary  String-Key/Real-Value pair to be passed to solver.

Description  Syntax: 'PARAM-REAL "some-name" VALUE 1.23' Primarily used for passing values to the ML package. ML has a large number of parameters which haven’t been individually incorporated into our parameter-parsing system. Can also be used to pass parameters to Aztec, e.g.:

PARAM-REAL "AZ_drop" VALUE 1.e-8

20.3.19  Param-String
Scope:  Aztec Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter_name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>String_value</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Key/Value string-pair to be passed to solver.

Description  Syntax: 'PARAM-STRING "some-name" VALUE "some-value"' (Primarily used for passing values to the ML package. ML has a large number of parameters which haven’t been individually incorporated into our parameter-parsing system.)

20.3.20  Polynomial Order
Scope:  Aztec Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Order</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Polynomial order of preconditioning method.

20.3.21  Preconditioner Subdomain Overlap
Scope:  Aztec Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Overlap</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Ovrlap of Schwarz subdomains (eg, 0,1 or 2).

20.3.22  Preconditioning Method
Scope:  Aztec Equation Solver
Preconditioning Method {=}|are|is} AztecPrecondMethods

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>AztecPrecondMethods</td>
<td>{dd-bilu</td>
<td>dd-icc</td>
</tr>
</tbody>
</table>

Summary  Selection of the equation preconditioning method.

Description  Valid values for this are contained in the AztecPrecondMethods enum.

20.3.23  Preconditioning Steps
Scope:  Aztec Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Steps</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Number of Jacobi, Gauss-Seidel, or other preconditioning methods’ applications per iteration.

20.3.24  Residual Norm Scaling
Scope:  Aztec Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>AztecResidualNormScaling</td>
<td>{anorm</td>
<td>none</td>
</tr>
</tbody>
</table>

Summary  Scaling method for the residual norm.

Description  Determines the residual expression used in convergence checks and printing.

20.3.25  Residual Norm Tolerance
Scope:  Aztec Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>ResidualNormTolerance</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Iterative solution method relative residual convergence tolerance.
20.3.26 Restart Iterations
Scope: Aztec Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Restart Iterations</td>
<td>Restart_iters</td>
<td>integer</td>
</tr>
</tbody>
</table>

Summary: Number of iterations between GMRES restarts.

20.3.27 Shared Ownership Rule
Scope: Aztec Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shared Ownership Rule</td>
<td>SharedOwnershipRule</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Controls the way owning processors are chosen for shared nodes in the FEI.

Description:
'low-numbered-proc' is the default.
'proc-with-local-elem' is another valid value.
'sierra_specifies' is the other valid value.

20.3.28 Solution Method
Scope: Aztec Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Solution Method</td>
<td>AztecSolverMethods</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Selection of the linear-system solution algorithm.

Description: The AztecSolverMethods enum contains valid values for this line-command. They are mostly iterative methods (Krylov subspace algorithms), except for 'lu' which is a serial direct solver provided by the 'Y12M' library.

20.3.29 Select Fei
Scope: Aztec Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Select Fei</td>
<td>WhichFEI</td>
<td>undefined</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>WhichFEI</td>
<td>{new,old}</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Summary  
Selection of old vs new fei.

Description  
This parameter will be deprecated, it is used as a transition aid during some FEI refactoring.

20.4  Gdsw Equation Solver
Scope:  Sierra

Begin Gdsw Equation Solver Solver Name
  Debug Output Level {=|are|is} Level
  Debug Output Path {=|are|is} DebugOutput
  Fei Error Behavior {=|are|is} FeiErrorBehavior
  Fei Output Level {=|are|is} FeiOutputLevels
  Maximum Iterations {=|are|is} Max_iters
  Param-Int Parameter_name Value integer_value
  Param-Real Parameter_name Value Real_value
  Param-String Parameter_name Value String_value
  Residual Norm Tolerance {=|are|is} ResidualNormTolerance
End

Summary  
A set of solver parameters for GDSW equation solver.

20.4.1  Debug Output Level
Scope:  Gdsw Equation Solver

Debug Output Level {=|are|is} Level

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Level</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  
Output level for debugging. Generally 0 means no solver screen output, and higher values of this parameter correspond to more screen output.

20.4.2  Debug Output Path
Scope:  Gdsw Equation Solver

Debug Output Path {=|are|is} DebugOutput

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>DebugOutput</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  
Specify path where debug-logs and other debug output files will be placed.
20.4.3 Fei Error Behavior
Scope: Gdsw Equation Solver

Fei Error Behavior \{=|are|is\} \textit{FeiErrorBehavior}

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{FeiErrorBehavior}</td>
<td>{abort</td>
<td>returncode}</td>
</tr>
</tbody>
</table>

Summary  Set FEI error behavior to abort (rather than the default which is to simply print a message and return an integer error code).

Description  This is becoming less relevant starting with FEI version 2.11, as the FEI begins to adopt exception handling and abandon the antiquated int-error-code interfaces.

20.4.4 Fei Output Level
Scope: Gdsw Equation Solver

Fei Output Level \{=|are|is\} \textit{FeiOutputLevels}

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{FeiOutputLevels}</td>
<td>{all</td>
<td>brief_logs</td>
</tr>
</tbody>
</table>

Summary  Control the amount of output produced by FEI.

Description  Output level controls the amount of debugging information (screen output, matrix/vector files and log files) produced by FEI. The FeiOutputLevels enum contains the valid values for this line-command. The location of files can be controlled by the ‘debug output path’ line-command.

20.4.5 Maximum Iterations
Scope: Gdsw Equation Solver

Maximum Iterations \{=|are|is\} \textit{Max_iters}

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{Max_iters}</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Maximum number of solution method iterations.

20.4.6 Param-Int
Scope: Gdsw Equation Solver

Param-Int \textit{Parameter_name} \textit{Value} integer \textit{integer_value}

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{Parameter_name}</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>\textit{integer_value}</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Summary  String-Key/Integer-Value pair to be passed to solver.

Description  Syntax: 'PARAM-INT "some-name" VALUE 123' Primarily used for passing values to the ML package. ML has a large number of parameters which haven't been individually incorporated into our parameter-parsing system. Can also be used to pass parameters to Aztec, e.g.: PARAM-REAL "AZ_ilut_fill" VALUE 3

20.4.7  Param-Real
Scope:  Gdsw Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter_name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>Real_value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  String-Key/Real-Value pair to be passed to solver.

Description  Syntax: 'PARAM-REAL "some-name" VALUE 1.23' Primarily used for passing values to the ML package. ML has a large number of parameters which haven't been individually incorporated into our parameter-parsing system. Can also be used to pass parameters to Aztec, e.g.: PARAM-REAL "AZ_drop" VALUE 1.e-8

20.4.8  Param-String
Scope:  Gdsw Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter_name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>String_value</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Key/Value string-pair to be passed to solver.

Description  Syntax: 'PARAM-STRING "some-name" VALUE "some-value"' (Primarily used for passing values to the ML package. ML has a large number of parameters which haven’t been individually incorporated into our parameter-parsing system.)

20.4.9  Residual Norm Tolerance
Scope:  Gdsw Equation Solver

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>ResidualNormTolerance</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Iterative solution method relative residual convergence tolerance.
Chapter 21

Postprocessing Operations

21.1 Overview

Aria supports postprocessing of solution information partially through Aria itself and alternatively through the Encore library [34].

Postprocessing through Aria itself is invoked using a single line like:

Postprocess density on block_1

This creates a field "pp->density" that is available for output from the Results Output Command Block 25.1. Any expression created during the calculation may be postprocessed, of particular interest are properties defined in the material block input 4 and source terms present for any equation. In addition single components of any tensor quantity can be postprocessed by appending the suffix for the desired component (e.g. "_XX", "_XY", etc.) to the end of the expression name used in the "Postprocess" line command. Similarly, the Frobenius norm of a tensor may be postprocessed by using the "_NORM" suffix.

In addition to the pp postprocessing command another means of postprocessing called solution options is also available. Whereas pp can be used for material parameters, the use of solution options usually implies that the postprocessing variable is the result of an algorithmic computation. Within the input file postprocessing command lines will appear at the Region scope as illustrated below.

Begin Procedure My_Aria_Procedure

Begin Aria Region My_Region

Post Processing related commands

Begin Solution Options

post process commands

End Solution Options

End
Aria postprocessing with Encore is usually accomplished using Encore specific commands [34]. Input commands for both pp and solution options postprocessing are included in what follows.

## 21.1.1 Postprocess

**Scope:**

```
Postprocess Expression [ {of|species} SpeciesName | {in|material_phase} MaterialPhaseName ]{@|at|for|in|on|over} MeshPartIdentifier [ Using ElementType | Averaging Method {=|are|is} AveragingMethod ]
```

### Parameter | Value | Default
---|---|---
Expression | string | undefined
SpeciesName | string | undefined
MaterialPhaseName | string | undefined
MeshPartIdentifier | string | undefined
ElementType | string | undefined
AveragingMethod | string | undefined

**Summary**

Postprocess Expression on a mesh part (block, surface, etc.). By default, the quantity is computed on the grid element nodes. If the optional arguments "using ElementType" are provided, the postprocessed field will be computed on that element. If the ElementType is P0 then the quantity is computed as a constant element value. By default the nodal field value is calculated using a weighted average of the value in each element at the node location where the element contributions are weighted by $\int w_{\text{node}} d\Omega$, This is the SUPPORT_WEIGHTED averaging method. Two alternative averaging methods are available. SUPPORT_AVERAGED integrates $\int \phi w_{\text{node}} d\Omega$ (where $\phi$ is the expression being postprocessed) over all supporting elements and normalizes by $\int w_{\text{node}} d\Omega$ over all supporting elements. SIMPLE just takes the mean of the element values at the node location.

## 21.2 Solution Options

**Scope:** Aria Region

```
Begin Solution Options OptionsName

Post Process PostProcessType VariableName On IoPartList... As OutputName [ Using WeightList... Weight ]

Post Process Flux fluxName On ioPartList... As outputName [ Restricted To restrictPartList... ]

Post Process Mass

Post Process Nodal PostProcessType VariableName On IoPartList... As OutputName [ Using WeightList... Weight ]

Post Process Volume

Begin Cvfem Algorithm Specification PStabName
End
```

676
Summary Specify information regarding the governing equations to be solved.

21.2.1 Post Process
Scope: Solution Options

Post Process PostProcessType VariableName On IoPartList... As OutputName [ Using WeightList... Weight ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>PostProcessType</td>
<td>integral</td>
<td>normalized}</td>
</tr>
<tr>
<td>VariableName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>IoPartList</td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td>OutputName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Post process an Aria variable and save it to a global variable.

Description Post process Aria variable VariableName and save it to a global variable OutputName. The global OutputName will be written to the log file and is also available for output to an ExodusII database. During post processing the variable a product is formed from VariableName and the evaluation of WeightList items. The keyword WEIGHT terminates the Weightlist items.

21.2.2 Post Process Flux
Scope: Solution Options

Post Process Flux fluxName On ioPartList... As outputName [ Restricted To restrictPartList... ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>fluxName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>ioPartList</td>
<td>string...</td>
<td>undefined</td>
</tr>
<tr>
<td>outputName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Post process an Aria flux and save the integrated results to a global Aria Field.

Description Post process Aria flux FluxName and save the integrated flux to a global Field variable OutputName. The resultant variable OutputName will be available for output to an ExodusII database. Examples of FluxName are heat_conduction and current_density.
21.2.3 Post Process Mass
Scope: Solution Options

Summary  Post process both the total mass for all blocks and per-block mass. By default these values will be output to log file.

Additionally these values are available as global variables which can be output to the ExodusII database. Naming convention of these values are TOTAL_MASS for the aggregate mass and MASS_mesh_entity_name (e.g. MASS_BLOCK_1) for individual blocks.

21.2.4 Post Process Nodal
Scope: Solution Options

Post Process Nodal PostProcessType VariableName On IoPartList... As OutputName [ Using WeightList... Weight ]

Parameter | Value | Default
---|---|---
PostProcessType | {integral|normalized} | undefined
VariableName | string | undefined
IoPartList | string... | undefined
OutputName | string | undefined

Summary  Post process an Aria nodal variable and save the results to another Aria Field.

Description  Post process Aria variable VariableName and save it to another nodal Field variable OutputName. The resultant variable OutputName will be available for output to an ExodusII database. PostProcessType is NORMALIZED so that during post processing the variable a product is formed from VariableName and the evaluation of WeightList items. The keyword WEIGHT terminates the Weightlist items.

21.2.5 Post Process Volume
Scope: Solution Options

Summary  Post process both the total volume and and per-block volume. By default these values will be output to log file.

Additionally these values are available as global variables which can be output to the ExodusII database. Naming convention of these values are TOTAL_VOLUME for the aggregate volume and VOLUME_mesh_entity_name (e.g. VOLUME_BLOCK_1) for individual blocks.

21.3 Residual Based Integrated Surface Flux Calculations

Oftentimes one needs to require the evaluation of surface flux over some portion of the problem domain. Examples of such evaluations include the net thermal energy through a surface or the net force on a body. While the evaluation can be carried out in a variety of ways one method of carrying out an accurate evaluation
of these quantities is to employ a residual approach. The general approach is described here for problems involving scalar variables but the technique applies equally as well to vector variables.

Governing partial differential equations for scalar potential \( u \) are often of the form

\[
C \frac{Du}{Dt} = \nabla \cdot q + Q \tag{21.1}
\]

where \( C \) is storage coefficient, \( q \) is the flux of \( u \), \( Q \) is a volumetric source and the substantial derivative \( \frac{D}{Dt} \) accounts for material motion. The flux of \( u \) is oftentimes a simple function of the potential gradient

\[
q = -k \nabla u \tag{21.2}
\]

where \( k \) is a diffusion tensor. In many cases \( k \) is isotropic so it is treated as a constant coefficient.

To obtain solutions of scalar PDE (21.1) using finite elements, one forms a weighted residual over a spatial discretization of elements of the problem domain \( \Omega \). Integration by parts, then yields a general form of the nodal residual

\[
R^i = \int_{\Omega} \left[ \left( C \frac{Du}{Dt} - Q \right) \phi^i - q \cdot \nabla \phi^i \right] d\Omega + \int_{\Gamma_n} n \cdot q \phi^i d\Gamma \tag{21.3}
\]

where \( i \) denotes a node of the meshed discretization, \( \phi^i \) is the finite element weight function for node \( i \). The collection of nodal residual equations over \( \Omega \) form a linear system of equations for which we seek to minimize the nodal residuals, \( R^i = 0 \) over the solution domain.

Dirichlet boundary conditions on (21.3) for specific nodes can be eliminated from the system and the unknown temperature is assigned directly. For flux boundary conditions, the term \( n \cdot q \) is replaced with a model of the surface normal flux. Surface contact boundary conditions can also be accounted for through a special implementation of this surface integral as outlined in the Aria user manual.

In some problems there is need of computing the surface flux \( f_n \) over a subset of the domain boundary \( \Gamma \) and one might consider evaluating

\[
f_n = \int_{\Gamma_n} n \cdot q \ d\Gamma_n \ . \tag{21.4}
\]

This approach requires the evaluation of finite element gradients on element boundaries as per (21.2) but is known to be problematic since gradients of the finite element solutions in terms of the basis functions \( \phi^i \) are of lower order than the finite element solution. An alternative is to evaluate the flux over \( \Gamma_n \) using the volumetric term of (21.3)

\[
f^i_n = -\int_{\Omega_n} \left[ \left( C \frac{Du}{Dt} - Q \right) \phi^i - q \cdot \nabla \phi^i \right] d\Omega_n \tag{21.5}
\]

which must be computed while forming the linear system of equations and is known to be second order accurate. Note that \( f^i \) is the integrated surface flux contribution at a node, not a flux per unit area. The area over which \( f^i_n \) is computed is of course mesh dependent and is given by the finite element support for node \( i \).

In order to obtain the nodal flux (21.5), we proceed with our typical FEM assembly procedure in the evaluation of (21.3) where the volume and surface contributions are handled in separate portions of the code. Using the `Save Residuals = Before_BCs` input command line in Aria, we request that intermediate values of the residuals be stored after the volumetric terms are calculated but before the surface contributions are computed. These integrated nodal flux values are stored in a field named `residual->dof_name`, where `dof_name` corresponds to the solution variable \( u \) for a given equation.

The total flux on a portion of the boundary \( \Gamma_n \) Computation of \( t \) is illustrated for the energy equation with solution variable \textit{temperature}. is obtained by summing the residual values for all nodes along the surface

\[
f_n = -\sum_{i \in \Gamma_n} \int_{\Omega_n} \left[ \left( C \frac{Du}{Dt} - Q \right) \phi^i - q \cdot \nabla \phi^i \right] d\Omega_n \ . \tag{21.6}
\]
Computation of the total flux over $\Gamma_n$ is illustrated for the energy equation with solution variable temperature. To perform this calculation using Aria one first needs to enable the saving of residuals. In the Aria Region block we add the command line

```
Save Residuals = Before_BCs
```

This instructs Aria to store the result of (21.3) in a field named residual->temperature. Next, to perform the summation in (21.6) on a portion of the surface, say sideset_12 one can add an Encore summation postprocessor block at the domain scope:

```
Begin Summation Postprocessor flux_12
Use Function residual->temperature
Surfaces surface_12
End
```

To execute this postprocessor, one adds an additional command line to the Aria Region scope:

```
Evaluate Postprocessor flux_12
```

This will create a global variable named flux_12 whose result will be written to the Aria log file for each successful solution step. Finally, one can output the result flux_12 to the Exodus results output database file by adding the following line to the appropriate output block:

```
global variables = flux_12
```

Similarly the global variable can be written to the heartbeat output file by adding the line

```
variable = global flux_12
```

to the Heartbeat file command block.
Chapter 22

Enclosure Radiation Reference

22.1 Enclosure Radiation

When energy radiates from one portion of a surface to another, and the intermediate medium is transparent (i.e., it does not absorb any energy), then enclosure radiation may be used to model the heat flux on the surface. Using the net radiation method [35], the normal flux at a particular location on the surface may be written as the difference between the emitted radiative heat flux leaving the surface, and the absorbed incident radiative flux due to the rest of the enclosure, namely

$$q_n = \sigma \epsilon T^4 - \alpha G,$$  \hfill (22.1)

where $T$ is the temperature, $\alpha$ denotes the absorptivity of the surface, and $G$ represents the surface irradiation. Under the additional assumption that the emissivity, absorptivity, and reflectivity are independent of direction and wavelength, we may write

$$\epsilon = \alpha = 1 - \rho,$$  \hfill (22.2)

where we have used the conventional symbol $\rho$ for reflectivity. In this section, $\rho$ always refers to reflectivity and not density.

Without loss of generality, we can regard the enclosure, $\Gamma_E$, as a union of $E$ surfaces,

$$\Gamma_E = \Gamma_1 \cup \Gamma_2, \ldots \Gamma_{E-1} \cup \Gamma_E$$

This situation is illustrated in Figure 22.1, where the the radiosity for surface $i$ in the enclosure is defined to be

$$J_i = \sigma \epsilon_i T_i^4 + \rho_i G_i,$$  \hfill (22.3)

where $u_i$ is the spatially constant temperature on $\Gamma_i$. The surface irradiation for surface $i$ is determined by the radiosity of all the other surfaces in the enclosure through the relation

$$G_i = \sum_{j=1}^{E} F_{ij} J_j,$$  \hfill (22.4)

where $F_{ij}$ denotes the geometric viewfactor of surface $i$ with respect to surface $j$. The viewfactor may be considered the fraction of energy that leaves surface $i$ and arrives at surface $j$.

Upon substitution of equations (22.4) and (22.2) into (22.3), the radiosity may be written as

$$J_i - (1 - \epsilon_i) \sum_{j=1}^{N} F_{ij} J_j = \sigma \epsilon_i T_i^4,$$  \hfill (22.5)

Finally, the first $J_i$ term in (22.5) may be moved inside the summation to yield

$$\sum_{j=1}^{N} [\delta_{ij} - (1 - \epsilon_i)F_{ij}] J_j = \sigma \epsilon_i T_i^4,$$  \hfill (22.6)
where
\[
\delta_{ij} = \begin{cases} 
1 & \text{if } i = j \\
0 & \text{if } i \neq j 
\end{cases} 
\] (22.7)

(22.6) is a nonlinear system of equations for the radiosities that must be solved simultaneously with (at least) the energy transport equations (see 5.3). Finally, we may rewrite (22.1) to express the normal flux boundary condition on surface \(i\) as
\[
q_n(T) = \sigma \epsilon T^4 - \epsilon G_i, 
\] (22.8)
where \(G_i\) is given by (22.4).

## 22.2 Enclosure Radiation Surface Flux

The enclosure radiation boundary condition given by (22.1) enters our weak statement in a manner similar to that of the surface radiation flux described in 9.2.16. The enclosure radiation flux was written as (22.1), which we repeat here for convenience:
\[
q_n(T) = \sigma \epsilon T^4 - \epsilon G_i, 
\] (22.9)

This flux and the surface radiation flux are both nonlinear, and vary as the fourth power of the temperature, but there are important differences. First, instead of a known reference temperature \(T_r\), the surface irradiation, \(G\), appears in (22.9), and must be calculated as part of the solution process. To begin, let \(\Gamma^e\) be an arbitrary finite element on the enclosure. We call \(\Gamma^e\) a *facet*, and let \(\Gamma^e\) consist of a total of \(E\) facets. Furthermore, the flux (22.9) is constant on a facet; to emphasize this, let us rewrite it as
\[
q^e_n(T) = \sigma \epsilon [T^e]^4 - \epsilon G_e(T^e), 
\] (22.10)
where \(T^e\) is some average facet temperature, and \(G_e(T^e)\) is the irradiation on facet \(e\), which is determined by the combined effects of all the other facets in the enclosure. Recall that this was expressed in Section 22.1.
as
\[ G_e(T^e) = \sum_{j=1}^{E} F_{ef} J_f(T^e). \] (22.11)

Details of our nonlinear solution strategy are not included here but it is important to note here that the unknown radiosities, \( J_f \), depend upon the solution of the temperature. We use a decoupled approach, and calculate the viewfactors, \( F_{ef} \), and radiosities, \( J_f \), via the Chaparral library \[36\]. Recall that the radiosities are obtained by solving the system of equations
\[ \sum_{j=1}^{N} \left[ \delta_{ef} - (1 - \epsilon_e) F_{ef} \right] J_f = \sigma \epsilon_e [T^e]^4 \] (22.12)

In (22.12), it is important to realize that the \( J_f \) and \( T^e \) are constant values associated with a given facet. But in our finite element approximation, the temperatures are associated with the nodes. Therefore, the facet temperature should be considered a projection, or averaging of the temperature found at the nodes of a given facet. Hence, we define the facet temperature as
\[ T^e = \frac{\sum_{i=1}^{N} T_i \int_{\Gamma_i^e} \psi_e^c d\Gamma}{J_{\psi_e^c} d\Gamma} \] (22.13)

It is now convenient to define the projection vector \( P^e \), where row \( i \) is defined by
\[ P_i^e = \frac{1}{A^e} \int_{\Gamma_i^e} \psi_e^c d\Gamma, \] (22.14)
and \( A^e \) is the area of facet \( e \). Thus, (22.13) may be written as
\[ T^e = \sum_{i=1}^{N} P_i^e T_i \] (22.15)

### 22.3 Solution Strategy

When using Newton’s method one can employ the surface radiation flux 22.9 directly to obtain Jacobian and residual contributions to the linear system of equations. Solution convergence of enclosure radiation problems by successive substitution is improved by employing a linearization of the surface radiation flux as described in what follows.

Now that we have appropriately defined the facet temperature, we may linearize the flux given in (22.10). Expanding \( q_n(T) \) in a Taylor series about a known state \( T^e_* \), we obtain
\[ q_n(T_* + \delta T) = \sigma \epsilon [T_*^e]\ 4 - \epsilon G_e(T_*^e) + 4 \sigma \epsilon [T_*^e]^3 \delta T \] (22.16)
where \( \delta T = T^e - T_*^e \), and we have chosen to evaluate the irradiation at the known state, \( T_*^e \). If we collect terms involving the known \( T_*^e \) and unknown \( T^e \), (22.16) may be rearranged and written as
\[ q_n(T_* + \delta T) = 4 \sigma \epsilon [T_*^e]^3 T^e - 3 \sigma \epsilon [T_*^e]^4 - \epsilon G_e(T_*^e) \] (22.17)

In order to apply this flux in our weak statement, we must integrate it against the test function, \( \psi_e^c \). Therefore, we have
\[ \int_{\Gamma_i^e} \psi_i^e q_n(T) d\Gamma = \int_{\Gamma_i^e} \psi_i^e \left\{ 4 \sigma \epsilon [T_*^e]^3 T^e - 3 \sigma \epsilon [T_*^e]^4 - \epsilon G_e(T_*^e) \right\} d\Gamma \] (22.18)
Exploiting the fact that all quantities on a given facet are constant, except for the shape functions, (22.18) becomes

\[
\int_{\Gamma} \psi_i^e q_n(T) \, d\Gamma = \int_{\Gamma} \psi_i^e \, d\Gamma \left\{ 4\sigma \varepsilon [T_\star^e]^3 T^e \right\} \\
- \int_{\Gamma} \psi_i^e \, d\Gamma \left\{ 3\sigma \varepsilon [T_\star^e]^4 + \varepsilon G_e(T_\star^e) \right\} 
\]

(22.19)

If we introduce the definitions (22.15) and (22.14) into (22.19), we see that the linearized form of the element matrix contribution is

\[
K_{ij}^e = 4\sigma \varepsilon [T_\star^e]^3 A^e P_i^e P_j^e 
\]

and the contribution to the source vector is

\[
F_i^e = A^e \varepsilon \left\{ 3\sigma [T_\star^e]^4 + G_e(T_\star^e) \right\} P_i^e 
\]

(22.20)

(22.21)

We remark that, when calculating terms in Equations (22.20) and (22.21), experience has shown that it is important to project the exponential power of the temperature, as opposed to projecting the temperature, then raising it to some exponential power. In other words, spurious oscillations are less likely to occur if

\[
[T^e]^4 = \sum_{i=1}^{N} P_i^e T_i^4 
\]

is used instead of

\[
[T^e]^4 = \left[ \sum_{i=1}^{N} P_i^e T_i \right]^4 
\]

22.4 Banded Wavelength Enclosure Radiation

The enclosure radiation model previously presented considers a net response over all wavelengths assuming that all surfaces behave as grey bodies. Here the surface response with regard to different wavelengths is implicitly included in the model by using temperature dependent emissivities and allowing the surfaces to emit as grey bodies. In this case the surface flux previously mentioned 22.1 can be alternatively expressed as

\[
q_n = \int_{0}^{\infty} q(\lambda) \, d\lambda . 
\]

(22.22)

In many applications engineering materials respond differently to different portions of the thermal energy spectrum. Thus modeling the thermal radiation response of these materials using the entire blackbody spectrum results in poor characterization of the enclosure response. For this purpose more specialized approaches for radiation modeling of wavelength dependent surfaces have been developed using continuous representations of emissivity as a function of wavelength and temperature while integrating over the wavelength spectrum. For numerical modeling the simplest approach involves discretization of the wavelength spectrum into a few representative bands \(N_b\) and integrating over each of the \(k\) bands. Using this approach the methods previously described follow directly except that the emissivity and emissive power now have independent representations in a wavelength band, each of which contributes some surface flux \(\Delta q_k\) thus

\[
q_n = \sum_{k=1}^{N_b} \Delta q_k \lambda . 
\]

(22.23)
Recalling that flux can also be expressed in terms of radiosity then similarly the radiosities are obtained by solving a system of equations

$$\sum_{k=1}^{N_n} \sum_{f=1}^{N_f} \left[ \delta(\epsilon_{ek}) - (1 - \epsilon_{ek}) F_{ef} \right] \Delta J_{fk} \Delta \lambda_k = \sum_{k=1}^{N_n} F \Delta \lambda_k T^4 \sigma \epsilon_{ek} [T^4] \Delta \lambda_k$$  \hspace{2cm} (22.24)

where $\Delta J_{fk}$ is the incremental radiosity and $F \Delta \lambda_k$ is the blackbody fraction for band $k$. Finally we note that given a fixed facet temperature field this solution can be carried out independently for each wavelength band and the radiosities $\Delta J_{fk}$ can be accumulated to obtain the net facet radiosity $J_f$ and net flux $q_n$. Thus the banded wavelength model very much resembles the more simplified enclosure radiation model 22.1. From a simulation point of view, the major difference being that the modeler must supply information describing the band discretization in addition to emissivity models for each band.

Construction of the banded wavelength model begins by first prescribing the emissivity variation for each surface of the enclosure as a function of wavelength $\lambda$. The overall band discretization is then obtained by collective consideration of the wavelengths at which the emissivity changes on any of the given surfaces. Thus the number of bands for the enclosure will likely be more than those for any single surface and is demonstrated in Figure 22.2 below. It is important to note that for each modeled band an emissivity model must be supplied for each surface, even when its emissivity is not changing.

![Emissivity Bands For Two-Surface Problem](image)

**Figure 22.2.** Emissivity Bands For Two-Surface Problem

### 22.5 Enclosure Radiation Modeling

Enclosure radiation modeling is normally performed in several steps,
• Compute the viewfactors or read in pre-computed viewfactors.
• Radiosity solve.
• Perform the coupled finite element solve.

The last step is handled implicitly once boundary conditions on the finite element model have been defined. To facilitate the first three steps the analyst must first define the enclosure. For complex enclosures smoothing (conditioning) of the viewfactors is often required to ensure the quality of the viewfactors before proceeding to the radiosity solve and a facility for performing this operation is also provided. As part of this enclosure definition details concerning both computation of viewfactors, and the radiosity solve must be included. Full details of enclosure definition, viewfactor calculation, viewfactor smoothing and radiosity solve are specified in separate input command blocks. Additional input command blocks must also be supplied in order to employ the banded-wavelength enclosure model. Individual directives appearing within the aforementioned command blocks are described in the sections which follow.

In all cases the enclosure is viewed as a closed surface. As illustrated in Figure 22.3 the discretization may need to be modified in order to properly define the closed surface.

![Figure 22.3. Discretization and Associated Closed Enclosure Surface.](image)

For problems in which the enclosure surface is not entirely meshed, it can be implicitly closed by defining a partial enclosure and its associated properties within the enclosure definition command block. The partial enclosure is demonstrated in Figure 22.4. Note that the partial enclosure approach is a model simplification and is no substitute for a fully-meshed enclosure. Best results with partial enclosures are obtained when most of the enclosure model facets have the same view of the partial enclosure area and temperature.

As part of the partial enclosure description one must supply a partial enclosure area but for many problems it may be difficult to initially compute or even estimate the partial enclosure area. For these cases provision is made within Aria to internally compute the minimum area that should be used. To obtain the minimum area one temporarily assigns a small number for the partial enclosure, starts the problem and terminates it after one step. The minimum partial enclosure area can then be obtained from the .output file. Further discussion of the partial enclosure is given in a section which follows 22.5.1.

![Figure 22.4. Discretization and Associated Partial Enclosure Surface.](image)

It is worth mentioning that the viewfactor between two facets is roughly proportional to \( \frac{1}{r^2} \) where \( r \) is the distance between facet centroids. Thus it follows directly that facets which are nearly parallel may adversely affect the viewfactor calculation. Similar statements apply to poorly equivalenced surface intersections leading to slivered surfaces becoming part of an enclosure. In the worst of circumstances the
viewfactor will simply fail when the distance between facets is too small. At the user’s request some attempt can be made to remedy these situations by preprocessing of the surface facets to remove these defects, via an algorithm denoted as DASH. This algorithm can be applied universally to all the enclosures or to each enclosure independently. The use of DASH for the construction of enclosures is functionally equivalent to the standard prescription of surfaces except in the case where partial enclosures are present. DASH presents a powerful mechanism for automatically generating enclosures but the user has to ensure that the problem being addressed is appropriate such as cases where the entire set of element blocks is used to generate the enclosures but the user is only interested in a subset.

One measure of how accurately the viewfactors are being computed is the rowsum value. For each surface facet in a fully-closed enclosure the rowsum value should approach one, thus the total rowsum value is always the number of facets. In cases where the target rowsum is not equal to one the surface may not be fully-closed or the view factor calculation may be inaccurate for some other reason. In either event the analyst should consider investigate possible causes of the discrepancy. The viewfactor calculation is often followed by a smoothing process which may offset some of the errors indicated by the rowsum values. Note that while use of the Dash algorithm can be useful to overcome inaccuracies, there is no substitute for repair of any mesh defects.

Spatial discretization of the enclosure consists of surfaces as illustrated in Figure 22.1 where the surfaces respect an inward facing normal orientation. For most cases the enclosure is devoid of interior mesh discretization. In the event that the enclosure interior is meshed, the interior mesh can be ignored by using the OMIT_VOLUME command line in the FINITE_ELEMENT_MODEL command block.

The objective in solving the enclosure radiation problem is to obtain flux values at the enclosure cavity surfaces for later application to the finite element model. In a special class of closed enclosure problems one may wish to superpose flux contributions from different enclosure radiation problems overlying the same surfaces as depicted in Figure 22.5. This situation often arises in cases where the enclosure thermal response is wavelength dependent. For any given enclosure, unique descriptions of the radiation enclosure surfaces are provided to the Chaparral library. When enclosure radiation surfaces overlap special provision must be made in order to uniquely define these surfaces for the Chaparral library. Here the distinction between surfaces is made by denoting an enclosure overlying another enclosure as an OVERLAPPING ENCLOSURE. In the example, two consistent definitions of OVERLAPPING ENCLOSURE are possible. Enclosure 2 can be denoted as the overlapping enclosure since it overlaps enclosures 1 and 3. Likewise enclosures 1 and 3 could be denoted as overlapping enclosures since they overlap Enclosure 2.

![Figure 22.5. Superposed Radiation Enclosures.](image-url)
In certain simulations one might require an interior mesh to represent other coupled physics within the cavity area of the enclosure. In this case one must use the MESHED_ENCLOSURE command within the enclosure definition command block to exclude the meshed block (and its surface) from the enclosure radiation problem. For simulations in which enclosure radiation approaches the optically-thick limit one might consider utilizing a meshed enclosure in conjunction with the OPTICALLY_THICK thermal conductivity model 4.52.6 in lieu of the enclosure radiation problem.

Once a simulation has begun the enclosure radiation problem is solved for each enclosure in the order they appear in the input file. First the viewfactors are obtained, then followed by a radiosity solve for that enclosure. If any errors are encountered while resolving viewfactors or during a radiosity solve, the solution is terminated with appropriate diagnostics. After all the individual enclosure radiation problems have been solved, those results are then coupled with the finite element solution. In most simulations involving enclosure radiation computation of the viewfactors represents a large portion of the overall solution time. If the same geometric model will be used for different analyses saving the the viewfactors to file and reusing them is highly recommended. Code facilities for writing the viewfactor files and for reading them in are provided specifically for this purpose.

After performing a simulation the analyst will often interpret the results by post-processing of the output data. To assist in this task the analyst is able to request output of the enclosure surface flux, \( q_n \) and irradiation \( G \) of equation 22.1 as well as the radiosity \( J \) 22.3. If these variables are of interest their output can be requested by including the names RAD_FLUX, IRRADIANCE, or RADIOSITY in the results output block described in a chapter 25. When using the banded-wavelength model the flux, irradiation and radiosity can be output on each band by prefacing the names with ENCLOSURE_EID, where \( EID \) is the enclosure index (the order index corresponding to where the enclosure definition appears in the input command file). In this case the values corresponding to each band will be output. All enclosure surface output is written as face variables in 3D and edge variables in 2D. Because solution variables for the partial enclosure, RAD_FLUX, IRRADIANCE, or RADIOSITY can have some relevance, their output can also be requested. However, unlike the variables corresponding to part of the surface mesh, partial enclosure variable output is requested from the enclosure definition command block and is written to a global variable.

### 22.5.1 Partial Enclosures

The concept of a partial enclosure is best demonstrated by use of a simplified view of radiative transfer system between surfaces 1 and 2 surrounded by free space as shown in Figure 22.6.

![Surface Normals](image)

**Figure 22.6.** Two-Body Radiative Transfer Model

If radiative transfer occurs between the two bodies and the view factor between the two bodies is readily accessible a simplified network representation of the interaction, Figure 22.7 may suffice but here the interest lies in analyzing the system using enclosure radiation.

Recall that the enclosure view factors \( F_{ij} \) are computed as

\[
F_{ij} = \frac{1}{A_i} \int_{A_i} \int_{A_j} \frac{\cos \theta_i \cos \theta_j}{\pi r^2} \delta_{ij} dA_i dA_j.
\]
Determination of the view factors is a compute intensive endeavor and within most code implementations extraneous calculations are eliminated based upon the geometry. One example of this would be excluding this calculation for surfaces which are not visible to each other. Moreover, from a geometric view of enclosure surfaces, Figure 22.8, one can conclude that legitimate interactions between surfaces are those for which \( n_i \) and \( n_j \) are opposed. Thus an important feature of the enclosure model is the notion of inward facing normals. This convention effectively defines the interaction between the enclosure subfacets. A notable metric of the view factor calculation for closed surfaces (watertight enclosures) is that for each facet \( i \) the row-sum over all surface facets \( k \), \( \sum_{j=0}^{j=k} F_{ij} = 1.0 \).

In many cases a more complete model of the two-body configuration will include radiative transfer with the surroundings. Including these interactions requires introduction of an additional enclosure surface as shown in Figure 22.9.

Surface 3 known as the "partial enclosure" is artificially added to the numerical model (i.e. not included in the meshed discretization). Introduction of the partial enclosure is necessary in order to apply conventional approaches for numerical evaluation of the system view factors. Here we note that algorithmically, failure to add the partial enclosure surface would be manifest in bad row-sum metrics.

Introduction of the partial enclosure leads to the thermal network representation shown in Figure 22.10 in which the partial enclosure interacts with the other bodies of the system. Since the partial enclosure is modeled in the same way as a real surface questions often arise concerning characterization of the partial enclosure in terms of its temperature, area and emissivity. The temperature must of course must represent the surroundings. The area should be an area that envelopes the true surfaces of the enclosures. The partial enclosure emissivity should be chosen in a manner consistent with the relationship of surface radiosity \( J \)
with the emission, $\sigma T^4_3$ and irradiance, $G$

$$J = \varepsilon \sigma T^4 - (1 - \varepsilon) G$$

As examples of this we consider the consequence of selecting a partial emissivity at two extremes, $\varepsilon = 0$ and $\varepsilon = 1$, both of which reduce the network model of Figure 22.10 to that of 22.11. For $\varepsilon = 0$ from

the network interaction between $\sigma T^4_3 - J_3$ we find that the resistance becomes infinite thus eliminating any system interaction with $T_3$ as all the incident energy is reflected. This results in a modified version of the two-surface network 22.7, albeit with more attenuation. For $\varepsilon = 1$ and the network interaction between $\sigma T^4_3 - J_3$ is eliminated. However, now the radiative transfer system sees the emissive power of the surroundings since from 22.5.1, $J = \sigma T^4$. Clearly there are no obvious choices for partial enclosure emissivity other than that it should be chosen to enable interaction of the system with the surroundings.

### 22.6 Viewfactor Calculation

Scope: Equation System
Begin Viewfactor Calculation \( V_f\text{calc} \)

Bsp Tree Max Depth \(\{=\mid are\mid is\}\ depth\) And Min List Length \(\{=\mid are\mid is\}\ l\)
Compute Rule \(\{=\mid are\mid is\}\ \text{VFComputeRule}\)
Geometric Tolerance \(\{=\mid are\mid is\}\ n\)
Hemicube Max Subdivides \(\{=\mid are\mid is\}\ n\)
Hemicube Min Separation \(\{=\mid are\mid is\}\ n\)
Hemicube Resolution \(\{=\mid are\mid is\}\ n\)
Number Of Rotations \(\{=\mid are\mid is\}\ n\)
Output Rule \(\{=\mid are\mid is\}\ \text{OutputRule}\)
Pairwise Monte Carlo Sample Rule \(\{=\mid are\mid is\}\ \text{AMCSampleRule}\)
Pairwise Monte Carlo Tol1 \(\{=\mid are\mid is\}\ \text{Real\_value}\)
Pairwise Monte Carlo Tol2 \(\{=\mid are\mid is\}\ \text{Real\_value}\)
Pairwise Number Of Monte Carlo Samples \(\{=\mid are\mid is\}\ n\)
Pairwise Number Of Visibility Samples \(\{=\mid are\mid is\}\ n\)
Pairwise Visibility Sample Rule \(\{=\mid are\mid is\}\ \text{AVSampleRule}\)
X-Y Plane Symmetry
X-Z Plane Symmetry
Y-Z Plane Symmetry

End

Summary

This block command specifies a radiation enclosure and is used to define a method for calculating view factors. The parameter for this block corresponds to an instance of a radiation enclosure mechanics.

22.6.1 Bsp Tree Max Depth

Scope: Viewfactor Calculation

Bsp Tree Max Depth \(\{=\mid are\mid is\}\ depth\) And Min List Length \(\{=\mid are\mid is\}\ l\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(depth)</td>
<td>integer</td>
<td>12</td>
</tr>
<tr>
<td>(l)</td>
<td>integer</td>
<td>25</td>
</tr>
</tbody>
</table>

Summary

This line command sets the BSP tree parameters.

Description

This line command sets the parameters for the Binary Space Partitioning (BSP) tree. This data structure is used to accelerate the geometric operations associated with visibility to calculate the view factors. The maximum depth of the tree will set the maximum number of partitioning boxes created for a given enclosure. The minimum list length will set the desired number of faces to be contained in a box. The depth of the tree will be increased until the minimum list length is met of the maximum depth is reached. It is suggested that these parameters not be adjusted from their default values.
### 22.6.2 Compute Rule
**Scope:** Viewfactor Calculation

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>VFComputeRule</td>
<td>{hemicube</td>
<td>pairwise</td>
</tr>
</tbody>
</table>

**Summary**
This line command sets the method for computing the view factors for this enclosure.

### 22.6.3 Geometric Tolerance
**Scope:** Viewfactor Calculation

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(n)</td>
<td>real</td>
<td>1.0e-6</td>
</tr>
</tbody>
</table>

**Summary**
Set the geometric tolerance

**Description**
This line command sets the spatial geometry tolerance for use in all geometry-related comparisons. This value should correspond to the known minimum geometric size of the element faces used to define the enclosure.

### 22.6.4 Hemicube Max Subdivides
**Scope:** Viewfactor Calculation

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(n)</td>
<td>integer</td>
<td>2</td>
</tr>
</tbody>
</table>

**Summary**
Set the upper limit of hemicube subdivides

**Description**
This line command sets the upper limit of the number of element-face subdivides that will occur when the proximity criteria limit is exceeded (see the HEMICUBE MIN SEPARATION line command).

### 22.6.5 Hemicube Min Separation
**Scope:** Viewfactor Calculation

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(n)</td>
<td>real</td>
<td>5</td>
</tr>
</tbody>
</table>

**Summary**
Set the hemicube minimum separation
This line command sets a nondimensional minimum-separation distance allowed between element faces for the HEMICUBE method before the face is subdivided. The accuracy of the hemicube method degrades rapidly if the element faces are in close proximity to each other. This tolerance is based on dividing the normal distance between the center of the element faces by the effective diameter of the element face.

### 22.6.6 Hemicube Resolution

**Scope:** Viewfactor Calculation

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hemicube Resolution {=</td>
<td>are</td>
<td>is} ( n )</td>
</tr>
</tbody>
</table>

**Summary**
Set the hemicube resolution

**Description**
This line command sets the number of uniform subpatches into which the hemicube will be divided.

### 22.6.7 Number Of Rotations

**Scope:** Viewfactor Calculation

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number Of Rotations {=</td>
<td>are</td>
<td>is} ( n )</td>
</tr>
</tbody>
</table>

**Summary**
Set the number of internal rotations for 2D axisymmetric geometry or 3D geometry with rotation symmetry.

### 22.6.8 Output Rule

**Scope:** Viewfactor Calculation

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Output Rule {=</td>
<td>are</td>
<td>is} ( OutputRule )</td>
</tr>
</tbody>
</table>

**Summary**
Toggle verbose reporting

**Description**
This line command determines the amount of information that is printed to the screen during the view-factor calculation.

### 22.6.9 Pairwise Monte Carlo Sample Rule

**Scope:** Viewfactor Calculation
Pairwise Monte Carlo Sample Rule {=|are|is} AMCSampleRule

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>AMCSampleRule</td>
<td>{halton</td>
<td>jitter</td>
</tr>
</tbody>
</table>

Description: This line command selects the method for the distribution of Monte Carlo integration sample points on the surface for view-factor quadrature.

22.6.10 Pairwise Monte Carlo Tol1

Scope: Viewfactor Calculation

<table>
<thead>
<tr>
<th>Parameter Monte Carlo Tol1 {=</th>
<th>are</th>
<th>is} Real_value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Real_value</td>
<td>Value</td>
<td>Default</td>
</tr>
<tr>
<td>real</td>
<td>real</td>
<td>1.0e-5</td>
</tr>
</tbody>
</table>

Summary: Set first of two convergence checks for Monte Carlo integration

Description: This line command sets one of the two convergence checks used by the Monte Carlo integration algorithm for the PAIRWISE method. If the standard deviation of the view factor divided by the value of the view factor is less than this tolerance, the Monte Carlo integration will terminate, i.e.

\[
\frac{\text{Std. Dev. of } F_{ij}}{F_{ij}} < tol_1
\]

22.6.11 Pairwise Monte Carlo Tol2

Scope: Viewfactor Calculation

<table>
<thead>
<tr>
<th>Parameter Monte Carlo Tol2 {=</th>
<th>are</th>
<th>is} Real_value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Real_value</td>
<td>Value</td>
<td>Default</td>
</tr>
<tr>
<td>real</td>
<td>real</td>
<td>1.0e-5</td>
</tr>
</tbody>
</table>

Summary: Set second of two convergence checks for Monte Carlo integration

Description: This line command sets one of the two convergence checks used by the Monte Carlo integration algorithm for the PAIRWISE method. If the standard deviation of the view factor is less than this tolerance, the Monte Carlo integration will terminate, i.e.

\[
\text{Std. Dev. of } F_{ij} < tol_2
\]

22.6.12 Pairwise Number Of Monte Carlo Samples

Scope: Viewfactor Calculation

<table>
<thead>
<tr>
<th>Parameter Number Of Monte Carlo Samples {=</th>
<th>are</th>
<th>is} n</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>Value</td>
<td>Default</td>
</tr>
<tr>
<td>integer</td>
<td>integer</td>
<td>1000</td>
</tr>
</tbody>
</table>
Set the pairwise number of Monte Carlo sample points when Monte Carlo integration is activated during the PAIRWISE view-factor-calculation algorithm.

**22.6.13 Pairwise Number Of Visibility Samples**

**Scope:** Viewfactor Calculation

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pairwise Number Of Visibility Samples</td>
<td>=</td>
<td>are</td>
</tr>
</tbody>
</table>

Summary: Set the pairwise number visibility sample points

Description: This line command will set the number of sample points to use for evaluating visibility between element faces for the PAIRWISE view-factor-calculation algorithm. Visibility is used to select the method for calculating the view factors. If the two element faces are completely visible to each other, the view factor is calculated using Gauss quadrature or an analytic method. If the element faces are partially visible, Monte Carlo integration is used.

**22.6.14 Pairwise Visibility Sample Rule**

**Scope:** Viewfactor Calculation

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pairwise Visibility Sample Rule</td>
<td>=</td>
<td>are</td>
</tr>
</tbody>
</table>

Summary: Set the pairwise visibility sample rule

Description: This line command selects the method for distributing the visibility sample points on the surface.

**22.6.15 X-Y Plane Symmetry**

**Scope:** Viewfactor Calculation

Summary: Specifies symmetry about the X-Y plane.

**22.6.16 X-Z Plane Symmetry**

**Scope:** Viewfactor Calculation

Summary: Specifies symmetry about the X-Z plane.
22.6.17 Y-Z Plane Symmetry
Scope: Viewfactor Calculation

Summary Specifies symmetry about the Y-Z plane.

22.7 Viewfactor Smoothing
Scope: Equation System

Begin Viewfactor Smoothing Vf_smooth
   Convergence Tolerance {=} Param1
   Maximum Iterations {=} n
   Method {=} VFSmoothMethod
   Output Rule {=} OutputRule
   Reciprocity Rule {=} VFMatrixSymmRule
   Weight Power {=} Param1
End

Summary Defines a view factor smoothing scheme and its associated parameters.

Description This block command is used to define a method for smoothing the view factors. The defined method will be used for view factor smoothing only when requested with the USE VIEWFACTOR SMOOTHING line command in the ENCLOSURE DEFINITION command block.

When VIEWFACTOR SMOOTHING is requested a two step sequence is initiated
1) Reciprocity enforcement using the specified RECIPROCITY RULE
2) Rowsum enforcement as per the specified METHOD

In both steps the viewfactor matrix will be modified as prescribed.

**Note**
In step 1 the appropriate modifications are always applied. However, if the rowsum enforcement criteria of step 2 are not satisfied the viewfactor matrix supplied to the enclosure radiation calculation will be the viewfactor matrix resulting from step 1 and -not- the raw viewfactor matrix.

22.7.1 Convergence Tolerance
Scope: Viewfactor Smoothing

Convergence Tolerance {=} Param1

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Param1</td>
<td>real</td>
<td>1.0e-8</td>
</tr>
</tbody>
</table>

696
Summary
Sets convergence tolerance.

Description
This line command sets the convergence tolerance for both the SIMPLE and LEAST-SQUARES smoothing algorithms based on the following:

1) SIMPLE: $|1 - \sum F_{ij}| < tol$
2) LEAST-SQUARES: $\|r\| < tol$,

where $r$ is the residual of the Lagrange multiplier matrix problem.

22.7.2 Maximum Iterations
Scope: Viewfactor Smoothing

Maximum Iterations {=|are|is} $n$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$n$</td>
<td>integer</td>
<td>500</td>
</tr>
</tbody>
</table>

Summary
Set maximum iterations.

Description
This line command sets the maximum number of iterations that the SIMPLE smoothing algorithm will take.

22.7.3 Method
Scope: Viewfactor Smoothing

Method {=|are|is} VFSmoothMethod

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>VFSmoothMethod</td>
<td>{least-squares</td>
<td>none</td>
</tr>
</tbody>
</table>

Summary
Defines view factor smoothing method.

Description
This line command defines the algorithm for smoothing the view factors. Smoothing is a term used to describe the process of enforcing the row-sum property of the view-factor matrix given by

$$\sum F_{ij} = 1.0$$

and the reciprocity property of view factors given by

$$A_i F_{ij} = A_j F_{ji}.$$ The HEMICUBE method guarantees the row-sum property, but not necessarily the reciprocity property. The PAIRWISE method will guarantee the reciprocity property (lower diagonal is calculated and the upper triangular is filled in using reciprocity), but not necessarily the row-sum property. If these two properties are not met by the view-factor matrix, energy conservation will not be achieved.
22.7.4 Output Rule

Scope: Viewfactor Smoothing

Output Rule \{=|are|is\} OutputRule

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>OutputRule</td>
<td>{more verbose</td>
<td>none</td>
</tr>
</tbody>
</table>

Summary       Set the amount of output.

Description   This line command sets the amount of information reported to the output screen about the view-factor smoothing.

22.7.5 Reciprocity Rule

Scope: Viewfactor Smoothing

Reciprocity Rule \{=|are|is\} VFMatrixSymmRule

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>VFMatrixSymmRule</td>
<td>{addition</td>
<td>average</td>
</tr>
</tbody>
</table>

Summary       Selects reciprocity enforcement rule.

Description   This line command determines the method for enforcing reciprocity,

\[ A_i F_{ij} = A_j F_{ji} \]

during smoothing. This command is only required for the HEMICUBE algorithm. The PAIRWISE algorithm uses reciprocity to fill in the lower triangular part of the view-factor matrix. If both of the view factors, \( F_{ij} \) and \( F_{ji} \), are nonzero, these factors are adjusted by averaging \( A_i F_{ij} \) and \( A_j F_{ji} \).

22.7.6 Weight Power

Scope: Viewfactor Smoothing

Weight Power \{=|are|is\} Param1

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Param1</td>
<td>real</td>
<td>2.0</td>
</tr>
</tbody>
</table>

Summary       Sets weight power.

Description   This line command sets the power, \( p \), used for the weights in the LEAST-SQUARES smoothing algorithm

\[ w_{ij} = F_{ij}^p. \]
22.8 Radiosity Solver

Scope: Equation System

Begin Radiosity Solver Boundary condition instance name
  Convergence Tolerance {=|are|is} Tolerance
  Coupling {=|are|is} RadCouplingRule
  Maximum Iterations {=|are|is} m
  Output Rule {=|are|is} OutputRule
  Solver {=|are|is} RadSolveRule
End

Summary Sets convergence tolerance.

Description This line command sets the convergence tolerance for the iterative solution of the radiosity system. The iterative solution will stop when the ratio of the current residual to the initial residual is less than the specified tolerance:

\[
\frac{\|r\|}{\|r_0\|} < tol
\]

22.8.1 Convergence Tolerance

Scope: Radiosity Solver

Convergence Tolerance {=|are|is} Tolerance

Parameter Value Default
Tolerance real 1.0e-6

Summary Sets convergence tolerance.

Description This line command sets the convergence tolerance for the iterative solution of the radiosity system. The iterative solution will stop when the ratio of the current residual to the initial residual is less than the specified tolerance:

\[
\frac{\|r\|}{\|r_0\|} < tol
\]

22.8.2 Coupling

Scope: Radiosity Solver

Coupling {=|are|is} RadCouplingRule

Parameter Value Default
RadCouplingRule {lagged|mason|smoothed} MASON

Summary Specifies linearization method.

Description This line command specifies the linearization method used to apply the radiative heat flux to the thermal model.
22.8.3 Maximum Iterations
Scope: Radiosity Solver

Maximum Iterations \( m \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( m )</td>
<td>integer</td>
<td>300</td>
</tr>
</tbody>
</table>

Summary: Sets maximum number of iterations.

Description: This line command sets the maximum number of iterations allowed for the iterative solution of the radiosity equations. If the maximum number of iterations is exceeded, the calculation will fail and the thermal simulation will stop.

22.8.4 Output Rule
Scope: Radiosity Solver

Output Rule \( OutputRule \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( OutputRule )</td>
<td>{more verbose</td>
<td>none</td>
</tr>
</tbody>
</table>

Summary: Sets the information reporting level.

Description: This line command will set the amount of information reported to the output screen about the radiosity solution.

22.8.5 Solver
Scope: Radiosity Solver

Solver \( RadSolveRule \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( RadSolveRule )</td>
<td>{chaparral cg</td>
<td>chaparral gmres</td>
</tr>
</tbody>
</table>

Summary: Chaparral solver selection for radiosity system.

Description: This line command selects the method for solving the linear system of radiosity equations. The currently available methods are iterative linear solvers.

22.9 Enclosure Definition
Scope: Equation System

Begin Enclosure Definition Boundary condition instance name
Activate Mean Beam Model With Bulk Node Model
Add Surface SurfaceList...
Area Output VariableName
Blocking Surfaces
Dash Closure Metric Samples {=|are|is} NumSamp [ Tol {=|are|is} Value ]
Dash Solve Enclosures {=|are|is} Values...
Dash Solve Enclosures Union {=|are|is} Values...
Database Name {=|are|is} Filename In VFILEFormat Format
Emissivity {=|are|is} Value [ On SurfaceName ]
Emissivity Function {=|are|is} FunctionName [ On SurfaceName ]
Emissivity Subroutine {=|are|is} MySub [ On SurfaceName ]
Emissivity Time Function {=|are|is} FunctionName [ On SurfaceName ]
Input Database Name {=|are|is} Filename
Integrated Flux Output VariableName
Integrated Power Output VariableName
Matched Flux On SurfaceName {=|are|is} FluxType [ Phase {=|are|is} MaterialPhaseName ]
Mean Beam Length {=|are|is} l [Parameters]...
Meshed Enclosure Is Blocklist...
Nonblocking Surfaces
Output Database Name {=|are|is} Filename In VFILEFormat Format
Overlapping Enclosure {=|are|is} Bool
Partial Enclosure Area {=|are|is} a
Partial Enclosure Area Subroutine {=|are|is} FName
Partial Enclosure Area Time Function {=|are|is} FName
Partial Enclosure Emissivity {=|are|is} e
Partial Enclosure Emissivity Subroutine {=|are|is} FName
Partial Enclosure Emissivity Time Function {=|are|is} FName
Partial Enclosure Flux Output VariableName
Partial Enclosure Irradiance Output VariableName
Partial Enclosure Radiosity Output VariableName
Partial Enclosure Temperature {=|are|is} t
Partial Enclosure Temperature Subroutine {=|are|is} FName
Partial Enclosure Temperature Time Function {=|are|is} FName
Preprocess Enclosures [ Options... ]
Radiosity Database Name {=|are|is} Filename
Rowsum Database Name {=|are|is} Filename
Topology Database Name {=|are|is} Filename
Use Banded Wavelength Model ModelName
Use Dash Enclosures [ type tol {=|are|is} Params... ]
Use Radiosity Solver Param0
Use Toggle Block ToggleName [ \{at\ for\ in\ on\over\} ElementBlockList... ]
Use Viewfactor Calculation Param0
Use Viewfactor Smoothing Param0
Viewfactor Update \{are\is\} UpdateMethod [ Using Params... ]
Viewfactor Update Start Time Is Start_time
Begin Band BandName
End

Summary     Specifies a radiation enclosure. Corresponds to an instance of radiation enclosure mechanics.
Description  This block command is used to define an enclosure for the thermal model. There may be
more than one enclosure defined for any thermal model. An enclosure definition includes the
geometric aspects (list of element faces forming the surface), material properties, and setting
of the algorithms for calculating the view factors between the element faces in the enclosure
and the radiosity solution. The name of the enclosure is specified by the user.
** NOTE ** Enclosure names cannot begin with any sequence of numeric symbols (i.e. 0 -
9). This will cause a model to fail with very little in the way of error reporting.

22.9.1 Activate Mean Beam Model With Bulk Node
Scope:    Enclosure Definition

Activate Mean Beam Model With Bulk Node Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary     Sets the bulk element representing the PMR material for the MBL (mean beam length)
enclosure.

22.9.2 Add Surface
Scope:    Enclosure Definition

Add Surface SurfaceList...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>SurfaceList</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary     Adds surfaces, by name, to a boundary condition’s extent.

Description  This line command is used to add surfaces to the extent of a boundary condition. In Exodus
II, surfaces are specified as side sets, that have a global integer identifier. For example, side
set 12 would be added by this line command using the surface name surface_12. Note that
in SIERRA, each element of an array of strings must be separated by whitespace.
22.9.3 Area Output
Scope: Enclosure Definition

<table>
<thead>
<tr>
<th>VariableName</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>VariableName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Output the area associated with this flux boundary condition.

Description This line triggers output of the enclosure area and volume.

22.9.4 Blocking Surfaces
Scope: Enclosure Definition

Summary Specifies a non-blocking enclosure

Description This line command informs Chaparral to consider blocking surfaces during the calculation of view factors. A blocking surface is defined as one or more element faces that occlude the view from any pair of element faces within the enclosure. If it is known that no blocking surfaces exist in the enclosure, this line command should not be included in the enclosure definition. This will help to reduce the compute time for view factors. Note that this line command should be included if there is any doubt whether there are blocking surfaces in the enclosure.

22.9.5 Dash Closure Metric Samples
Scope: Enclosure Definition

<table>
<thead>
<tr>
<th>NumSamp</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>integer</td>
<td>undefined</td>
<td></td>
</tr>
</tbody>
</table>

Summary Specify number of samples for DASH closure metric and possibly an acceptable tolerance.

Description When using DASH enclosures one may wish to assign the number of sample points used in determining closure. Optionally, one can set the acceptable value of closure metric associated with closure.

22.9.6 Dash Solve Enclosures
Scope: Enclosure Definition

<table>
<thead>
<tr>
<th>Values</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>integer</td>
<td>undefined</td>
<td></td>
</tr>
</tbody>
</table>

Dash Solve Enclosures {are|is} Values...
Summary
List of integer index values for Dash enclosures to perform solution. The index values must be obtained by a preprocessing step using both PREPROCESS ENCLOSURE TOPOLOGY command line at the Region level and the PREPROCESS ENCLOSURES command line in the Enclosure Definition command block.

22.9.7 Dash Solve Enclosures Union
Scope: Enclosure Definition

Dash Solve Enclosures Union {=|are|is} Values...

Parameter | Value | Default
---|---|---
Values | integer... | undefined

Summary
List of integer index values for Dash enclosures to perform solution. The index values must be obtained by a preprocessing step using both PREPROCESS ENCLOSURE TOPOLOGY command line at the Region level and the PREPROCESS ENCLOSURES command line in the Enclosure Definition command block.

22.9.8 Database Name
Scope: Enclosure Definition

Database Name {=|are|is} Filename In VFileFormat Format

Parameter | Value | Default
---|---|---
Filename | string undefined | undefined
VFileFormat | {ascii|binary} undefined | undefined

Summary
Specifies common filename to read/write viewfactors

Description
This line command specifies the database name for both input and output of the view factors for this enclosure. The file name is provided by the user, and one of two file formats is specified.

The ASCII format should be used only for small enclosures. This format is useful when the analyst wants to examine the values of the view factors that have been evaluated.

The BINARY format will produce a machine-dependent file. This view-factor file can only be read on the same machine in subsequent calculations.

22.9.9 Emissivity
Scope: Enclosure Definition

Emissivity {=|are|is} Value [ On SurfaceName ]

Parameter | Value | Default
---|---|---
Value | real | undefined

Summary
Sets a constant value of emissivity for a defined surface. If the optional parameters are not included, then this is the default emissivity for the enclosure. Otherwise it is only applied to the indicated surface.
**Note that if a surface is called out more than once, the emissivity definition it is overwritten: last one in wins.**

### 22.9.10 Emissivity Function

**Scope:** Enclosure Definition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>FunctionName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Sets an emissivity function for a defined surface. If the optional parameters are not included, then this is the default emissivity for the enclosure. Otherwise it is only applied to the indicated surface.

**Note that if a surface is called out more than once, the emissivity definition it is overwritten: last one in wins.**

### 22.9.11 Emissivity Subroutine

**Scope:** Enclosure Definition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>MySub</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Sets an emissivity user subroutine for a defined surface. If the optional parameters are not included, then this is the default emissivity for the enclosure. Otherwise it is only applied to the indicated surface.

Also, the software supports using locally scoped user data for most user subroutines, but I haven’t figured out a syntax for it here yet. So it is not yet supported. If you need to get data into this subroutine, use the region’s "REAL DATA" and "INTEGER DATA" line commands.

**Note that if a surface is called out more than once, the emissivity definition it is overwritten: last one in wins.**

### 22.9.12 Emissivity Time Function

**Scope:** Enclosure Definition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>FunctionName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Sets an emissivity function of time for a defined surface. If the optional parameters are not included, then this is the default emissivity for the enclosure. Otherwise it is only applied to the indicated surface.

**Note that if a surface is called out more than once, the emissivity definition it is overwritten: last one in wins.**
22.9.13 Input Database Name

Scope: Enclosure Definition

Input Database Name {=|are|is} Filename

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Filename</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies filename to read viewfactors from

Description: This line command provides the name of the view-factor file. If this line command is present, the file containing the given view-factor matrix will be opened and read. The calculation of view factors will not take place. Note that you must also specify a value of READ for the COMPUTE RULE command in the associated VIEWFACTOR CALCULATION command block.

22.9.14 Integrated Flux Output

Scope: Enclosure Definition

Integrated Flux Output VariableName

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>VariableName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Calculate the average flux associated with this flux boundary condition.

Description: This line command specifies that, as a postprocess, the normal flux associated with this boundary condition be integrated over the surface to obtain the total power. This power is then divided by the total area of the surface to obtain the average flux on the surface, and stored in a global variable named "variableName". This global variable may then be output to history files, or accessed in user subroutines, etc.

22.9.15 Integrated Power Output

Scope: Enclosure Definition

Integrated Power Output VariableName

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>VariableName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Calculate the total power associated with this flux boundary condition.

Description: This line command specifies that, as a postprocess, the normal flux associated with this boundary condition be integrated over the surface to obtain the total power which is then stored into a global variable named "variableName". This global variable may then be output to history files, or accessed in user subroutines, etc.
22.9.16  Matched Flux On
Scope:  Enclosure Definition

Matched Flux On SurfaceName {=|are|is} FluxType [ Phase {=|are|is} MaterialPhaseName ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>SurfaceName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>FluxType</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Indicates an equivalent matching flux for surfaces within the enclosure that border element
blocks which do not have the ENERGY equation defined but do have an equivalent equation
(such as ENTHALPY) defined. The primary use for this is in OMD simulations and should
be considered at the same reliability level as OMD.

22.9.17  Mean Beam Length
Scope:  Enclosure Definition

Mean Beam Length {=|are|is} l [Parameters]...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>l</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Parameters</td>
<td>[string]...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Sets the expression for the radiant decay length L for the MBL (mean beam length) enclosure.

22.9.18  Meshed Enclosure
Scope:  Enclosure Definition

Meshed Enclosure Is BlockList...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>BlockList</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Adds blocks, by name, to the meshed enclosure extent.

Description  In the event one wishes to solve a conduction problem within an enclosure as well as the
radiation problem the interior discretization must be defined. This line command is used to
add blocks to the extent of a enclosure interior. In Exodus II, surfaces are specified as side
sets, that have a global integer identifier. For example, block 12 would be added by this line
command using the surface name block_12. Note that in SIERRA, each element of an array
of strings must be separated by whitespace.

22.9.19  Nonblocking Surfaces
Scope:  Enclosure Definition

Summary  Specifies a blocking enclosure
### 22.9.20 Output Database Name

**Scope:** Enclosure Definition

Output Database Name `{=|are|is} Filename In VFileFormat Format

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Filename</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>VFileFormat</td>
<td>{ascii</td>
<td>binary}</td>
</tr>
</tbody>
</table>

**Summary**  Specify filename to write viewfactors to

**Description** This line command activates the output of the view factors calculated during a simulation. The file name is provided by the user, and one of three file formats is specified. The ASCII format should be used only for small enclosures. This format is useful when the analyst wants to examine the values of the view factors that have been evaluated. The BINARY format will produce a machine-dependent file. This view-factor file can only be read on the same machine in subsequent calculations.

### 22.9.21 Overlapping Enclosure

**Scope:** Enclosure Definition

Overlapping Enclosure `{=|are|is} Bool

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bool</td>
<td>{false</td>
<td>true}</td>
</tr>
</tbody>
</table>

**Summary** Designates this enclosure as overlapping other enclosures.

**Description** On occasion one might want to define an enclosure radiation problem as a superposition of several enclosures. In this case one must designate one of the enclosures as overlapping the others in order to distinguish it from the other enclosures. This command adds a qualifier to the enclosure definition to enable making this distinction possible.

### 22.9.22 Partial Enclosure Area

**Scope:** Enclosure Definition

Partial Enclosure Area `{=|are|is} a

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary** Constant value for the partial enclosure area associated with this enclosure radiation flux boundary condition.

### 22.9.23 Partial Enclosure Area Subroutine

**Scope:** Enclosure Definition
Partial Enclosure Area Subroutine \{=|are|is\} **FName**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>FName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: User-defined function name for the partial enclosure area associated with this enclosure radiation flux boundary condition.

### 22.9.24 Partial Enclosure Area Time Function

**Scope:** Enclosure Definition

| Partial Enclosure Area Time Function \{=|are|is\} **FName**
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>FName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Time-dependent function name for the partial enclosure area associated with this enclosure radiation flux boundary condition.

### 22.9.25 Partial Enclosure Emissivity

**Scope:** Enclosure Definition

| Partial Enclosure Emissivity \{=|are|is\} **e**
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>e</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Constant value for the partial enclosure emissivity associated with this enclosure radiation flux boundary condition.

### 22.9.26 Partial Enclosure Emissivity Subroutine

**Scope:** Enclosure Definition

| Partial Enclosure Emissivity Subroutine \{=|are|is\} **FName**
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>FName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: User-defined function name for the partial enclosure emissivity associated with this enclosure radiation flux boundary condition.

### 22.9.27 Partial Enclosure Emissivity Time Function

**Scope:** Enclosure Definition

| Partial Enclosure Emissivity Time Function \{=|are|is\} **FName**
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>FName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Summary  Time-dependent function name for the partial enclosure emissivity associated with this enclosure radiation flux boundary condition.

22.9.28  Partial Enclosure Flux Output
Scope:  Enclosure Definition

Partial Enclosure Flux Output  **VariableName**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>VariableName</strong></td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Output the flux associated with the partial enclosure.

Description  During the simulation the flux associated with the partial enclosure is output to the ExodusII file as a global variable.

22.9.29  Partial Enclosure Irradiance Output
Scope:  Enclosure Definition

Partial Enclosure Irradiance Output  **VariableName**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>VariableName</strong></td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Output the irradiance associated with the partial enclosure.

Description  During the simulation the irradiance associated with the partial enclosure is output to the ExodusII file as a global variable.

22.9.30  Partial Enclosure Radiosity Output
Scope:  Enclosure Definition

Partial Enclosure Radiosity Output  **VariableName**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>VariableName</strong></td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Output the radiosity associated with the partial enclosure.

Description  During the simulation the radiosity associated with the partial enclosure is output to the ExodusII file as a global variable.
22.9.31 Partial Enclosure Temperature

Scope: Enclosure Definition

Partial Enclosure Temperature \( t \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( t )</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Constant value for the partial enclosure temperature associated with this enclosure radiation flux boundary condition.

22.9.32 Partial Enclosure Temperature Subroutine

Scope: Enclosure Definition

Partial Enclosure Temperature Subroutine \( FName \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( FName )</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
User-defined function name for the partial enclosure temperature associated with this enclosure radiation flux boundary condition.

22.9.33 Partial Enclosure Temperature Time Function

Scope: Enclosure Definition

Partial Enclosure Temperature Time Function \( FName \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( FName )</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Time-dependent function name for the partial enclosure temperature associated with this enclosure radiation flux boundary condition.

22.9.34 Preprocess Enclosures

Scope: Enclosure Definition

Preprocess enclosures subject to options: ORIGINAL_FACES, ENCLOSURE_FACETS, ENCLOSURE_FACET_EDGES, CONTACT_FACETS, CONTACT_FACET_EDGES and COARSE_MESH. When invoked at Region preprocessing is performed and the simulation will terminate.

Description
For debug and inspection purposes the surfaces defined in the enclosure definitions will be processed, then output to the exodus database before terminating the simulation. If no options are chosen the default is to select all the available information. Options are: ORIGINAL_FACES, ENCLOSURE_FACETS, ENCLOSURE_FACET_EDGES, CONTACT_FACETS, CONTACT_FACET_EDGES and COARSE_MESH. Facets are output as triangle elements, edges as bars and faces as shell elements.
22.9.35 Radiosity Database Name

Scope: Enclosure Definition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Radiosity Database Name {=</td>
<td>are</td>
<td>is} Filename</td>
</tr>
</tbody>
</table>

Summary: Specify filename to write enclosure rowsum error and radiosity results to

Description: This line command activates the output of the view factor rowsum error calculated during a simulation. Additionally, this file will be used to also output the radiosity and flux values during subsequent calls to the radiosity solver. The file name is provided by the user and the output format is ExodusII. This file is useful when the analyst wants to where in the enclosure the maximum errors are occurring. The rowsum error is given by

\[ 1 - \sum_{j=1}^{N_{\text{faces}}} F_{ij} \]

for each row i in the view-factor matrix (which corresponds to a face in the enclosure). Use of this option will negatively affect performance and is intended for use by developers.

22.9.36 Rowsum Database Name

Scope: Enclosure Definition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rowsum Database Name {=</td>
<td>are</td>
<td>is} Filename</td>
</tr>
</tbody>
</table>

Summary: Specify filename to write enclosure rowsum error results to

Description: This line command activates the output of the view factor rowsum error calculated during a simulation. The file name is provided by the user and the output format is ExodusII. This file is useful when the analyst wants to where in the enclosure the maximum errors are occurring. The rowsum error is given by

\[ 1 - \sum_{j=1}^{N_{\text{faces}}} F_{ij} \]

for each row i in the view-factor matrix (which corresponds to a face in the enclosure).

22.9.37 Topology Database Name

Scope: Enclosure Definition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Topology Database Name {=</td>
<td>are</td>
<td>is} Filename</td>
</tr>
</tbody>
</table>
**Summary** Specify filename to write enclosure topology to

**Description** This line command activates the output of the enclosure topology, taking into account all specified symmetry. The file name is provided by the user and the output format is ExodusII.

### 22.9.38 Use Banded Wavelength Model

**Scope:** Enclosure Definition

**Use Banded Wavelength Model**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>ModelName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary** Use a specified banded wavelength model for use in enclosure radiation.

**Description** Requests that a specific banded wavelength model named modelName, be associated with an enclosure, where the modelName is defined outside of the enclosure definition block. Only one BANDED WAVELENGTH MODEL can be used for an enclosure.

### 22.9.39 Use Dash Enclosures

**Scope:** Enclosure Definition

**Summary** Perform DASH contact search on the enclosure surfaces.

**Description** Performing a DASH contact search on the enclosure surfaces will prevent any faces in contact from being processed as part of an enclosure. Optionally one may define a relative tolerance based upon the characteristic element length, or absolute tolerances in positive and negative directions from the enclosure surface.

### 22.9.40 Use Radiosity Solver

**Scope:** Enclosure Definition

**Use Radiosity Solver**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Param0</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary** Specifies which radiosity solver to use.

**Description** This line command is used to identify the radiosity solver for use with this enclosure radiation problem. The details of the radiosity solver are defined using the RADIOSITY SOLVER command block. The parameter for this line command is the parameter used with the RADIOSITY SOLVER block command.
22.9.41 Use Toggle Block
Scope:  Enclosure Definition

Use Toggle Block $ToggleName$ [ ${@|at|for|in|on|over}$ $ElementBlockList$... ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$ToggleName$</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Specification for toggling entities in the computational model based on Toggle Block parameters. When used at the region level, the list of element blocks to be toggled must be provided. Otherwise a listing of entities is not needed as the Toggle Block will be associated with the command line or the enclosing command block.

22.9.42 Use Viewfactor Calculation
Scope:  Enclosure Definition

Use Viewfactor Calculation $Param0$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$Param0$</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Specifies which view factor calculation to use.

Description  This line command is used to specify the name of a method for calculating view factors defined using the VIEWFACTOR CALCULATION command block. The parameter for this line command is the parameter used to name the VIEWFACTOR CALCULATION block command.

22.9.43 Use Viewfactor Smoothing
Scope:  Enclosure Definition

Use Viewfactor Smoothing $Param0$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$Param0$</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Specifies which view factor smoother to use.

Description  This line command is used to specify the name of a method for smoothing view factors defined using the VIEWFACTOR SMOOTHING command block. The parameter for this line command is the parameter used with the VIEWFACTOR SMOOTHING block command.

22.9.44 Viewfactor Update
Scope:  Enclosure Definition

Viewfactor Update {=|are|is} $UpdateMethod$ [ Using $Params$... ]
### Update Method

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Method</td>
<td>{affected_area</td>
<td>continuous</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Summary</th>
<th>Specify the viewfactor re-compute strategy.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Description</td>
<td>The view factor calculation oftentimes dominates the computational cost of a thermal analysis. Thus for cases in which only the geometry changes due to surface deformation and the analyst views the change to be small enough so as to not affect the view factors greatly, one can select one of the strategies below for update of the view factors. STANDARD is the default re-compute strategy, update on mesh change. FREEZE means the view factor update is deactivated for subsequent updates. Thus the first view factor calculation determines the view factors used throughout the simulation. CONTINUOUS means the view factors will be computed every time step of a transient simulation. AFFECTED_AREA is relevant only when used with Element Death. For every view factor calculation the enclosure surface area is computed and stored as a reference area. At subsequent solution steps the area corresponding to elements which are candidates for Element Death is accumulated. The ratio of candidate death element area to the original area is used as the criteria of whether Element Death and a new view factor update will occur. If AFFECTED_AREA is chosen and Element Death is not present the re-compute strategy will revert to STANDARD. Here the Params value is the ratio of affected area to total area for which element death can occur. INTERVAL specifies the time interval between view factor updates. Here Params is a single value time increment. TIMES selects specific times at which the view factors are recalculated. Here Params can be several discrete time values. For INTERVAL and TIMES, one can also select a time at which view factor updates are considered using the VIEW FACTOR UPDATE START TIME command line.</td>
</tr>
</tbody>
</table>

### 22.9.45 Viewfactor Update Start Time

**Scope:** Enclosure Definition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Start_time</td>
<td>real</td>
<td>0.0</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Summary</th>
<th>Specify start time for possible re-compute of viewfactors.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Description</td>
<td>Specify the time after which viewfactor re-compute can occur. Command is active only for UPDATE FACTOR method = TIMES or INTERVAL.</td>
</tr>
</tbody>
</table>

### 22.10 Banded Wavelength Model

**Scope:** Equation System
Begin Banded Wavelength Model *ModelName*

Band *BandName* {=} {are|is} *LowValue* *HighValue*

End

**Summary**
This command block specifies the wavelength distribution for a banded wavelength model.

**Description**
Defines a piecewise constant banded wavelength discretization model of the thermal wavelength spectrum and associate it with name *modelName*. The same *modelName* can be used for different boundary conditions.

### 22.10.1 Band
**Scope:** Banded Wavelength Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>BandName</em></td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td><em>LowValue</em></td>
<td>real</td>
<td>0.0</td>
</tr>
<tr>
<td><em>HighValue</em></td>
<td>real</td>
<td>REAL_MAX</td>
</tr>
</tbody>
</table>

**Summary**
Specify the range of a wavelength band in microns μm for a surface or collection of surfaces undergoing radiative heat transfer.

**Description**
Defines the upper and lower limits of a wavelength band in microns μm. Upper and lower limit values can intersect those of adjacent bands but cannot overlap into previously a defined BAND.

### 22.11 Band
**Scope:** Enclosure Definition

Begin Band *BandName*

Emissivity {=} {are|is} *Value* [ On *SurfaceName* ]

Emissivity Function {=} {are|is} *FunctionName* [ On *SurfaceName* ]

Emissivity Subroutine {=} {are|is} *MySub* [ On *SurfaceName* ]

Emissivity Time Function {=} {are|is} *FunctionName* [ On *SurfaceName* ]

End

**Summary**
This nested command block specifies the emissivity distribution for the surfaces in the enclosure for the given band.

**Description**
Defines the emissivity distribution for the surfaces and/or default values for the band with name *bandName*.
22.11.1 Emissivity

Scope: Band

| Emissivity {=} |are| is} Value [ On SurfaceName ] |
|----------------|--------------------------------|
| **Parameter** | **Value** | **Default** |
| Value | real | undefined |

Summary

Sets a constant value of emissivity for a defined surface. If the optional parameters are not included, then this is the default emissivity for the enclosure. Otherwise it is only applied to the indicated surface.

**Note that if a surface is called out more than once, the emissivity definition it is overwritten: last one in wins.**

22.11.2 Emissivity Function

Scope: Band

| Emissivity Function {=} |are| is} FunctionName [ On SurfaceName ] |
|-------------------------|-------------------------------------|
| **Parameter** | **Value** | **Default** |
| FunctionName | string | undefined |

Summary

Sets a emissivity function for a defined surface. If the optional parameters are not included, then this is the default emissivity for the enclosure. Otherwise it is only applied to the indicated surface.

**Note that if a surface is called out more than once, the emissivity definition it is overwritten: last one in wins.**

22.11.3 Emissivity Subroutine

Scope: Band

| Emissivity Subroutine {=} |are| is} MySub [ On SurfaceName ] |
|---------------------------|--------------------------------|
| **Parameter** | **Value** | **Default** |
| MySub | string | undefined |

Summary

Sets a emissivity user subroutine for a defined surface. If the optional parameters are not included, then this is the default emissivity for the enclosure. Otherwise it is only applied to the indicated surface.

Also, the software supports using locally scoped user data for most user subroutines, but I haven’t figured out a syntax for it here yet. So it is not yet supported. If you need to get data into this subroutine, use the region’s "REAL DATA" and "INTEGER DATA" line commands.

**Note that if a surface is called out more than once, the emissivity definition it is overwritten: last one in wins.**
22.11.4  Emissivity Time Function

Scope:  Band

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>FunctionName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Sets a emissivity function of time for a defined surface. If the optional parameters are not included, then this is the default emissivity for the enclosure. Otherwise it is only applied to the indicated surface.

**Note that if a surface is called out more than once, the emissivity definition it is overwritten; last one in wins.

22.11.5  Dash Closure Metric Samples

Scope:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>NumSamp</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Specify number of samples for DASH closure metric and possibly an acceptable tolerance.

Description  When using DASH enclosures one may wish to assign the number of sample points used in determining closure. Optionally, one can set the acceptable value of closure metric associated with closure.

22.11.6  Dash Closure Metric Samples

Scope:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>NumSamp</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Specify number of samples for DASH closure metric and possibly an acceptable tolerance.

Description  When using DASH enclosures one may wish to assign the number of sample points used in determining closure. Optionally, one can set the acceptable value of closure metric associated with closure.

22.11.7  Preprocess Enclosures

Scope:
**Summary**  Preprocess enclosures subject to options: ORIGINAL_FACES, ENCLOSURE_FACETS, ENCLOSURE_FACET_EDGES, CONTACT_FACETS, CONTACT_FACET_EDGES and COARSE_MESH. When invoked at Region preprocessing is performed and the simulation will terminate.

**Description**  For debug and inspection purposes the surfaces defined in the enclosure definitions will be processed, then output to the exodus database before terminating the simulation. If no options are chosen the default is to select all the available information. Options are: ORIGINAL_FACES, ENCLOSURE_FACETS, ENCLOSURE_FACET_EDGES, CONTACT_FACETS, CONTACT_FACET_EDGES and COARSE_MESH. Facets are output as triangle elements, edges as bars and faces as shell elements.

### 22.11.8 Use Dash Enclosures

**Scope:**

**Summary**  Perform DASH contact search on the enclosure surfaces.

**Description**  Performing a DASH contact search on the enclosure surfaces will prevent any faces in contact from being processed as part of an enclosure. Optionally one may define a relative tolerance based upon the characteristic element length, or absolute tolerances in positive and negative directions from the enclosure surface.
Chapter 23

SPn Radiation Reference

23.1 Radiative Transport

Radiative transport, which is another mode of heat transfer, describes the spatial variation of radiative intensity corresponding to a given direction and at a given wavelength within a radiatively participating medium. This is governed by the Boltzmann Equation or Radiative Transport Equation (RTE) which represents a balance between absorption, emission and scattering. For the purpose of thermal analysis, we make certain assumptions that simplify the RTE. These are

1. Gray Media: The optical properties of the material medium are not a function of wavelength i.e no spectral banding.
2. Steady State: Energy transport is propagated by photons which travel at the speed of light. This means that there are no transient effects relative to the time scale of other physics.
3. Isotropic Scattering: Since the scattering phase function is rarely known, we simply assume isotropic scattering where the probability of intensity being propagated in a given direction has equal probability of being scattered in another direction.

Given the above assumptions, we may write the RTE as

\[ \Omega \cdot \nabla I(\Omega) + (\sigma_A + \sigma_S) I(\Omega) = \sigma_A I_b + \frac{\sigma_S}{4\pi} G, \]  \hspace{1cm} (23.1)

where \( \sigma_A \) is the absorption coefficient, \( \sigma_S \) is the scattering coefficient, \( I(\Omega) \) is the intensity along the direction \( \Omega \), \( T \) is the temperature and the angle-integrated intensity is \( G = \int_{4\pi} I(\Omega) d\Omega \). The black body radiation, \( I_b \) is defined by \( \sigma T^4/\pi \). Note that for situations in which the scattering coefficient is zero, the RTE reduces to a set of linear, decoupled equations for each intensity to be solved. Discrete Ordinate (DO) methods such as \( S_N \) have been developed to help solve this set of coupled equations by selecting a set of prescribed directions \( \Omega_k \), solving for the discrete intensities \( I_k \) and combining these intensities to represent the radiative field. These methods suffer from high computational cost due to the fact that a large number of ordinate directions is required to adequately resolve the intensity field and avoid “ray effects”. For a given quadrature order \( N \), the number of ordinate directions scales as \( N^2 \).

23.1.1 Simplified Spherical Harmonics (\( SP_N \)) Equations

The simplified spherical harmonics (\( SP_N \)) approximation to the radiative transport equation (RTE) was first proposed by Gelbard for reactor analysis in the early 1960s. The initial derivation involved replacing the spatial derivatives in the 1D spherical harmonics (\( P_N \)) approximation with their 3D analogs. It was later shown that a more rigorous theoretical basis was possible and that the \( SP_N \) equations may be derived as either an asymptotic correction to the diffusion limit or from the use of certain trial functions in the self-adjoint variational characterization of the even-parity form of the RTE. There are a number of equivalent
forms for the $SP_N$ equations in the literature. We choose to use the “canonical” form which is derived from
the 1D even-parity discrete ordinates equations.

$$- \nabla \cdot \left( \frac{\mu_n^2}{\sigma_T} \nabla I_n \right) + \sigma_T I_n = 4\pi \sigma_S \sum_{m=1}^{N+1} w_m I_m + \sigma_A I_b$$
(23.2)

where

• $\sigma_T$ is the extinction coefficient, $\sigma_T = \sigma_A + \sigma_S$
• $\mu_n$ is the nth quadrature point in a $(N+1)$-point Gauss set on $[-1, 1]$
• $I_n$ is the angular intensity at quadrature point $n$
• $w_n$ is the $n$th quadrature weight

The appropriate boundary conditions for the “canonical” $SP_n$ equations are derived from the boundary
conditions for the 1D even-parity discrete ordinates equations. We consider a Mark boundary condition of
the 1st order intensity BC

$$I_n = \epsilon I_b + (1 - \epsilon) \sum_{\vec{n} \cdot \vec{\Omega}_j < 0} I_j w_j \vec{n} \cdot \vec{\Omega}_j$$
(23.3)

such that in canonical form, it yields

$$- \frac{\mu_n}{\sigma_T} \nabla I_n \cdot \vec{n} = \frac{\epsilon}{2 - \epsilon} (I_n - I_b) + \frac{1 - \epsilon}{2 - \epsilon} \left[ \sum_k \frac{I_k - \frac{\mu_k}{\sigma_T} \nabla I_k \cdot \vec{n}}{\sum \mu_k w_k} \mu_k w_k - I_n + \frac{\mu_n}{\sigma_T} \nabla I_n \cdot \vec{n} \right]$$
(23.4)

where

• $\epsilon$ is the emissivity
• $\vec{n}$ is the surface normal

The $SP_N$ equations are solved for the unknown intensities to provide an approximate solution to the RTE.
The angle-integrated intensity is then found and used in the source term for the material energy equation.
It is to be noted from the form of the $SP_N$ equations that this cannot be used in vacuum or near-vacuum
situations where $\sigma_T 0$. Also analysis by Zheng et al shows that the existence of a unique solution may be
proved when absorption effects are non-negligible and the geometry is small.
Chapter 24

Contact Reference

24.1 Contact Overview

Contact is a term that is used when a simulation is to be performed on a nonconformal mesh. One example of a nonconformal mesh is one that has adjacent blocks and there appears to be a single node at a certain location in space on the interfaces between blocks, but in fact there are distinct nodes that have identical, or nearly identical coordinates. In other words, the mesh nodes have not been merged or equivalenced. More generally, a nonconformal mesh does not require that mesh lines match at the interface between block boundaries.

There are two modeling situations that lead to the use of non conformal meshes in thermal analysis. The first occurs when compatibility conditions are enforced at the interface to require that the temperature be continuous across the interface. This situation arises as a convenient way to generate the mesh for particularly complex geometries, or because a person wants to have a particular part be meshed independently for the convenience of changing its orientation, etc. The second modeling situation that occurs is not a matter of convenience but of physics. It may be that two parts are known to fit imperfectly, so that there is a gap between the two parts that has a certain known resistance. In this case, the enforcement is known as gap conductance (or resistance) and enforces a discontinuous solution across the gap by modeling the energy flux that is "lost" across the gap.

There are three C0 enforcement strategies currently in Aria:

1. Multipoint constraints (MPC), which are implemented using the ACME search. This is a first order accurate implementation that makes the slave surface the interpolant of the master surface. This method is first order accurate and is not conservative.

2. Generalized Contact, which is implemented using the ACME search. This method uses a discontinuous Galerkin approach to write two surface integrals along the contact interface. One integral penalizes the difference in the temperature, the other penalizes the difference in the fluxes, and so it attempts to match both the temperature values across the interface, as well as the flux. If the mesh aligns perfectly at the interface so that the quadrature points of the two facets match, then this method can be shown to be conservative. This method has the same order of accuracy as the interior discretization.

3. Dash Contact, which is implemented with the Dash search. This method uses a surface integral that penalizes the difference in the temperatures across the interface. It can be viewed as a specialization of Generalized Contact that drops the penalty term on the fluxes and has a specific integration rule. This integration rule uses a one-point quadrature rule on the subtriangle areas returned by Dash. This method is not conservative and is first order accurate.

The user supplied information to define contact is divided into two parts: surface specification and enforcement. Surfaces can either be specified as Exodus sidesets or blocks, or they can be automatically found via a block skinning approach. Enforcement is the strategy that is used to model the conditions
at the interface. The generalized contact algorithm, denoted as tied in the enforcement, are written to mathematically model both continuity of temperature and resistance contact simultaneously. The modeling of contact resistance is denoted in the enforcement as gap_conductance and is not currently available when using Dash contact.

### 24.2 Contact Definition

**Scope:** Equation System

```plaintext
Begin Contact Definition Contact_name
    Contact Surface Surface_name Contains List_of_instances...
    Disable Opposed Normal Interaction Culling
    Initial Overlap Removal {=|are|is} Option
    Multiple Interactions {=|are|is} Option [List_of_instances]...
    Opposed Normal Threshold Angle {=|are|is} Angle
    Output Rule {=|are|is} OutputRule
    Reinitialize Contact On Restart {=|are|is} Option
    Search {=|are|is} Type
    Update Search Every n Steps
    Visualize Contact
    Begin Contact Surface Contact_surface_name End
    Begin Debug Debug_name End
    Begin Developer Command: Enclosure Debug_name End
    Begin Enforcement Enforcement_name End
    Begin Interaction Interaction_name End
    Begin Interaction Defaults End
    Begin Nonconformal NonConformal_name End
    Begin Search Options Search_options_name End
    Begin Surface Options Shell_lofting_name End
End```

**Summary**
Contains the commands needed to define contact for an analysis.

**Description**
This block command defines a scenario for the contact model. The contact surfaces are defined, the interactions are set, and the type of contact enforcement is defined.
24.2.1 Contact Surface

Scope: Contact Definition

Contact Surface Surface_name Contains List_of_instances...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surface_name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>List_of_instances</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary

Defines a surface made up of a set of surfaces in the mesh file to consider for contact. Specification of an element block implies that block skinning of the element block is to take place. The specific implementation of block skinning depends on the application.

24.2.2 Disable Opposed Normal Interaction Culling

Scope: Contact Definition

Summary

This line command disables the culling of interactions that do not have opposed normals.

Description

This is a temporary developer command

24.2.3 Initial Overlap Removal

Scope: Contact Definition

Summary

Turns on or off the computation of contact variables. Turning off contact variables may provide a performance increase.

24.2.4 Multiple Interactions

Scope: Contact Definition

Multiple Interactions {=|are|is} Option [List_of_instances]...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>List_of_instances</td>
<td>[string]...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary

This command turns the processing of multiple interactions per node on or off.

24.2.5 Opposed Normal Threshold Angle

Scope: Contact Definition

Opposed Normal Threshold Angle {=|are|is} Angle

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Angle</td>
<td>real</td>
<td>90.0</td>
</tr>
</tbody>
</table>
Summary: Assign the angle, in degrees, for which surface normals are considered to be opposed.

Description: Prescribed angle must be greater or equal to 90 degrees.
Warning: This option is not recommended for general use but serves as a temporary patch that can be used for some problems.

### 24.2.6 Output Rule
Scope: Contact Definition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>OutputRule</td>
<td>{more verbose</td>
<td>none</td>
</tr>
</tbody>
</table>

Summary: Toggle amount of information output from the search.

### 24.2.7 Reinitialize Contact On Restart
Scope: Contact Definition

Summary: ***DEVELOPER ONLY COMMAND*** This flag indicates if contact data stored on restart should be read in on restart or if we should just reinitialize from scratch.

### 24.2.8 Search
Scope: Contact Definition

Summary: Specify contact search algorithm. ACME-use acme node/face contact library DASH-use dash face/face contact methods inside of Presto. ARS-use ars node/face contact methods inside Adagio/Presto.

### 24.2.9 Update Search Every
Scope: Contact Definition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Force an updating of the contact search every N steps.

Description: This command is useful if, for example, Aria is coupled to Adagio and the likelihood is strong that the mesh coordinates have changed significantly. Because contact surfaces may also appear in the definition of boundary conditions the application must selectively apply contact relations and boundary conditions appropriately. These issues are resolved automatically within the application code using the contact surface definitions and internal data masks.
24.2.10 Visualize Contact
Scope: Contact Definition

Summary Output contact surfaces with facets marked with 1 where contact is applied and 0 otherwise.

Description Enables output of surfaces where contact is active. Surface facets will be marked by whether contact constraints are applied on the facet - 1 or 0 otherwise.
Note: Visualization is enabled by default for contact defined using surface sidesets.
Warning: Use of this option is primarily for debugging purposes as the Results Output and History file will contain a skinned version of the contact element blocks. This may constitute a large amount of information for discontiguously meshed models.

24.3 Enforcement
Scope: Contact Definition

```
Begin Enforcement Enforcement_name
  Conductance Coefficient {=|are|is} Coeff
  Conductance Coefficient Contact Pressure Function {=|are|is} FName Using VarName
  Conductance Coefficient For EquationName {=|are|is} Model [ ModelParams... ]
  Conductance Coefficient Fortran Subroutine {=|are|is} FName
  Conductance Coefficient Subroutine {=|are|is} FName
  Conductance Coefficient Temperature Function {=|are|is} FName
  Conductance Coefficient Voltage Function {=|are|is} FName
  Dash Penalty Factor {=|are|is} value
  Enforcement For EquationName {=|are|is} EnforcementModel [ ModelParams... ]
  Integer Data Values...
  Real Data Values...
  Slave Variable Slave_field To Master_field
  Use Toggle Block ToggleName [ {@|at|for|in|on|over} ElementBlockList... ]
End
```

Summary Contains the commands needed to define contact enforcement for an analysis in Aria.

Description This block command is used to define the type of contact between the surfaces. The name of the enforcement is specified by the user. When the contact includes some contact resistance, the conductance coefficient for the contact between the two surfaces must be specified.
24.3.1 Conductance Coefficient

**Scope:** Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coeff</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** Set the constant valued contact conductance coefficient used by the GAP_CONDUCTANCE and CONDUCTANCE contact models.

**Description:** This line command defines the value of the effective heat-transfer coefficient, $h_c$, for the interface condition between the two surfaces. The temperature drop across the interface, due to the contact resistance, is modeled using

$$ q_i n_i|_{\Gamma_m} = h_c (T_{\Gamma_m} - T_{\Gamma_{m2}}) = q_i n_i|_{\Gamma_{m2}} $$

where the contact coefficient or contact conductance, $h_c(x_i, t, T_{\Gamma_m}, T_{\Gamma_{m2}})$, is modeled using a correlation that is dependent upon the two surface temperatures, location, and interface conditions. Note that smaller values of CONDUCTANCE COEFFICIENT correspond to resistance contact while larger values tend toward tied contact.

24.3.2 Conductance Coefficient Contact Pressure Function

**Scope:** Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>FName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>VarName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** Gap conductance coefficient from a user function in VarName

**Description:** See CONDUCTANCE COEFFICIENT line command.

24.3.3 Conductance Coefficient For

**Scope:** Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>EquationName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Model</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** Choose the model for the conductance coefficient for the specified equation to be used in generalized contact.

**Description:** Choose the model for the conductance coefficient for the specified equation to be used in generalized contact.
24.3.4 Conductance Coefficient Fortran Subroutine

Scope: Enforcement

```
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>
```

Summary Specify the name of a FORTRAN user subroutine that will be used to calculate the conductance coefficient.

24.3.5 Conductance Coefficient Subroutine

Scope: Enforcement

```
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>FName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>
```

Summary Set the user defined contact conductance coefficient. See CONDUCTANCE COEFFICIENT line command for details.

Description The subroutine must satisfy the element signature. The subroutine is passed a pair of faces that are in contact, and is expected to populate a single value for the coefficient. The temperature at the current integration point and its projection onto the face across the interface is provided. The contact interface is processed twice—one from the left side and once from the right. **Note that to ensure conservation, the subroutine should calculate the same value of the gap conductance on both sides of the interface.**

24.3.6 Conductance Coefficient Temperature Function

Scope: Enforcement

```
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>FName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>
```

Summary Set the temperature dependent contact conductance coefficient.

Description See CONDUCTANCE COEFFICIENT line command.

24.3.7 Conductance Coefficient Time Function

Scope: Enforcement

```
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>FName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>
```
24.3.8 Conductance Coefficient Voltage Function

**Scope:** Enforcement

```
Conductance Coefficient Voltage Function {=|are|is} FName
```

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>FName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Set the temperature dependent contact conductance coefficient.

**Description**
See CONDUCTANCE COEFFICIENT line command.

24.3.9 Dash Penalty Factor

**Scope:** Enforcement

```
Dash Penalty Factor {=|are|is} value
```

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Defines a penalty factor to use for the dash constraints.

24.3.10 Enforcement For

**Scope:** Enforcement

```
Enforcement For EquationName {=|are|is} EnforcementModel [ ModelParams... ]
```

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>EquationName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>EnforcementModel</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Defines the model for enforcement for the command block interactions, either tied or resistance/conductance contact. Tied contact can be of type generalized or Lagrange multiplier/MPC, where generalized contact is the preferred contact algorithm. 24.4

**Description**
This command line sets the type of contact problem for interactions defined in the current CONTACT DEFINITION command block and the equation to which the enforcement applies. The Generalized contact enforcement model can be either GAP_CONDUCTANCE, CONDUCTANCE or TIED_DOF. In the case of resistance contact one must define a CONDUCTANCE COEFFICIENT model. To specify that TIED contact will be enforced for DOF my_DOF_name, the enforcement model will be TIED_my_DOF_name, where the solution DOF for EquationName is defined in the EQ command line.
Lagrange multiplier enforcement is specified simply as model TIED and must be accompanied by a SLAVE VARIABLE command line within the ENFORCEMENT command block. MPC contact requires that one supply MASTER and SLAVE surface definitions in the INTERACTION command block. Additionally one should consider using the MATRIX REDUCTION = FEI-REMOVE-SLAVES in the equation solver command block for MPC constraint reduction.

DASH contact enforcement will be enforced only in a TIED sense.

### 24.3.11 Integer Data

**Scope:** Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Values</td>
<td>integer...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  List of integer data values to be used by the FORTRAN user subroutine. Copies of these values are provided to the subroutine hence changes to these values within the subroutine are not saved.

### 24.3.12 Real Data

**Scope:** Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Values</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  List of real data values to be used by the FORTRAN user subroutine. Copies of these values are provided to the subroutine hence changes to these values within the subroutine are not saved.

### 24.3.13 Slave Variable

**Scope:** Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Slave_field</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Master_field</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  Defines the variables involved in enforcement for this interaction.

**Description**  This line command sets the master and slave variables involved in the current tied contact definition.
24.3.14 Use Toggle Block
Scope: Enforcement

Use Toggle Block *ToggleName* [ {@|at|for|in|on|over} *ElementBlockList...* ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>ToggleName</em></td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specification for toggling entities in the computational model based on Toggle Block parameters. When used at the region level, the list of element blocks to be toggled must be provided. Otherwise a listing of entities is not needed as the Toggle Block will be associated with the command line or the enclosing command block.

24.4 Enforcement Models

24.4.1 Enforcement For Continuity = Tied Ip Continuity
Scope: Enforcement

24.4.2 Enforcement For Current = Bulter Volmer
Scope: Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Toggle</em></td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td><em>F</em></td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td><em>R</em></td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td><em>Alpha_A</em></td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td><em>Alpha_C</em></td>
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<td>undefined</td>
</tr>
</tbody>
</table>

24.4.3 Enforcement For Current = Conductance
Scope: Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Toggle</em></td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Applies a contact current flux of the form \(C(V_1 - V_2)/2\) to sides 1 and 2 of the contact interface where \(C\) is a user specified conductance coefficient. Used in conjunction with a CONDUCTANCE COEFFICIENT model.

24.4.4 Enforcement For Current = Gap Conductance
Scope: Enforcement
Summary: Applies a contact current flux of the form \( \tilde{C}(V_1 - V_2)/2 \) to sides 1 and 2 of the contact interface. With \( C \), a user defined coefficient and \( \tilde{C} \), an average conductance coefficient, then \( \tilde{C} = C \) for \( C \ll \tilde{C} \) and \( \tilde{C} \) otherwise. Used in conjunction with a CONDUCTANCE COEFFICIENT model.

### 24.4.5 Enforcement For Current = Tied Voltage

**Scope:** Enforcement

Summary: Applies a contact current flux of the form \((\sigma_1 \nabla V_1 \cdot n_1 + \sigma_2 \nabla V_2 \cdot n_2 + \tilde{C}(V_1 - V_2))/2\) to sides 1 and 2 of the contact interface where \( \sigma \) is the electrical conductivity and \( \tilde{C} \) is an average conductance coefficient.

### 24.4.6 Enforcement For Cvfem Continuity = Pressure Jump

**Scope:** Enforcement

### 24.4.7 Enforcement For Cvfem Continuity = Tied Continuity

**Scope:** Enforcement

### 24.4.8 Enforcement For Cvfem Energy = Conductance

**Scope:** Enforcement

Summary: Applies a contact heat flux of the form \( C(T_1 - T_2)/2 \) to sides 1 and 2 of the contact interface where \( C \) is a user specified conductance coefficient. Used in conjunction with a CONDUCTANCE COEFFICIENT model.
24.4.9 Enforcement For Cvfem Energy = Gap Conductance
Scope: Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>toggle</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Applies a contact heat flux of the form $\tilde{h}(T_1 - T_2)/2$ to sides 1 and 2 of the contact interface. With $h$, the user defined coefficient and $\bar{h}$, an average conductance coefficient, then $\tilde{h} = h$ for $h \ll \bar{h}$ and $\bar{h}$ otherwise. Used in conjunction with a CONDUCTANCE COEFFICIENT model.

24.4.10 Enforcement For Cvfem Energy = Tied Temperature
Scope: Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>toggle</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Applies a contact heat flux of the form $(k_1 \nabla T_1 \cdot n_2 + k_2 \nabla T_2 \cdot n_2 + \bar{h}(T_1 - T_2))/2$ to sides 1 and 2 of the contact interface where $k$ is the thermal conductivity and $\bar{h}$ is an average conductance coefficient.

24.4.11 Enforcement For Cvfem Lumped Projection = Tied Gradp
Scope: Enforcement

24.4.12 Enforcement For Cvfem Momentum = Tied Momentum
Scope: Enforcement

24.4.13 Enforcement For Cvfem Projection = Tied Gradp
Scope: Enforcement

24.4.14 Enforcement For Cvfem Specific Dissipation Rate = Tied Sdr
Scope: Enforcement

Summary: Tied_Sdr contact flux; CVFEM pressure stabilization included
24.4.15 Enforcement For Cvfem Turbulent Kinetic Energy = Tied Tke
Scope: Enforcement

Summary Tied_Tke contact flux; CVFEM pressure stabilization included

24.4.16 Enforcement For Energy = Cht Robin
Scope: Enforcement

Enforcement For Energy = Cht Robin [ Toggle = toggle |Opposite_Material_Phase = opposite_material_phase |Opposite_Flux_Name = opposite_flux_name |Coeff_Scaling = coeff_scaling ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
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<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>opposite_material_phase</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>opposite_flux_name</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>coeff_scaling</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Applies a Robin-style BC for the energy equation: \( F_{\text{local}} = F_{bc} + \alpha (T_{\text{local}} - T_{bc}) \).

24.4.17 Enforcement For Energy = Conductance
Scope: Enforcement

Enforcement For Energy = Conductance [ Toggle = toggle ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>toggle</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Applies a contact heat flux of the form \( C(T_1 - T_2)/2 \) to sides 1 and 2 of the contact interface where \( C \) is a user specified conductance coefficient. Used in conjunction with a CONDUCTANCE COEFFICIENT model.

24.4.18 Enforcement For Energy = Conductance Salgon
Scope: Enforcement

Enforcement For Energy = Conductance Salgon [ Toggle = toggle |A = a |B = b |Xi = xi ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>toggle</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
<tr>
<td>a</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>b</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>xi</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Applies a contact heat flux of the form \((2ka)/(\pi b^2(1 - 1.14\xi))(T_1 - T_2)\) to sides 1 and 2 of the contact interface where \( k \) is the thermal conductivity, \( \hat{k} = (k_1 + k_2)/(k_1k_2) \) and \( \xi \) is a user specified parameter (Salgon,1997).
24.4.19 Enforcement For Energy = Contact Resistance

Scope: Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>toggle</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Applies a contact heat flux of the form \( h(T - T_{pp}) \) where \( h \) is the gap conductance coefficient defined for the surface material and the material phase of \( T_{pp} \) is optionally specified.

24.4.20 Enforcement For Energy = Gap Conductance

Scope: Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>toggle</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Applies a contact heat flux of the form \( \bar{h}(T_1 - T_2)/2 \) to sides 1 and 2 of the contact interface. With \( h \), a user defined coefficient and \( \bar{h} \), an average conductance coefficient, then \( \bar{h} = h \) for \( h \ll \bar{h} \) and \( \bar{h} \) otherwise. Used in conjunction with a CONDUCTANCE COEFFICIENT model.

24.4.21 Enforcement For Energy = Phase Change

Scope: Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>toggle</td>
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</tr>
<tr>
<td>wrt</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Applies an energy flux to an interface based on phase change, as \( q = \rho v (T_s C_{p,s} - T_g C_{p,g} + L) \) to sides \( s \) and \( g \) of the contact interface where \( v \) is the velocity, \( \rho \) is the interface density, \( C_p \) is the specific heat and \( L \) is the latent heat of fusion.

24.4.22 Enforcement For Energy = Tied D Style Temperature

Scope: Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
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</tr>
<tr>
<td>lambda_scaling</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Summary

Applies a contact heat flux of the form \( \lambda \bar{h}(T_1 - T_2)/2 \) to sides 1 and 2 of the contact interface where \( \lambda \) is a penalty parameter, \( \bar{h} \) is an average conductance coefficient. This model is similar to the DASH_TIED model.

24.4.23 Enforcement For Energy = Tied Ip Temperature

Scope: Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>toggle</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary

Applies a contact heat flux of the form \( (k_1 \nabla T_1 \cdot n_1 + k_2 \nabla T_2 \cdot n_2 + \bar{h}(T_1 - T_2))/2 \) to sides 1 and 2 of the contact interface where \( k \) is the thermal conductivity and \( \bar{h} \) is an average conductance coefficient. Additionally an interior penalty jump term is also applied.

24.4.24 Enforcement For Energy = Tied Robin Style Temperature

Scope: Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>toggle</td>
<td>&quot;string&quot;</td>
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</tr>
</tbody>
</table>

Summary

Applies a contact heat flux of the form \( (k_2 \nabla T_2 \cdot n_2 + \bar{h}(T_1 - T_2))/2 \) to sides 1 and 2 of the contact interface where \( k \) is the thermal conductivity and \( \bar{h} \) is an average conductance coefficient.

24.4.25 Enforcement For Energy = Tied Temperature

Scope: Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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<tbody>
<tr>
<td>toggle</td>
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</tbody>
</table>

Summary

Applies a contact heat flux of the form \( (k_1 \nabla T_1 \cdot n_1 + k_2 \nabla T_2 \cdot n_2 + \bar{h}(T_1 - T_2))/2 \) to sides 1 and 2 of the contact interface where \( k \) is the thermal conductivity and \( \bar{h} \) is an average conductance coefficient.

24.4.26 Enforcement For Hfem Continuity = Tied Ip Continuity

Scope: Enforcement
24.4.27 Enforcement For Hfem Momentum = Tied Ip Momentum
Scope: Enforcement

24.4.28 Enforcement For Hfem Momentum = Tied Momentum
Scope: Enforcement

24.4.29 Enforcement For Lubrication Height = Deforming
Scope: Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$h_{min}$</td>
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</tr>
</tbody>
</table>

Summary Lubrication height that deforms by the displacement of the adjoining volume region.

24.4.30 Enforcement For Mass Balance = Bulk Node Mass Open
Scope: Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk_Node</td>
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</tr>
</tbody>
</table>

Summary Open BC between a porous medium and a bulk node

24.4.31 Enforcement For Mass Balance = Bulk Node Open
Scope: Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk_Node</td>
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</tr>
</tbody>
</table>

24.4.32 Enforcement For Mass Balance = Bulk Node Species Open
Scope: Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk_Node</td>
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<td>undefined</td>
</tr>
</tbody>
</table>
24.4.33 Enforcement For Mass Balance = Conserved Mass
Scope: Enforcement

Summary Continuous mass balance and capillary pressure, discontinuous nonwetting phase saturation.

24.4.34 Enforcement For Mesh = Lubrication Pressure
Scope: Enforcement

Summary Lubrication pressure mesh flux

24.4.35 Enforcement For Momentum = Tied Ip Momentum
Scope: Enforcement

24.4.36 Enforcement For Momentum = Tied Momentum
Scope: Enforcement

24.4.37 Enforcement For Porous Enthalpy = Bulk Node Open
Scope: Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>bulk_node</td>
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</tr>
</tbody>
</table>

24.4.38 Enforcement For Porous Enthalpy = Bulk Node Open Uncoupled
Scope: Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>bulk_node</td>
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</tr>
</tbody>
</table>

24.4.39 Enforcement For Porous Enthalpy = Contact Resistance
Scope: Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>opposite_material_phase</td>
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</tr>
<tr>
<td>bulk_node</td>
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<td>undefined</td>
</tr>
</tbody>
</table>
Summary

Applies a heat flux of the form \( J \cdot n = h(T - T_{opp}) \) where \( h \) is the gap conductance coefficient defined for the surface material and the material phase of \( T_{opp} \) is optionally specified.

24.4.40 Enforcement For Porous Species = Bulk Node Open All Inflow
Scope: Enforcement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>bulk_node</td>
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</tr>
</tbody>
</table>

24.5 Interaction
Scope: Contact Definition

Summary
Contains the commands needed to define a surface-surface interaction.

24.5.1 Master
Scope: Interaction

Summary
Defines a list of surfaces that are master to each surface in the slave list

24.5.2 Normal Tolerance
Scope: Interaction

Summary

Normal Tolerance \( \{=|are|is\} Distance \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Distance</td>
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<td>undefined</td>
</tr>
</tbody>
</table>
Summary  Set distance of normal tolerance.

24.5.3 Overlap Normal Tolerance
Scope: Interaction

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Overlap Normal Tolerance (=</td>
<td>)are(=</td>
<td>)is (Distance)</td>
</tr>
<tr>
<td>Distance</td>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary  defines distance allowed for normal tolerance of overlap

24.5.4 Overlap Tangential Tolerance
Scope: Interaction

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Overlap Tangential Tolerance (=</td>
<td>)are(=</td>
<td>)is (Distance)</td>
</tr>
<tr>
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<td>undefined</td>
</tr>
</tbody>
</table>

Summary  defines distance allowed for tangential tolerance of overlap

24.5.5 Slave
Scope: Interaction

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Slave (=</td>
<td>)are(=</td>
<td>)is (Slave)...</td>
</tr>
<tr>
<td>Slave</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Defines a list of surfaces that are slave to each surface in the master list

24.5.6 Surfaces
Scope: Interaction

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
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</thead>
<tbody>
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<td>)are(=</td>
<td>)is (Surfaces)...</td>
</tr>
<tr>
<td>Surfaces</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Defines a pairwise set of surfaces for which to turn on interactions

24.5.7 Tangential Tolerance
Scope: Interaction
Tangential Tolerance `{]|are|is}` *Distance*

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Distance</em></td>
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<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Set distance of tangential tolerance
Chapter 25

Output Reference

25.1 Output Overview

Sierra Framework services provide various controls for output data. Output is divided into three major categories, Results output, Heartbeat output and Restart output. Results output contains binary format simulation results in a form suitable for visualization. While the Results output usually corresponds to the entire model, it can also be applied to portions of the model. Heartbeat output is generally written to a text file and provides a convenient means of monitoring intermediate simulation results. Restart output (Restart Data) is specialized for use in analyses that may not complete in a single job submission but might complete when continued from a previous termination time.

When using Results output or Heartbeat output one should note that the output request must be made using the internal Field name 2.9, most frequently solution->field_name or pp->postprocess_name. If the internal Field name is not used in these requests output cannot be generated for that Field and a warning to that effect will be written to the log file. Global variables may also be output to the Results and Heartbeat output files. A list of all fields and global variables available for output in a particular model is output to the log file immediately after the mesh is read. This list can be previewed by running aria with the –check-input option that will load verify the input syntax, read the mesh, print the variables available for output, and then exit.

Depending upon the application one may wish to define multiple Results output or Heartbeat output command blocks. In doing so one must recognize that output time or output interval specifications are cumulative over the various output requests. Note that because time stepping accommodates simulation output requests, inconsistency of output frequency specification among multiple Results output blocks can produce unexpected/undesirable behavior in the solution time stepping.

Commands pertinent to the definition of Restart are included in this section. These commands describe management of restart files and content of the files. The intent of restart is to begin a new sequence of calculations using the entire content of the restart file. Those wishing to restart a simulation using only part of the results may wish to consider using the IC READ_FILE option 8.7. The invocation of restart within a simulation (specifying when it should start) is outlined elsewhere in this document 3.10.

Within the input file Output command blocks will appear at the Region scope as illustrated below.
25.2 Results Output

Scope: Encore Region

Begin Results Output Label
  Additional Steps {=|are|is} List_of_steps...
  Additional Times {=|are|is} List_of_times...
  At Step n Option {=|are|is} m
  At Time Dt1 Option {=|are|is} Dt2
  Component Separator Character {=|are|is} Separator
  Database Name {=|are|is} StreamName
  Database Type {=|are|is} DatabaseType
  Edge [ VariableList... ]
  Edge Variables {=|are|is} [ VariableList... ]
  Element [ VariableList... ]
  Element Variables {=|are|is} [ VariableList... ]
  Exclude {=|are|is} [ ElementBlockList... ]
  Exists Option1 Option2
  Face [ VariableList... ]
  Face Variables {=|are|is} [ VariableList... ]
  Global [ Variables... ]
  Global Variables {=|are|is} [ Variables... ]
  Include {=|are|is} [ ElementBlockList... ]
  Nodal [ VariableList... ]
Summary  
Describes the location and type of the output stream used for outputting results for the enclosing region.

### 25.2.1 Additional Steps

**Scope:** Results Output

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>List_of_steps</td>
<td>integer...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  
Additional simulation steps when output should occur.

### 25.2.2 Additional Times

**Scope:** Results Output

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>List_of_times</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  
Additional simulation times when output should occur.
25.2.3  **At Step**  
**Scope:** Results Output

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( n )</td>
<td>integer</td>
<td>undefined</td>
</tr>
<tr>
<td>( m )</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  
Specify an output interval in terms of the internal iteration step count. The first step specifies the step count at the beginning of this interval and the second step specifies the output frequency to be used within this interval.

25.2.4  **At Time**  
**Scope:** Results Output

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( Dt1 )</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>( Dt2 )</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  
Specify an output interval in terms of the internal simulation time. The first time specifies the time at the beginning of this interval and the second time specifies the output frequency to be used within this interval.

25.2.5  **Component Separator Character**  
**Scope:** Results Output

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( Separator )</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  
The separator is the single character used to separate the output variable basename (e.g. "stress") from the suffixes (e.g. "xx", "yy") when displaying the names of the individual variable components. For example, the default separator is ",", which results in names similar to "stress_xx", "stress_yy", ... "stress_zx". To eliminate the separator, specify an empty string (""") or NONE.

25.2.6  **Database Name**  
**Scope:** Results Output

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( StreamName )</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Summary: The base name of the database containing the output results. If the filename begins with the '/' character, it is an absolute path; otherwise, the path to the current directory will be prepended to the name. If this line is omitted, then a filename will be created from the basename of the input file with a ".e" suffix appended.

25.2.7 Database Type
Scope: Results Output

Summary: The database type/format to be used for the output results.

25.2.8 Edge
Scope: Results Output

Summary: Define the variables that should be written to the results database. If "variable" is entered, then its name will be used on the output database. If "variable as db_name" is entered, then "db_name" will be the name used on the database for the internal variable "variable". Multiple "variable" or "variable as db_name" entries are allowed on the same line. The entities that this variable are written to can also be limited or specified with "exclude list_of_entities" or "include list_of_entities". Edge variables are not supported for all database types.

25.2.9 Edge Variables
Scope: Results Output

Summary: Define the variables that should be written to the results database. If "variable" is entered, then its name will be used on the output database. If "variable as db_name" is entered, then "db_name" will be the name used on the database for the internal variable "variable". Multiple "variable" or "variable as db_name" entries are allowed on the same line. The entities that this variable are written to can also be limited or specified with "exclude list_of_entities" or "include list_of_entities". Edge variables are not supported for all database types.

25.2.10 Element
Scope: Results Output

Summary: Define the variables that should be written to the results database. If "variable" is entered, then its name will be used on the output database. If "variable as db_name" is entered, then "db_name" will be the name used on the database for the internal variable "variable". Multiple "variable" or "variable as db_name" entries are allowed on the same line. The entities that this variable are written to can also be limited or specified with "exclude list_of_entities" or "include list_of_entities"
25.2.11 Element Variables
Scope: Results Output

Summary Define the variables that should be written to the results database. If "variable" is entered, then its name will be used on the output database. If "variable as db_name" is entered, then "db_name" will be the name used on the database for the internal variable "variable". Multiple "variable" or "variable as db_name" entries are allowed on the same line. The entities that this variable are written to can also be limited or specified with "exclude list_of_entities" or "include list_of_entities"

25.2.12 Exclude
Scope: Results Output

Summary Specify that the results file will only contain a subset of the element blocks in the analysis model. The element_block_list lists only the blocks which will not be output to the results database.

25.2.13 Exists
Scope: Results Output

Summary Specify the behavior when creating this database and there is an existing file with the same name. The default behavior is "OVERWRITE" which deletes the existing file and creates a new file of the same name. "APPEND" will (if possible) append the new data to the end of the existing file. "ABORT" will print an error message and end the analysis. "ADD_SUFFIX" will add a -s???? suffix where the ???? is replaced by a sequential number starting at 0002.

25.2.14 Face
Scope: Results Output

Summary Define the variables that should be written to the results database. If "variable" is entered, then its name will be used on the output database. If "variable as db_name" is entered, then "db_name" will be the name used on the database for the internal variable "variable". Multiple "variable" or "variable as db_name" entries are allowed on the same line. The entities that this variable are written to can also be limited or specified with "exclude list_of_entities" or "include list_of_entities". Face variables are not supported for all database types.

25.2.15 Face Variables
Scope: Results Output

Summary Define the variables that should be written to the results database. If "variable" is entered, then its name will be used on the output database. If "variable as db_name" is entered, then
"db_name" will be the name used on the database for the internal variable "variable". Multiple "variable" or "variable as db_name" entries are allowed on the same line. The entities that this variable are written to can also be limited or specified with "exclude list_of_entities" or "include list_of_entities". Face variables are not supported for all database types.

25.2.16 Global
Scope: Results Output

Summary Define the global variables that should be written to the results database. If "variable" is entered, then its name will be used on the output database. If "variable as db_name" is entered, then "db_name" will be the name used on the database for the internal variable "variable". Multiple "variable" or "variable as db_name" entries are allowed on the same line.

25.2.17 Global Variables
Scope: Results Output

Summary Define the global variables that should be written to the results database. If "variable" is entered, then its name will be used on the output database. If "variable as db_name" is entered, then "db_name" will be the name used on the database for the internal variable "variable". Multiple "variable" or "variable as db_name" entries are allowed on the same line.

25.2.18 Include
Scope: Results Output

Summary Specify that the results file will only contain a subset of the element blocks in the analysis model. The element_block_list lists only the blocks which will be output to the results database.

25.2.19 Nodal
Scope: Results Output

Summary Define the nodal variables that should be written to the results database. If "variable" is entered, then its name will be used on the output database. If "variable as db_name" is entered, then "db_name" will be the name used on the database for the internal variable "variable". Multiple "variable" or "variable as db_name" entries are allowed on the same line.

25.2.20 Nodal Variables
Scope: Results Output

749
25.2.21 Node
Scope: Results Output

Define the nodal variables that should be written to the results database. If "variable" is entered, then its name will be used on the output database. If "variable as db_name" is entered, then "db_name" will be the name used on the database for the internal variable "variable". Multiple "variable" or "variable as db_name" entries are allowed on the same line.

25.2.22 Node Variables
Scope: Results Output

Define the nodal variables that should be written to the results database. If "variable" is entered, then its name will be used on the output database. If "variable as db_name" is entered, then "db_name" will be the name used on the database for the internal variable "variable". Multiple "variable" or "variable as db_name" entries are allowed on the same line.

25.2.23 Nodeset
Scope: Results Output

Define the variables that should be written to the results database. If "variable" is entered, then its name will be used on the output database. If "variable as db_name" is entered, then "db_name" will be the name used on the database for the internal variable "variable". Multiple "variable" or "variable as db_name" entries are allowed on the same line. The entities that this variable are written to can also be limited or specified with "exclude list_of_entities" or "include list_of_entities". Nodeset variables are not supported for all database types.

25.2.24 Nodeset Variables
Scope: Results Output

Define the variables that should be written to the results database. If "variable" is entered, then its name will be used on the output database. If "variable as db_name" is entered, then "db_name" will be the name used on the database for the internal variable "variable". Multiple "variable" or "variable as db_name" entries are allowed on the same line. The entities that this variable are written to can also be limited or specified with "exclude
list_of_entities" or "include list_of_entities". Nodeset variables are not supported for all database types.

25.2.25 Output Mesh
Scope: Results Output

Output Mesh {=|are|is} OutputMesh

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>OutputMesh</td>
<td>{exposed surface</td>
<td>refined</td>
</tr>
</tbody>
</table>

Summary: Use this command to turn on "unrefined" as the output mesh. The default behavior is "refined", in which field variables are output on the current mesh, which may have been refined (either uniformly or adaptively) or had its topology altered in some way (e.g., dynamic load balancing) with respect to the original mesh read from the the input file. By specifying "Output Mesh = unrefined", all output variables are output only on the original mesh objects read from the input file.

25.2.26 Output On Signal
Scope: Results Output

Output On Signal {=|are|is} Signals

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Signals</td>
<td>{sigabrt</td>
<td>sigalrm</td>
</tr>
</tbody>
</table>

Summary: When the specified signal is raised, the output stream associated with this block will be output.

25.2.27 Overwrite
Scope: Results Output

Summary: (DEPRECATED, Use EXISTS) Specify whether the database should be overwritten if it exists. The default behavior is to overwrite unless this command is specified in the output block and either off, false, or no is specified.

25.2.28 Property
Scope: Results Output

Property PropertyName {=|are|is} PropertyValue

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>PropertyName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>PropertyValue</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Define a database property named "PropertyName" with the value "PropertyValue". If PropertyValue consists of all digits, it will define an integer property. If PropertyValue is "true" or "yes" or "false" or "no", it will define a logical property; otherwise it will define a string property. Supported properties are typically database dependent; Current properties are: COMPRESS_LEVEL = [0..9] (off) COMPRESS本朴 SHUFFLE = true|false|on|off (off) FILE_TYPE = netcdf4 (forces use of netcdf-4 hdf5-based file) (netcdf3)
INTEGER_SIZE_DB = 4|8 (4) INTEGER_SIZE_API = 4|8 (4) REAL_SIZE_DB = 4|8 (8 is default) LOGGING = true|false|on|off (off) MAX_NAME_LENGTH = value (32)

25.2.29 Sideset
Scope: Results Output

Define the variables that should be written to the results database. If "variable" is entered, then its name will be used on the output database. If "variable as db_name" is entered, then "db_name" will be the name used on the database for the internal variable "variable". Multiple "variable" or "variable as db_name" entries are allowed on the same line. The entities that this variable are written to can also be limited or specified with "exclude list_of_entities" or "include list_of_entities". Face variables are not supported for all database types.

25.2.30 Sideset Variables
Scope: Results Output

Define the variables that should be written to the results database. If "variable" is entered, then its name will be used on the output database. If "variable as db_name" is entered, then "db_name" will be the name used on the database for the internal variable "variable". Multiple "variable" or "variable as db_name" entries are allowed on the same line. The entities that this variable are written to can also be limited or specified with "exclude list_of_entities" or "include list_of_entities". Face variables are not supported for all database types.

25.2.31 Start Time
Scope: Results Output

Start Time {=} real Start_time

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Start_time</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Specify the time to start outputting results from this output request block. This time overrides all 'at time' and 'at step' specifications.

25.2.32 Surface
Scope: Results Output
25.2.33 Surface Variables

Scope: Results Output

Summary
Define the variables that should be written to the results database. If "variable" is entered, then its name will be used on the output database. If "variable as db_name" is entered, then "db_name" will be the name used on the database for the internal variable "variable". Multiple "variable" or "variable as db_name" entries are allowed on the same line. The entities that this variable are written to can also be limited or specified with "exclude list_of_entities" or "include list_of_entities". Face variables are not supported for all database types.

25.2.34 Synchronize Output

Scope: Results Output

Summary
In an analysis with multiple regions, it is sometimes desirable to synchronize the output of results data between the regions. This can be done by adding the SYNCHRONIZE OUTPUT command line to the results output block. If a results block has this set, then it will write output whenever a previous region writes output. The ordering of regions is based on the order in the input file, algorithmic considerations, or by solution control specifications.

Although the USE OUTPUT SCHEDULER command line can also synchronize output between regions, the SYNCHRONIZE OUTPUT command line will synchronize the output with regions where the output frequency is not under the direct control of the Sierra IO system. Examples of this are typically coupled applications where one or more of the codes are not Sierra-based applications such as Alegra and CTH. A results block with SYNCHRONIZE OUTPUT specified will also synchronize its output with the output of the external code.

The SYNCHRONIZE OUTPUT command can be used with other output scheduling commands such as time-based or step-based output specifications.

25.2.35 Termination Time

Scope: Results Output

Summary
Specify the time to stop outputting results from this output request block.

### Termination Time

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Final_time</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specify the time to stop outputting results from this output request block.
25.2.36 Timestep Adjustment Interval
Scope: Results Output

Timestep Adjustment Interval \( N_{\text{steps}} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( N_{\text{steps}} )</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Specify the number of steps to 'look ahead' and adjust the timestep to ensure that the specified output times or simulation end time will be hit 'exactly'.

25.2.37 Title
Scope: Results Output

Summary
Specify the title to be used for this specific output block.

25.2.38 Use Output Scheduler
Scope: Results Output

Use Output Scheduler \( \text{Timer}_\text{name} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \text{Timer}_\text{name} )</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Associates a predefined output scheduler with this output block (results, restart, heartbeat, or history).

25.3 Heartbeat
Scope: Encore Region

Begin Heartbeat \( \text{Label} \)

Additional Steps \( \text{List}_\text{of}_\text{steps} \)
Additional Times \( \text{List}_\text{of}_\text{times} \)
Append \( \text{Option} \)
At Step \( m \) \( \text{Option} \)
At Time \( \text{Dt}_1 \) \( \text{Option} \)
Edge \( \text{VariableList} \)
Element \( \text{VariableList} \)
Exists \( \text{Option}_1 \) \( \text{Option}_2 \)
Face \( \text{VariableList} \)
Format \( \text{StreamTypes} \)
Global [ Variables... ]
Labels {=|are|is} Option
Legend {=|are|is} Option
Monitor Equals Option
Nodal [ VariableList... ]
Node [ VariableList... ]
Nodeset [ VariableList... ]
Output On Signal {=|are|is} Signals
Precision {=|are|is} Precision
Start Time {=|are|is} Start_time
Stream Name {=|are|is} OutputFilename
Synchronize Output
Termination Time {=|are|is} Final_time
Timestamp Format
Timestep Adjustment Interval {=|are|is} Nsteps
Use Output Scheduler Timer_name
Variable {=|are|is} Option [ Variable_list... ]

Summary
Describes the location and type of the output stream used for outputting the heartbeat information for the enclosing region.

### 25.3.1 Additional Steps
Scope: Heartbeat

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>List_of_steps</td>
<td>integer...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Additional simulation steps when output should occur.

### 25.3.2 Additional Times
Scope: Heartbeat

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>List_of_times</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Additional simulation times when output should occur.
25.3.3 Append
Scope: Heartbeat

Summary Specifies whether the heartbeat file is appended if it exists. By default, the file is appended if restart is requested and not if restart is not requested. This option does not work for automatic restarts because a new heartbeat file is written with each auto restart.

25.3.4 At Step
Scope: Heartbeat

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>integer</td>
<td>undefined</td>
</tr>
<tr>
<td>m</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specify an output interval in terms of the internal iteration step count. The first step specifies the step count at the beginning of this interval and the second step specifies the output frequency to be used within this interval.

25.3.5 At Time
Scope: Heartbeat

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dt1</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>Dt2</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specify an output interval in terms of the internal simulation time. The first time specifies the time at the beginning of this interval and the second time specifies the output frequency to be used within this interval.

25.3.6 Edge
Scope: Heartbeat

Summary Define the edge variables that should be written to the heartbeat database. The syntax is:
"edge internal_name at edge id as DBname" or "edge internal_name nearest location X, Y, Z as DBname".
Where internal_name is the name of the variable in the Sierra application; and DBname is the name as it should appear on the heartbeat database.

25.3.7 Element
Scope: Heartbeat
Define the element variables that should be written to the heartbeat database. The syntax is: "element internal_name at element id as DBname" or "element internal_name nearest location X, Y, Z as DBname".

Where internal_name is the name of the variable in the Sierra application; and DBname is the name as it should appear on the heartbeat database.

25.3.8 Exists
Scope: Heartbeat

Specify the behavior when creating this database and there is an existing file with the same name. The default behavior is "OVERWRITE" which deletes the existing file and creates a new file of the same name. "APPEND" will (if possible) append the new data to the end of the existing file. "ABORT" will print an error message and end the analysis.

25.3.9 Face
Scope: Heartbeat

Define the face variables that should be written to the heartbeat database. The syntax is: "face internal_name at face id as DBname" or "face internal_name nearest location X, Y, Z as DBname".

Where internal_name is the name of the variable in the Sierra application; and DBname is the name as it should appear on the heartbeat database.

25.3.10 Format
Scope: Heartbeat

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>StreamTypes</td>
<td>{default</td>
<td>spyhis}</td>
</tr>
</tbody>
</table>

The stream type/format to be used for the output results. The only two options at this time are 'Default' which is the normal Sierra heartbeat format; and 'SpyHis' which mimics the CTH Spyhis history output format.

25.3.11 Global
Scope: Heartbeat

Define the global/reduction variables that should be written to the heartbeat database. The syntax is: "global internal_name as DBname".

Where internal_name is the name of the variable in the Sierra application; and DBname is the name as it should appear on the heartbeat database.
25.3.12 Labels
Scope: Heartbeat

Summary Specifies whether labels will be displayed or just the value of the variable. Labels will be shown if this line is not present.

25.3.13 Legend
Scope: Heartbeat

Summary Specifies whether a legend will be displayed prior to outputting any variables. The legend will not be shown unless this line is present. The legend shows the names of the variables that will be written to the heartbeat output stream. If the variable has multiple components, then the component count is shown after the variable e.g., velocity(3).

25.3.14 Monitor
Scope: Heartbeat

Summary Specifies whether a line will be written to the heartbeat stream when either the results, history, and/or restart data are output.

25.3.15 Nodal
Scope: Heartbeat

Summary Define the nodal variables that should be written to the heartbeat database. The syntax is: "nodal internal_name at node id as DBname" or "nodal internal_name nearest location X, Y, Z as DBname".

Where internal_name is the name of the variable in the Sierra application; and DBname is the name as it should appear on the heartbeat database.

25.3.16 Node
Scope: Heartbeat

Summary Define the nodal variables that should be written to the heartbeat database. The syntax is: "node internal_name at node id as DBname" or "node internal_name nearest location X, Y, Z as DBname".

Where internal_name is the name of the variable in the Sierra application; and DBname is the name as it should appear on the heartbeat database.
### 25.3.17 Nodeset
**Scope:** Heartbeat

**Summary**
Define the nodeset variables that should be written to the heartbeat database. The syntax is: "nodeset internal_name at node id as DBname" or "nodeset internal_name nearest location X, Y, Z as DBname".

Where internal_name is the name of the variable in the Sierra application; and DBname is the name as it should appear on the heartbeat database.

### 25.3.18 Output On Signal
**Scope:** Heartbeat

**Output On Signal {=|are|is} Signals**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Signals</td>
<td>{sigabrt</td>
<td>sigalrm</td>
</tr>
</tbody>
</table>

**Summary**
When the specified signal is raised, the output stream associated with this block will be output.

### 25.3.19 Precision
**Scope:** Heartbeat

**Precision {=|are|is} Precision**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Precision</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
The precision to be used for the output of real variables.

### 25.3.20 Start Time
**Scope:** Heartbeat

**Start Time {=|are|is} Start_time**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Start_time</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specify the time to start outputting results from this output request block. This time overrides all 'at time' and 'at step' specifications.

### 25.3.21 Stream Name
**Scope:** Heartbeat
Stream Name {=|are|is} OutputFilename

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>OutputFilename</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  
The filename of where the heartbeat data should be written. If the filename begins with the '/' character, it is an absolute path; otherwise, the path to the current directory will be prepended to the name. In addition, there are several predefined streams that can be specified. The predefined streams are 'cout' or 'stdout' specifies standard output; 'cerr', 'stderr', 'clog', or 'log' specifies standard error; 'output' or 'outputP0' specifies Sierra’s standard output which is redirected to the file specified by the '-o' option on the command line. If the file already exists, it is overwritten. If this line is omitted, then a filename will be created from the basename of the input file with a ".hrt" suffix appended.

25.3.22 Synchronize Output
Scope: Heartbeat

Summary  
In an analysis with multiple regions, it is sometimes desirable to synchronize the output of results data between the regions. This can be done by adding the SYNCHRONIZE OUTPUT command line to the results output block. If a results block has this set, then it will write output whenever a previous region writes output. The ordering of regions is based on the order in the input file, algorithmic considerations, or by solution control specifications.

Although the USE OUTPUT SCHEDULER command line can also synchronize output between regions, the SYNCHRONIZE OUTPUT command line will synchronize the output with regions where the output frequency is not under the direct control of the Sierra IO system. Examples of this are typically coupled applications where one or more of the codes are not Sierra-based applications such as Alegra and CTH. A results block with SYNCHRONIZE OUTPUT specified will also synchronize its output with the output of the external code.

The SYNCHRONIZE OUTPUT command can be used with other output scheduling commands such as time-based or step-based output specifications.

25.3.23 Termination Time
Scope: Heartbeat

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Final_time</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  
Specify the time to stop outputting results from this output request block.

25.3.24 Timestamp Format
Scope: Heartbeat

Summary  
The format to be used for the timestamp. See 'man strftime' for more information.
25.3.25 Timestep Adjustment Interval
Scope: Heartbeat

Timestep Adjustment Interval \( N_{\text{steps}} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( N_{\text{steps}} )</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify the number of steps to 'look ahead' and adjust the timestep to ensure that the specified output times or simulation end time will be hit 'exactly'.

25.3.26 Use Output Scheduler
Scope: Heartbeat

Use Output Scheduler \( \text{Timer}_\text{name} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \text{Timer}_\text{name} )</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Associates a predefined output scheduler with this output block (results, restart, heartbeat, or history).

25.3.27 Variable
Scope: Heartbeat

Summary: Define the variables that should be written to the heartbeat output. The user can request that the values of certain variables be output on the heartbeat line. These variables are limited to region and framework control data currently. The syntax is:

\[
\text{variable} = \{\text{entity_type}\} \{\text{internal_name}\} \text{ at} \\
\{\text{entity_type}\} \{\text{entity_id}\} \text{ as} \{\text{external_name}\}
\]

\[
\text{variable} = \{\text{entity_type}\} \{\text{internal_name}\} \text{ nearest location} \\
\{x, y, z\} \text{ as} \{\text{external_name}\}
\]

For global variables, use:

\[
\text{variable} = \text{global} \{\text{internal_name}\} \text{ [as} \{\text{external_name}\}\text{]}
\]

Where:

- \text{entity}_\text{type} = \text{node, element, face, edge, global}
- \text{internal}_\text{name} = \text{Sierra variable name}
- \text{entity}_\text{id} = \text{id of the node, element, face, edge that you want the specified variable output at.}
- \text{external}_\text{name} = \text{name of variable on the database.}

The names 'timestep', and 'time' can be specified as variables also. They are the current timestep and simulation time. This line can appear multiple times.
25.4 History Output

Scope: Encore Region

Begin History Output Label

Additional Steps {=|are|is} List_of_steps...
Additional Times {=|are|is} List_of_times...
At Step n Option {=|are|is} m
At Time Dt1 Option {=|are|is} Dt2
Database Name {=|are|is} StreamName
Database Type {=|are|is} DatabaseTypes
Debug
Edge [ VariableList...
Element [ VariableList...
Exists Option1 Option2
Face [ VariableList...
Global [ Variables...
Nodal [ VariableList...
Node [ VariableList...
Nodeset [ VariableList...
Output On Signal {=|are|is} Signals
Overwrite Option1 Option2
Property PropertyName {=|are|is} PropertyValue
Start Time {=|are|is} Start_time
Synchronize Output
Termination Time {=|are|is} Final_time
Timestep Adjustment Interval {=|are|is} Nsteps
Title
Use Output Scheduler Timer_name
Variable {=|are|is} Option [ Variable_list... ]

End

Summary  Describes the location and type of the output stream used for outputting history for the enclosing region.

25.4.1 Additional Steps

Scope: History Output

Additional Steps {=|are|is} List_of_steps...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>List_of_steps</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Additional simulation steps when output should occur.
### 25.4.2 Additional Times

**Scope:** History Output

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>List_of_times</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Additional simulation times when output should occur.

### 25.4.3 At Step

**Scope:** History Output

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>integer</td>
<td>undefined</td>
</tr>
<tr>
<td>m</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specify an output interval in terms of the internal iteration step count. The first step specifies the step count at the beginning of this interval and the second step specifies the output frequency to be used within this interval.

### 25.4.4 At Time

**Scope:** History Output

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dt1</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>Dt2</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specify an output interval in terms of the internal simulation time. The first time specifies the time at the beginning of this time interval and the second time specifies the output frequency to be used within this interval.

### 25.4.5 Database Name

**Scope:** History Output

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>StreamName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
The base name of the database containing the output history. If the filename begins with the '/' character, it is an absolute path; otherwise, the path to the current directory will be prepended to the name. If this line is omitted, then a filename will be created from the basename of the input file with a ".h" suffix appended.
25.4.6 Database Type

Scope: History Output

| Database Type {=|are|is} DatabaseTypes |
|--------------------------------------|
| Parameter Value Default              |
| DatabaseTypes {catalyst|dof|dof_exodus|exodus|exodusii|generated|genesis|parallel_exodus|xdmf} undefined |

Summary
The database type/format to be used for the output history.

25.4.7 Debug

Scope: History Output

Summary
Turn on debugging output.

25.4.8 Edge

Scope: History Output

Summary
Define the edge variables that should be written to the history database. The syntax is: "edge internal_name at edge id as DBname" or "edge internal_name nearest location X, Y, Z as DBname".

Where internal_name is the name of the variable in the Sierra application; and DBname is the name as it should appear on the history database.

25.4.9 Element

Scope: History Output

Summary
Define the element variables that should be written to the history database. The syntax is: "element internal_name at element id as DBname" or "element internal_name nearest location X, Y, Z as DBname".

Where internal_name is the name of the variable in the Sierra application; and DBname is the name as it should appear on the history database.

25.4.10 Exists

Scope: History Output

Summary
Specify the behavior when creating this database and there is an existing file with the same name. The default behavior is "OVERWRITE" which deletes the existing file and creates a new file of the same name. "APPEND" will (if possible) append the new data to the end of the existing file. "ABORT" will print an error message and end the analysis. "ADD_SUFFIX" will add a -s???? suffix where the ???? is replaced by a sequential number starting at 0002.
25.4.11 Face
Scope: History Output

Summary Define the face variables that should be written to the history database. The syntax is: "face internal_name at face id as DBname" or "face internal_name nearest location X, Y, Z as DBname".

Where internal_name is the name of the variable in the Sierra application; and DBname is the name as it should appear on the history database.

25.4.12 Global
Scope: History Output

Summary Define the global/reduction variables that should be written to the history database. The syntax is: "global internal_name as DBname".

Where internal_name is the name of the variable in the Sierra application; and DBname is the name as it should appear on the history database.

25.4.13 Nodal
Scope: History Output

Summary Define the nodal variables that should be written to the history database. The syntax is: "nodal internal_name at node id as DBname" or "nodal internal_name nearest location X, Y, Z as DBname".

Where internal_name is the name of the variable in the Sierra application; and DBname is the name as it should appear on the history database.

25.4.14 Node
Scope: History Output

Summary Define the nodal variables that should be written to the history database. The syntax is: "node internal_name at node id as DBname" or "node internal_name nearest location X, Y, Z as DBname".

Where internal_name is the name of the variable in the Sierra application; and DBname is the name as it should appear on the history database.

25.4.15 Nodeset
Scope: History Output
Define the nodeset variables that should be written to the history database. The syntax is:
"nodeset internal_name at node id as DBname" or "nodeset internal_name nearest location X, Y, Z as DBname".
Where internal_name is the name of the variable in the Sierra application; and DBname is the name as it should appear on the history database.

### 25.4.16 Output On Signal

**Scope:** History Output

<table>
<thead>
<tr>
<th>Output On Signal</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>{=</td>
<td>are</td>
<td>is} Signals</td>
</tr>
</tbody>
</table>

Summary: When the specified signal is raised, the output stream associated with this block will be output.

### 25.4.17 Overwrite

**Scope:** History Output

Summary: (DEPRECATED, Use EXISTS) Specify whether the database should be overwritten if it exists. The default behavior is to overwrite unless this command is specified in the output block and either off, false, or no is specified.

### 25.4.18 Property

**Scope:** History Output

<table>
<thead>
<tr>
<th>Property</th>
<th>PropertyName</th>
<th>PropertyValue</th>
</tr>
</thead>
<tbody>
<tr>
<td>PropertyName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>PropertyValue</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Define a database property named "PropertyName" with the value "PropertyValue". If PropertyValue consists of all digits, it will define an integer property. If PropertyValue is "true" or "yes" or "false" or "no", it will define a logical property; otherwise it will define a string property. Supported properties are typically database dependent; Some history-related properties are: VARIABLE_NAME_CASE = upper|lower MAX_NAME_LENGTH = value (32)
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Start_time</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify the time to start outputting results from this output request block. This time overrides all 'at time' and 'at step' specifications.

25.4.20 Synchronize Output

Scope: History Output

Summary: In an analysis with multiple regions, it is sometimes desirable to synchronize the output of results data between the regions. This can be done by adding the SYNCHRONIZE OUTPUT command line to the results output block. If a results block has this set, then it will write output whenever a previous region writes output. The ordering of regions is based on the order in the input file, algorithmic considerations, or by solution control specifications.

Although the USE OUTPUT SCHEDULER command line can also synchronize output between regions, the SYNCHRONIZE OUTPUT command line will synchronize the output with regions where the output frequency is not under the direct control of the Sierra IO system. Examples of this are typically coupled applications where one or more of the codes are not Sierra-based applications such as Alegra and CTH. A results block with SYNCHRONIZE OUTPUT specified will also synchronize its output with the output of the external code.

The SYNCHRONIZE OUTPUT command can be used with other output scheduling commands such as time-based or step-based output specifications.

25.4.21 Termination Time

Scope: History Output

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Final_time</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify the time to stop outputting results from this output request block.

25.4.22 Timestep Adjustment Interval

Scope: History Output

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nsteps</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify the number of steps to 'look ahead' and adjust the timestep to ensure that the specified output times or simulation end time will be hit 'exactly'.
25.4.23 Title
Scope: History Output

Summary Specify the title to be used for this specific output block.

25.4.24 Use Output Scheduler
Scope: History Output

Use Output Scheduler Timer_name

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Timer_name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Associates a predefined output scheduler with this output block (results, restart, heartbeat, or history).

25.4.25 Variable
Scope: History Output

Summary Define the variables that should be written to the history database. The syntax is: "variable = entity internal_name at entity id as DBname" or "variable = entity internal_name nearest location X, Y, Z as DBname" or "variable = entity internal_name at location X, Y, Z as DBname".

Where entity is 'node', 'element', 'face', or 'edge'; internal_name is the name of the variable in the Sierra application; and DBname is the name as it should appear on the history database.

25.5 Data Probe
Scope: Equation System

Begin Data Probe probeName

Nodal_Id fieldName Node_Id {=|are|is} node_id [ Label label ]
Nodal_Location fieldName Location {=|are|is} location1 location2 [ location3 ] [ Label label ]
Parametric Tolerance parametric_tolerance...
Restriction Parts restriction_parts...
At Step startingStep Increment {=|are|is} increment
Output To File [ fileName ]
End
The Aria Data Probe capability allows the analyst to query data for fields of interest (e.g. temperature, pressure) during a simulation at times and locations specified by the analyst. The user can instantiate multiple data probes each with multiple probe points. This command creates an Aria Data Probe with the name given by probeName.

25.5.1 Nodal_Id

Scope: Data Probe

**Nodal_Id fieldName Node_Id {=|are|is} node_id [ Label label ]**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>fieldName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>node_id</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: If used, the data probe will include a point specified by the global id of a node.

Description: Sets the nodal field name, node global id, and optionally the label for each probe point. If a label is provided, a global variable with that name will be created. If no label is provided, a global variable will be created with a name that is the concatenation of the Data Probe name and the index of the probe point. The field_data at the specified node (and only that node) is associated with the probe point and its corresponding global variable.

25.5.2 Nodal_Location

Scope: Data Probe

**Nodal_Location fieldName Location {=|are|is} location1 location2[ location3] [ Label label ]**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>fieldName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>location</td>
<td>real_1 real_2[ real_3]</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: If used, the data probe will include a point specified by location (x,y,z).

Description: Sets the nodal field name, spatial location, and optionally the label for this probe point. If a label is provided, a global variable with that name will be created. If no label is provided, a global variable will be created with a name that is the concatenation of the Data Probe name and the index of the probe point. If the point lies within an element within the restricted set of blocks (see below), the given field will be interpolated in that element. If the point lies outside the mesh, then it is projected to the nearest point on the nearest element (face, edge, or vertex), and interpolation is used to compute the value of the field, unless the point lies too far outside the closest element (as indicated by the parametric tolerance below).

25.5.3 Parametric Tolerance

Scope: Data Probe

**Parametric Tolerance parametric_tolerance...**
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>parametric_tolerance</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  
The user may specify a parametric tolerance that is used to determine whether or not the data probe point is close enough to an element for the probe point to be considered valid. The parametric tolerance is a numeric value that indicates the maximum allowed parametric coordinate of the data probe point with respect to the closest element found during the search. For instance, if the parametric tolerance is set to 2.0, the data probe point is allowed to have a parametric coordinate of up to 2.0 with respect to the closest element that is found. However, if the parametric coordinate of the point with respect to this element is $>2.0$, the tolerance check will fail and the simulation will end with an error describing this failure. For convenience, all points within a probe are checked for the parametric tolerance condition, and if any failures are found, they are all reported together rather than the simulation halting on the first failure of this condition.

### 25.5.4 Restriction Parts
**Scope:** Data Probe

**Summary**  
Mesh parts for which data probe will be applied. Only elements within these parts will be searched for data probe points. If no restriction is used, all mesh parts will be searched.

### 25.5.5 At Step
**Scope:** Data Probe

**Summary**  
Write the results of the data probe beginning at step (startingStep) and output every specified number of steps (increment).

### 25.5.6 Output To File
**Scope:** Data Probe

**Summary**  
Enable data output to file specified by filename. Data Probe Point values are additionally sent to the Aria log file.
25.6 Restart Overview

Sierra Framework services provide convenient utilities for restarting an analysis from previous results. The most general capability supplements the results of a previous analysis with internal state variables to continue an analysis. In this case the input mesh is supplied from the Input Database Name from the Finite Element Model command block 3.1 and the restart information is obtained from the the Input Database Name from the Restart Data command block. Continuation of a job from a solution plane is invoked using the RESTART TIME command line.

Note that special considerations are warranted when initializing problems utilizing bulk volume elements 31.4 with restart files not containing the bulk volume elements.

25.7 Restart Data

Scope: Encore Region

Begin Restart Data Label

Additional Steps \{=|are|is\} List_of_steps...
Additional Times \{=|are|is\} List_of_times...
At Step n Option \{=|are|is\} m
At Time Dt1 Option \{=|are|is\} Dt2
Component Separator Character Option Separator
Cycle Count \{=|are|is\} Count
Database Name \{=|are|is\} StreamName
Database Type \{=|are|is\} DatabaseTypes
Debug Dump
Dump All
Decomposition Method \{=|are|is\} Method
Exists Option1 Option2
File Cycle Count \{=|are|is\} Count
Input Database Name \{=|are|is\} StreamName
Optional
Output Database Name \{=|are|is\} StreamName
Output On Signal \{=|are|is\} Signals
Overlay Count \{=|are|is\} Count
Overwrite Option1 Option2
Property PropertyName \{=|are|is\} PropertyValue
Restart \{=|are|is\} Option
Restart Time \{=|are|is\} Time
Start Time \{=|are|is\} Start_time
Synchronize Output
Termination Time \{=|are|is\} Final_time
Timestep Adjustment Interval \{=|are|is\} Nsteps
Use Output Scheduler Timer_name
End

Summary Describes the data required to output and input restart data for the enclosing region.

25.7.1 Additional Steps
Scope: Restart Data

Additional Steps \{=|are|is\} List_of_steps...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>List_of_steps</td>
<td>integer...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Additional simulation steps when output should occur.

25.7.2 Additional Times
Scope: Restart Data

Additional Times \{=|are|is\} List_of_times...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>List_of_times</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Additional simulation times when output should occur.

25.7.3 At Step
Scope: Restart Data

At Step \text{n} Option \{=|are|is\} m

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>integer</td>
<td>undefined</td>
</tr>
<tr>
<td>m</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specify an output interval in terms of the internal iteration step count. The first step specifies the step count at the beginning of this interval and the second step specifies the output frequency to be used within this interval.

25.7.4 At Time
Scope: Restart Data

At Time \text{Dt1} Option \{=|are|is\} \text{Dt2}
Parameter | Value | Default
---|---|---
$Dt1$ | real | undefined
$Dt2$ | real | undefined

Summary
Specify an output interval in terms of the internal simulation time. The first time specifies the time at the beginning of this time interval and the second time specifies the output frequency to be used within this interval.

### 25.7.5 Component Separator Character

Scope: Restart Data

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Component Separator Character</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
The separator is the single character used to separate the output variable basename (e.g. "stress") from the suffices (e.g. "xx", "yy") when displaying the names of the individual variable components. For example, the default separator is ",", which results in names similar to "stress_xx", "stress_yy", ... "stress_zx". To eliminate the separator, specify an empty string ("") or NONE.

### 25.7.6 Cycle Count

Scope: Restart Data

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cycle Count</td>
<td>Count</td>
<td>integer</td>
</tr>
</tbody>
</table>

Summary
Specify the number of restart steps which will be written to the restart database before previously written steps are overwritten. For example, if the cycle count is 5 and restart is written every 0.1 seconds, the restart system will write 0.1, 0.2, 0.3, 0.4, 0.5 to the database. It will then overwrite the first step with data from time 0.6, the second with time 0.7. At time 0.8, the database would contain data at times 0.6, 0.7, 0.8, 0.4, 0.5. Note that time will not necessarily be monotonically increasing on a database that specifies the cycle count.

### 25.7.7 Database Name

Scope: Restart Data

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Database Name</td>
<td>StreamName</td>
<td>string</td>
</tr>
</tbody>
</table>

Summary
The database containing the input and/or output restart data. If this analysis is being restarted, restart data will be read from this file. If the analysis is writing restart data, the data will be written to this file. It will be overwritten if it exists (after being read if
applicable). If the filename begins with the '/' character, it is an absolute path; otherwise, the path to the current directory will be prepended to the name. See also the 'Input Database' and 'Output Database' commands.

25.7.8 Database Type
Scope: Restart Data

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Database Types</td>
<td>catalyst</td>
<td>dof</td>
</tr>
</tbody>
</table>

Summary: The database type/format used for the restart file.

25.7.9 Debug Dump
Scope: Restart Data

Summary: Specify whether the the restart system will write the restart data immediately after reading the restart data if the run is restarting. The output data can be compared with the restart input data to determine whether they match.

25.7.10 Dump All
Scope: Restart Data

Summary: Specify that the restart system should treat all variables as needed for restart whether they are persistent, temporary, or constant. Used only for debugging restart.

25.7.11 Decomposition Method
Scope: Restart Data

Summary: The decomposition algorithm to be used to partition elements to each processor in a parallel run.

25.7.12 Exists
Scope: Restart Data

Summary: Specify the behavior when creating this database and there is an existing file with the same name. The default behavior is "OVERWRITE" which deletes the existing file and creates a new file of the same name. "APPEND" will (if possible) append the new data to the end of the existing file. "ABORT" will print an error message and end the analysis.
25.7.13 File Cycle Count
Scope: Restart Data

File Cycle Count {=|are|is} Count

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Count</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Each restart dump will be written to a separate file suffixed with A,B, ... The count specifies how many separate files are used before the cycle repeats. For example, if "FILE CYCLE COUNT = 3" is specified, the restart dumps would be written to file-A.rs, file-B.rs, file-C.rs, file-A.rs, ... The maximum value for the cycle count is 26.

25.7.14 Input Database Name
Scope: Restart Data

Input Database Name {=|are|is} StreamName

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>StreamName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: The database containing the input restart data. If this analysis is being restarted, restart data will be read from this file. See also the 'Database' and 'Output Database' commands.

25.7.15 Optional
Scope: Restart Data

Summary: The database will be read if it exists, but it is not an error if there is no restart database to read for this region during a restarted analysis.

25.7.16 Output Database Name
Scope: Restart Data

Output Database Name {=|are|is} StreamName

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>StreamName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: The database containing the output restart data. If the analysis is writing restart data, the data will be written to this file. It will be overwritten if it exists. See also the 'Database' and 'Input Database' commands.

25.7.17 Output On Signal
Scope: Restart Data
### Output On Signal

Parameter: Signals  

<table>
<thead>
<tr>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>{sigabrt, sigalrm, sigfpe, sighup, sigill, sigint, sigkill, sigpipe, sigquit, sigsev, sigterm, sigusr1, sigusr2}</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** When the specified signal is raised, the output stream associated with this block will be output.

### 25.7.18 Overlay Count

Scope: Restart Data

Parameter: Count  

<table>
<thead>
<tr>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** Specify the number of restart outputs which will be overlayed on top of the last written step. For example, if restarts are being output every 0.1 seconds and the overlay count is specified as 2, then restart will write times 0.1 to step 1 of the database. It will then write 0.2 and 0.3 also to step 1. It will then increment the database step and write 0.4 to step 2; overlay 0.5 and 0.6 on step 2... At the end of the analysis, assuming it runs to completion, the database would have times 0.3, 0.6, 0.9, ... However, if there were a problem during the analysis, the last step on the database would contain an intermediate step.

### 25.7.19 Overwrite

Scope: Restart Data

**Summary:** (DEPRECATED, Use EXISTS) Specify whether the restart database should be overwritten if it exists. The default behavior is to overwrite unless this command is specified in the restart block and either off, false, or no is specified.

### 25.7.20 Property

Scope: Restart Data

Parameter: PropertyName PropertyValue  

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>PropertyName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>PropertyValue</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** Define a database property named "PropertyName" with the value "PropertyValue". If PropertyValue consists of all digits, it will define an integer property. If PropertyValue is "true" or "yes" or "false" or "no", it will define a logical property; otherwise it will...
define a string property. If PropertyName consists of multiple strings, they will be con-
cated together with "_" separating the individual words. Supported properties are typi-
cally database dependent; Current properties are: COMPRESSION_LEVEL = [0..9] COM-
PRESSION_SHUFFLE = true|false|on|off FILE_TYPE = netcdf4 (forces use of netcdf-4 
hdf5-based file) INTEGER_SIZE_DB = 4|8 INTEGER_SIZE_API = 4|8 LOGGING = 
true|false|on|off MAX_NAME_LENGTH = value

25.7.21 Restart
Scope: Restart Data

Summary Specify that the analysis should be restarted from the last common time on all restart
databases for each Region in the analysis. In addition to this line command, each Region
in the analysis (strictly, only the region(s) that will be restarted) must have a restart block
specifying the database to read the restart state data.

25.7.22 Restart Time
Scope: Restart Data

Restart Time \{=}|are|is\ Time

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specify the time that the analysis will be restarted. In addition to this line command, each
Region in the analysis (strictly, only the region(s) that will be restarted) must have a restart block
specifying the database to read the restart state data. The restart 'time' must be
greater than zero and less than or equal to the termination time.

25.7.23 Start Time
Scope: Restart Data

Start Time \{=}|are|is\ Start_time

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Start_time</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specify the time to start outputting results from this output request block. This time overrides
all 'at time' and 'at step' specifications.

25.7.24 Synchronize Output
Scope: Restart Data

Summary In an analysis with multiple regions, it is sometimes desirable to synchronize the output of
results data between the regions. This can be done by adding the SYNCHRONIZE OUTPUT
command line to the results output block. If a results block has this set, then it will write
output whenever a previous region writes output. The ordering of regions is based on the
order in the input file, algorithmic considerations, or by solution control specifications.
Although the USE OUTPUT SCHEDULER command line can also synchronize output be-
tween regions, the SYNCHRONIZE OUTPUT command line will synchronize the output
with regions where the output frequency is not under the direct control of the Sierra IO sys-
tem. Examples of this are typically coupled applications where one or more of the codes are
not Sierra-based applications such as Alegra and CTH. A results block with SYNCHRONIZE
OUTPUT specified will also synchronize its output with the output of the external code.
The SYNCHRONIZE OUTPUT command can be used with other output scheduling com-
mands such as time-based or step-based output specifications.

### 25.7.25 Termination Time

**Scope:** Restart Data

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Final_time</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specify the time to stop outputting results from this output request block.

### 25.7.26 Timestep Adjustment Interval

**Scope:** Restart Data

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nsteps</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specify the number of steps to 'look ahead' and adjust the timestep to ensure that the specified
output times or simulation end time will be hit 'exactly'.

### 25.7.27 Use Output Scheduler

**Scope:** Restart Data

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Timer_name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Associates a predefined output scheduler with this output block (results, restart, heartbeat,
or history).

### 25.7.28 Restart Time

**Scope:**

778
### Summary
Specify the time that the analysis will be restarted. In addition to this line command, each Region in the analysis (strictly, only the region(s) that will be restarted) must have a restart block specifying the database to read the restart state data. The restart 'time' must be greater than zero and less than or equal to the termination time.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Chapter 26

Input Output Region Reference

26.1 Input_Output Region Overview

For some coupled simulations one can approximate part of the problem physics independent of the entire problem physics. In order to facilitate this type of loose application coupling the Sierra Framework provides the ability to input datasets that include the output of other simulations. An application can then make requests of information from these datasets. In fulfilling these requests, data can be extracted from these datasets and be copied or interpolated to another problem domain. Moreover these requests can be satisfied by data interpolated through time. The mechanism provided to achieve this end goal is known as the Input_Output Region and its usage is described in what follows.

The input_output region works in tandem with transfer 15.1 and solution control 14. Here transfer carries out the communication of data and solution control provides synchronization of the data transfer. Note that just like other Sierra Regions the input_output region must have its own Finite Element model command block defined.

As an example, let us assume that an input mesh for an Input_Output Region contains a nodal variable ConvCoeff that we wish to use in another Region. In this case an outline for one-way transfer of ConvCoeff to a Region, second_region, in a steady-state problem would be:

```sierra
Begin Sierra
  
  Begin Finite Element Model input_transfer
    
    End
    
  Begin Transfer my_first_transfer
    
    transfer commands for input_output_region to second_region
    
    SEND field hNd state none TO ConvCoeff state none
    
    End
    
  Begin Procedure My_Aria_Procedure
    
    Begin Solution Control Description
      Use System Main
      Begin System Main
        Begin Sequential MySolveBlock
          Advance io_region
          transfer my_first_transfer
```
Advance second_region
End
End
End

Begin Input_Output io_region
    USE FINITE ELEMENT MODEL my_input_transfer
End

Begin Aria Region second_region
    .
    use Finite Element Model input_transfer
    .
    USER FIELD REAL NODE SCALAR ConvCoeff on surface_1
    .
End
End
End

End Sierra

26.2 Input_Output Region

Scope: Procedure

Begin Input_Output Region Parameter_block_name
    Create Nodal Field Field_name Of Type Option And Dimension Dimension
    Fixed Time [ {=}is Fixed_time ]
    Offset Time {=} Period_offset_time
    Periodicity Time {=} Periodicity_time
    Start Time {=} Start_time
    Use Finite Element Model ModelName [ Model Coordinates Are Nodal_variable_name ]
    Begin Results Output Label
    End

End

Summary
BEGIN INPUT TRANSFER model_name USE FINITE ELEMENT MODEL fred START
TIME is 0 OFFSET TIME is 1 PERIODICITY TIME is 10 END INPUT TRANSFER
model_name

26.2.1 Create Nodal Field

Scope: Input_Output Region

Create Nodal Field Field_name Of Type Option And Dimension Dimension
### Summary

Creates a Nodal Field name field_name on the region.

#### 26.2.2 Fixed Time

**Scope:** Input_Output Region

**Summary**

The line specifies that the database will be read for a single, fixed time. Specifying the actual time is optional. If the time is not specified, the final time plane in the database will be read.

NOTE: This option takes precedence over the periodic specifications given by START TIME, PERIODICITY TIME, and OFFSET TIME.

if FIXED TIME is specified then if FIXED TIME value is given then (eg., FIXED TIME is 1.) DATABASE TIME = FIXED TIME else (eg., FIXED TIME) DATABASE TIME = last time in database else if PERIODICITY TIME greater than 0 then if APPLICATION TIME less than or equal to START TIME then DATABASE TIME = APPLICATION TIME else DATABASE TIME = START TIME + (APPLICATION TIME - START TIME) modulo PERIODICITY TIME else DATABASE TIME = APPLICATION TIME now add OFFSET TIME to the computed DATABASE TIME

#### 26.2.3 Offset Time

**Scope:** Input_Output Region

**Summary**

This value is added to the application time to determine what database time slice to input. If OFFSET TIME were 15 than at application time 0 database time slice 15 would be read from the file and used for the initial values. At application time 1, database time slice 16 would be read. NOTE: The OFFSET TIME is added in after the START TIME and PERIODICITY TIME are used. The FIXED TIME option takes precedence over this option.

if FIXED TIME is specified then if FIXED TIME value is given then (eg., FIXED TIME is 1.) DATABASE TIME = FIXED TIME else (eg., FIXED TIME) DATABASE TIME = last time in database else if PERIODICITY TIME greater than 0 then if APPLICATION TIME less than or equal to START TIME then DATABASE TIME = APPLICATION TIME else DATABASE TIME = START TIME + (APPLICATION TIME - START TIME) modulo PERIODICITY TIME else DATABASE TIME = APPLICATION TIME now add OFFSET TIME to the computed DATABASE TIME

#### 26.2.4 Periodicity Time

**Scope:** Input_Output Region

**Summary**

This value is added to the application time to determine what database time slice to input. If PERIODICITY TIME were 15 than at application time 0 database time slice 15 would be read from the file and used for the initial values. At application time 1, database time slice 16 would be read. NOTE: The PERIODICITY TIME is added in after the START TIME and PERIODICITY TIME are used. The FIXED TIME option takes precedence over this option.

if FIXED TIME is specified then if FIXED TIME value is given then (eg., FIXED TIME is 1.) DATABASE TIME = FIXED TIME else (eg., FIXED TIME) DATABASE TIME = last time in database else if PERIODICITY TIME greater than 0 then if APPLICATION TIME less than or equal to START TIME then DATABASE TIME = APPLICATION TIME else DATABASE TIME = START TIME + (APPLICATION TIME - START TIME) modulo PERIODICITY TIME else DATABASE TIME = APPLICATION TIME now add OFFSET TIME to the computed DATABASE TIME

---

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Field_name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Dimension</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

---

**Parameter**

Field_name

**Value**

string

**Default**

undefined

---

**Parameter**

Period_offset_time

**Value**

real

**Default**

undefined

---

**Parameter**

Periodicity_time

**Value**

Periodicity_time

---

783
Parameter | Value | Default
--- | --- | ---
Periodicity time | real | undefined

**Summary**

START TIME and PERIODICITY TIME taken together give the time frame from the input database to use to initialize the application values. If START TIME is 25 and PERIODICITY TIME is 10, then time slices from 25 to 35 will be used over and over again as the application time runs from 0 to whatever. In general DATABASE TIME is (APPLICATION TIME - START TIME) modulo PERIODICITY TIME after the application time reaches the START TIME.

NOTE: The OFFSET TIME is added in after the START TIME and PERIODICITY TIME are used. The FIXED TIME option take precedence over this option.

if FIXED TIME is specified then if FIXED TIME value is given then (eg., FIXED TIME is 1.) DATABASE TIME = FIXED TIME else (eg., FIXED TIME) DATABASE TIME = last time in database else if PERIODICITY TIME greater than 0 then if APPLICATION TIME less than or equal to START TIME then DATABASE TIME = APPLICATION TIME else DATABASE TIME = START TIME + (APPLICATION TIME - START TIME) modulo PERIODICITY TIME else DATABASE TIME = APPLICATION TIME now add OFFSET TIME to the computed DATABASE TIME

### 26.2.5 Start Time

**Scope:** Input_Output Region

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Start_time</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

The time in which to start applying PERIODICITY TIME. If PERIODICITY TIME is not specified then START TIME is ignored.

NOTE: The OFFSET TIME is added in after the START TIME and PERIODICITY TIME are used. The FIXED TIME option take precedence over this option.

if FIXED TIME is specified then if FIXED TIME value is given then (eg., FIXED TIME is 1.) DATABASE TIME = FIXED TIME else (eg., FIXED TIME) DATABASE TIME = last time in database else if PERIODICITY TIME greater than 0 then if APPLICATION TIME less than or equal to START TIME then DATABASE TIME = APPLICATION TIME else DATABASE TIME = START TIME + (APPLICATION TIME - START TIME) modulo PERIODICITY TIME else DATABASE TIME = APPLICATION TIME now add OFFSET TIME to the computed DATABASE TIME

### 26.2.6 Use Finite Element Model

**Scope:** Input_Output Region

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>ModelName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Associates a predefined finite element model with this region.
Chapter 27

Log File Output

27.1 Introduction

Aria, as well as all other Sierra codes, provides log file output capability which aids users in performing such actions as viewing run information, runtime/performance monitoring, and verifying input file accuracy. While much of the log file output is standardized for the Sierra codes, there are sections, especially in runtime monitoring, that are specific to Aria. This chapter will discuss the different sections of the log file and give examples of these sections in order to help the user assimilate and make the best use of the information that is output.

For this explanation, the Aria Foam Decomposition example problem will be used and can be found on the Sandia Computational Simulation webpage. This example was chosen due to its use of several different types of physics, such as element death, enclosure radiation, and chemistry modeling, as well as its use of adaptive time stepping.

27.2 Preamble

The log file preamble contains useful information about run information, input file parsing, and model setup.

27.2.1 Run Information

At the very beginning of the log file is printed information about the selected Sierra executable, where and when the model is run, and specific version information. Below is a section of the log file preamble which shows this information.

+----+----+----+----+----+----+----+----+----+----+----+----+----+----+----+----+----+----+

**Aria**

Coupled multiphysics including Navier-Stokes, elasticity, energy transport, species transport, electrostatics; free and moving boundaries; transient or steady state.

Version 4.33.1-398-gaeb9bfbb

With coupled mechanics support for
Aria - Coupled multiphysics
Encore - Solution Verification Analysis Region

Sandia National Laboratories
Albuquerque, New Mexico and Livermore, California

Please email questions and comments to
sierra-help@sandia.gov

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+----+----+----+----+----+----+----+----+----+----+----+----+----+----+----+----+
 Directory /home/rpshaw/foamDecomposition/
 Executable /projects/sierra/linux_rh6/install/master/bin/aria
    Built May 12 2014 18:46:01
 Build Options linux intel-12.1 release
 Run Started May 14 2014 11:04:22
    User rpshaw
 Architecture cee-build001
   Host cee-build001
 Hardware x86_64
 Running Linux
 Processors 1
 Processing foamDecomposition.i

<table>
<thead>
<tr>
<th>Product</th>
<th>Version</th>
<th>Qualifier</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACME</td>
<td>2.9.0</td>
<td></td>
</tr>
<tr>
<td>Aria</td>
<td>4.33.1-398-gaeb9bfbb</td>
<td></td>
</tr>
<tr>
<td>Chaparral</td>
<td>3.3.1 development</td>
<td>unreleased</td>
</tr>
<tr>
<td>Encore</td>
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<td></td>
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<tr>
<td>FEI</td>
<td>2.24.02</td>
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<tr>
<td>GDSW Dev</td>
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</tr>
<tr>
<td>UtilityLib</td>
<td>4.33.1-398-gaeb9bfbb</td>
<td></td>
</tr>
<tr>
<td>Zoltan</td>
<td>3.8</td>
<td></td>
</tr>
</tbody>
</table>

786
Here, it can be seen that Aria version 4.33.1-398-gaeb9bfbb is used, and further, it can be seen that the master (i.e., Version-of-the-Day/VOTD) version is used based on the executable directory:

Executable /projects/sierra/linux_rh6/install/master/bin/aria

It can also be seen that the run was started at May 14 2014 11:04:22 on cee-build001 with 1 processor in /home/rpshaw/foamDecomposition/.

More specific version information for each product that Aria uses is shown at the bottom of this section, for those who are interested in or concerned with more detail about how their model is being run.

27.2.2 Input File

The log file next contains a complete copy of the input file which is used to run the model, directly following the lines shown below:

Reading foamDecomposition.i May 14 2014 11:04:26

Note that the input file shown is the one that is used for parsing, so any pre-processing operations such as aprepro will have already been completed in the input file shown in the log file. Also, this section will be where parsing errors show up, if there are any in the input file.

In the event of misplacement of an input file, one can also extract the input file from the log file, which can save a considerable amount of time.

27.2.3 Instantiation

After the input file has been parsed, Aria will then instantiate all systems required to run the input file and will output the results of the instantiation to the log file. The instantiation begins with the lines

Instantiating system May 14 2014 11:04:26

This section of the log file is not often used by analysts, but can be useful in debugging non-obvious issues. The following is a list of common items that may be seen in the instantiation section:

- Setting of default values for unspecified variables
  
  *Example* DEFAULTING FAILED_TIME_STEP_SIZE_RATIO = 0.500000 for region AriaRegion in time block Transient_time_block_1

- Configuration of mesh parts as needed by different inputs
  
  *Example* Adding death bc iopart for element death with name surface_death_temp1

- Creation of edge/element mechanics
  
  *Example* Configuring Element mechanics for region "AriaRegion" equation system "main"

- Configuration of boundary/initial condition mechanics
Example Configuring Convective Flux for region "AriaRegion" equation system "main"...

- Configuration of other mechanics such as enclosure radiation and chemistry

Example Configuring Enclosure Radiation BC mechanics for region "AriaRegion" equation system "main"...

The very end of this section of the log file contains the number of errors and warnings that were found during the parsing of the input file. This can be helpful in helping users pinpoint if they incorrectly set up some parameter that still allows the simulation to continue.

### 27.2.4 Procedure Initialization

This section of the log file begins with the following lines:

+----+----+----+----+----+----+----+----+----+----+----+----+----+----+----+----+

Running procedure AriaProcedure May 14 2014 11:04:26
Reading mesh May 14 2014 11:04:26

The most useful portion of this section is the list of variables available for output. These variables are shown for each Region that is found in the input file and are separated by the type of field on which they reside, e.g., nodes, elements, edges, and faces. Global variables that may be output are also listed.

### 27.3 Run-time Reporting

After the preamble of the log file comes the actual execution of the model, including linear and nonlinear solver information, time stepping, and variable calculation. The information contained in this section can assist the user in determining whether the model is stable and if the correct results are being obtained. To illustrate this section of the log file, one time step of the run is shown below and will be further explained in the following sections.

+----+----+----+----+----+----+----+----+----+----+----+----+----+----+----+----+

Minimum Time Step Selection:

<table>
<thead>
<tr>
<th>min</th>
<th>max</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.000e-02</td>
<td>5.287e+00</td>
</tr>
</tbody>
</table>

AriaRegion

Transient Time_Block_2: dt = 5.287

---------------------------------------------------

Transient Time_Block_2, step 169, time 2.1985e+03, time step 5.2875e+00, 61.06% complete

---------------------------------------------------

Advance AriaRegion, time 2198, time step 5.287

Memory Usage: current = 209186816 (199.5 M), high-water-mark = 217268224 (207.2 M)

Equation System AriaRegion->main:

* Step : Transient, Strategy: NEWTON, Time: 2.20e+03, Step: 5.29e+00
* Mesh : Processor 0 of 1: 1253 of 1253 elems, 1363 of 1363 nodes
* Computing View Factors for enclosure space

NONLINEAR LINEAR

788
### Step Resid Delta Itns Status Resid Asm/Slv Time

<table>
<thead>
<tr>
<th>Step</th>
<th>Resid</th>
<th>Delta</th>
<th>Itns</th>
<th>Status</th>
<th>Resid</th>
<th>Asm/Slv Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>6.37e-02</td>
<td>1.10e+01</td>
<td>59</td>
<td>ok</td>
<td>8.23e-07</td>
<td>7.0e-03/2.0e-03</td>
</tr>
<tr>
<td>2</td>
<td>1.10e-04</td>
<td>8.18e-03</td>
<td>58</td>
<td>ok</td>
<td>1.19e-09</td>
<td>7.0e-03/2.0e-03</td>
</tr>
<tr>
<td>3</td>
<td>3.06e-07</td>
<td>8.81e-05</td>
<td>61</td>
<td>ok</td>
<td>3.68e-12</td>
<td>6.0e-03/2.0e-03</td>
</tr>
</tbody>
</table>

Termination reason: 8.80515e-05 < nonlinear_correction_tolerance(0.001)

### Field min @ id max @ id max-chg @ id pred-err pred-min @ id pred-max @ id

<table>
<thead>
<tr>
<th>Field</th>
<th>min</th>
<th>id</th>
<th>id</th>
<th>max</th>
<th>id</th>
<th>id</th>
<th>max-chg</th>
<th>id</th>
<th>pred-err</th>
<th>id</th>
<th>pred-min</th>
<th>id</th>
<th>pred-max</th>
<th>id</th>
</tr>
</thead>
<tbody>
<tr>
<td>TEMPERATURE</td>
<td>3.8650e+02</td>
<td>412</td>
<td>8.4155e+02</td>
<td>512</td>
<td>3.93e+00</td>
<td>1443</td>
<td>3.53e-04</td>
<td>3.8650e+02</td>
<td>412</td>
<td>8.4155e+02</td>
<td>512</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

DT_TEMPERATURE : MIN(value,loc) = (-1.18e+00, 1443) : MAX(value,loc) = (9.68e-01, 249)

Segregated solution procedure converged after 1 iterations.

Global predictor error = 3.531e-04

Time step selection: dt <= 2.326e+01 (based on Predictor-Corrector Tolerance).

Time step selection: dt <= 1.057e+01 (based on Maximum Time Step Size Ratio).

Time step selection: dt <= 6.000e+01 (based on Maximum Time Step Size).

Time step selection: dt = 5.277e+00 (Minimum Chemistry timestep block_6).

Time step selection: dt <= 1.277e+01 (based on Stability limit for dt Ratio).

Time step selection: dt = 1.057e+01 (Adaptive time stepping result).

Killed 1 elements this timestep, for a total of 87 dead.

Global Variable Value

| Conv_ipo | 189.224 |
| Conv_ipo_surface_block_2_edge2_2 | 0 |
| Conv_ipo_surface_block_3_edge2_2 | 0 |
| killed_elements | 1 |
| space_area | 0.155955 |
| Tmax | 386.498 841.553 |
| total_dead_elements | 87 |

Region::execute() time for AriaRegion: 3.537e+00 sec.

### 27.3.1 Current Temporal Information

The first part of the above section deals with the time step and temporal information about the running model. This section can be seen below:

Minimum Time Step Selection:

<table>
<thead>
<tr>
<th>min</th>
<th>max</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.000e-02</td>
<td>5.287e+00</td>
</tr>
</tbody>
</table>

Transient Time_Block_2: dt = 5.287

Advance AriaRegion, time 2198, time step 5.287

Aria determined the current time step size based on factors from the previous time step. Here, the maximum allowed time step size is 5.287, which is based on the minimum time step allowed for the chemistry...
model. It can also be seen that the run is currently in Time_Block_2 as is set up in Solution Control.

The information contained within the dotted lines allows one to gauge how far along a simulation is. In that one line, one can find the current time block, the iteration number, current simulation time, current time step, and the percent completion of the current time block.

### 27.3.2 Memory Reporting

One factor that determines the efficiency of a simulation is the amount of memory that is utilized. Each Sierra application will periodically output the current memory utilization - an example of this is shown below:

Memory Usage: current = 209186816 (199.5 M), high-water-mark = 217268224 (207.2 M)

One can see that at this time, 199.5 MB is currently being utilized, with a historical maximum of 207.2 MB during the course of the simulation.

### 27.3.3 Linear/Nonlinear Solves

The next part of the runtime information deals with the linear and nonlinear solution:

**Equation System AriaRegion->main:**

- **Step**: Transient, Strategy: NEWTON, Time: 2.20e+03, Step: 5.29e+00
- **Matrix**: Solver: "Solve_Temperature", Unknowns: 1360, Nonzeros: 11578
- **Mesh**: Processor 0 of 1: 1253 of 1253 elems, 1363 of 1363 nodes
- **Computing View Factors for enclosure space**

<table>
<thead>
<tr>
<th>NONLINEAR</th>
<th>LINEAR</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Step</th>
<th>Resid</th>
<th>Delta</th>
<th>Itns</th>
<th>Status</th>
<th>Resid</th>
<th>Asm/Slv</th>
<th>Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>6.37e-02</td>
<td>1.10e+01</td>
<td>59</td>
<td>ok</td>
<td>8.23e-07</td>
<td>7.0e-03/2.0e-03</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>1.10e-04</td>
<td>8.18e-03</td>
<td>58</td>
<td>ok</td>
<td>1.19e-09</td>
<td>7.0e-03/2.0e-03</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>3.06e-07</td>
<td>8.81e-05</td>
<td>61</td>
<td>ok</td>
<td>3.68e-12</td>
<td>6.0e-03/2.0e-03</td>
<td></td>
</tr>
</tbody>
</table>

Termination reason: 8.80515e-05 < nonlinear_correction_tolerance(0.001)

- **FIELD**

<table>
<thead>
<tr>
<th>Field</th>
<th>min</th>
<th>@ id</th>
<th>max</th>
<th>@ id</th>
<th>max-chg</th>
<th>@ id</th>
<th>pred-err</th>
<th>pred-min</th>
<th>@ id</th>
<th>pred-max</th>
<th>@ id</th>
</tr>
</thead>
<tbody>
<tr>
<td>TEMPERATURE</td>
<td>3.8650e+02</td>
<td>412</td>
<td>8.4155e+02</td>
<td>512</td>
<td>3.93e+00</td>
<td>1443</td>
<td>3.53e-04</td>
<td>3.8650e+02</td>
<td>412</td>
<td>8.4155e+02</td>
<td>512</td>
</tr>
<tr>
<td>DT_TEMPERATURE</td>
<td>MIN(value,loc) = (-1.18e+00, 1443)</td>
<td>MAX(value,loc) = (9.68e-01, 249)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Segregated solution procedure converged after 1 iterations.

Each bullet point here gives some information as to the solution time and strategy, as well as some information about the matrix being solved. The line *Computing View Factors* ... indicates that enclosure radiation viewfactors are being solved - more information about enclosure radiation diagnostics can be found in Section 27.3.6.

Each column of the NONLINEAR/LINEAR table deserves some explanation.

- **Step** The nonlinear step being taken - here there are three nonlinear steps in the timestep.
Resid  Nonlinear residual at the end of each nonlinear step and before the linear steps are taken
Delta  Difference in the solution variable(s) compared to the last nonlinear solution
Itns  Number of linear iterations per nonlinear step
Status  Whether linear solver completed successfully (output is either ok or fail)
Resid  Linear residual at the end of the linear iterations

A more complete description of this table is included in the Nonlinear Solution Specifications chapter refer:nlsolv.

As an example of how to interpret the output from this table, nonlinear step 2 will be translated to text:
“During nonlinear step 2, the iterative linear solver took 7.0ms to assemble and 2.0ms to solve the linear system. It completed the solve in 58 iterations and exited with a linear residual of 1.19e-09. The nonlinear residual at the end of the nonlinear step was 1.10e-04, which was a change of 8.18e-03 from the previous nonlinear step.” This section of the log file also shows the reason why the time step exited - usually due to reaching some desired tolerance or failure to converge. Here, the solution was reached when the change in temperature from the previous nonlinear step (Delta = 8.81e-05) was less than the nonlinear correction tolerance of 0.001. For more information on what linear and nonlinear residuals mean, refer to the Aria Theory Manual for how the equations and linear matrix are solved.

The lines below the table also show useful output on the fields being solved - here, for temperature. Here, we can see that the minimum temperature is 386.5 at node 412, and the maximum temperature is 841.6 at node 512. Information about the predictor-corrector errors are also shown, as well as the time derivative of temperature.

### 27.3.4 Timestep Determination

Based on the solution at the current step, the timestep is re-calculated. This is shown in the following lines:

```
---------- For equation system name: main ----------
Global predictor error = 3.531e-04
Time step selection: dt <= 2.326e+01 (based on Predictor-Corrector Tolerance).
Time step selection: dt <= 1.057e+01 (based on Maximum Time Step Size Ratio).
Time step selection: dt <= 6.000e+01 (based on Maximum Time Step Size).
Time step selection: dt = 5.277e+00 (Minimum Chemistry timestep block_6).
Time step selection: dt <= 1.277e+01 (based on Stability limit for dt Ratio).
Time step selection: dt = 1.057e+01 (Adaptive time stepping result).
```

These restrictions will change based on the method in which the solution is solved as well as the physics being solved. Here, the timestep may be limited by adaptive time stepping restrictions (predictor-corrector tolerance, maximum timestep ratio, and maximum time step size), chemistry timestep, and BDF2 stability limit. To put this block in sentence format, “The next timestep must be: less than or equal to 23.3s based on the predictor-corrector tolerance, less than or equal to 10.57s based on the maximum timestep ratio, less than or equal to 60s based on the maximum time step size, equal to 5.28s based on the ChemEQ solution, and less than or equal to 12.77s based on timestep stability.” After looking at it in this manner, one can see that the Chemistry timestep of 5.28s is the limiting timestep, and this will be the timestep size which is used in the next timestep, similar to what was explained in Section 27.3.1.
27.3.5 Global Variable Output

At the end of each timestep, a summary is given of the global variables in the model. If there are global variables that the user does not wish to output the "GLOBAL VARIABLES EXCLUDED FROM LOG FILE = [list of variables]" input deck line may be used.

Killed 1 elements this timestep, for a total of 87 dead.

<table>
<thead>
<tr>
<th>Global Variable</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conv_ipo</td>
<td>189.224</td>
</tr>
<tr>
<td>Conv_ipo_surface_block_2_edge2_2</td>
<td>0</td>
</tr>
<tr>
<td>Conv_ipo_surface_block_3_edge2_2</td>
<td>0</td>
</tr>
<tr>
<td>killed_elements</td>
<td>1</td>
</tr>
<tr>
<td>space_area</td>
<td>0.155955</td>
</tr>
<tr>
<td>Tmax</td>
<td>386.498 841.553</td>
</tr>
<tr>
<td>total_dead_elements</td>
<td>87</td>
</tr>
</tbody>
</table>

Region::execute() time for AriaRegion: 3.537e+00 sec.

This section is useful for run-time monitoring of key values, as well as how long the execution time was, in order to locate any slow-running parts of the model through time.

27.3.6 Chaparral Output - Enclosure Radiation

When enclosure radiation is defined and set up in the input file, an external library called Chaparral is utilized. Chaparral output is not automatically sent to the Sierra log file, but rather is sent to standard output (on Sandia HPC platforms, the slurm output file). Chaparral will output useful information regarding the viewfactor calculation, matrix smoothing, and the radiosity solver. Each section below will describe the output one may expect from Chaparral.

Instantiation

This output only occurs at the very start of the Chaparral output, but can help a user remember the number of processors used and the number of enclosures in the model:

*********************************************************
CHAPARRAL -- Version 3.3.1 development -- unreleased
*********************************************************

Initializing for:
number of processors = 1
number of enclosures = 1
max number of patches = 1000

Viewfactor Calculation

At the beginning of the viewfactor calculation, a banner will be displayed which shows the current simulation step and time, which is useful in correlating the Chaparral output to Aria output, as well as an estimate of the rowsum for each enclosure:
The next section actually deals with the calculation of the radiation enclosure viewfactors:

Calculating viewfactors for enclosure <space>
enclosure geometry: axisymmetric
enclosure type: partial (area=100), blocking
# of rotations: 16
# of patches: 62
# of facets: 976
# of nodes: 63
spatial tolerance: 1e-06
BSP target max depth: 10
BSP target min length: 50
BSP Tree num leafs: 28
BSP Tree max depth: 5
BSP Tree min length: 27
BSP Tree max length: 45
output level: 2

Data segment memory size = 0.00Mb

Calling VF_Hemicube()...
resolution = 400
max subdivisions = 4
min separation = 5

Minimum Partial Enclosure Area = 0.151296

Minimum effective surface radius = 0.00916667
Minimum separation distance = 0.00558985
Maximum desired surface subdivision = 9, 46
Actual maximum surface subdivision = 4

Data segment memory size = 0.00Mb

Elapsed time = 3.92

While some of this output is merely a summary of user-specified or default tolerances, some information is useful during the simulation or for debugging. First, the user-defined name of the enclosure is shown in < > brackets (i.e., <space>). The number of facets and nodes that participate in the enclosure can also be useful in determining if the enclosure is complete in some instances. Last, if one is specifying a partial enclosure, then the Minimum Partial Enclosure Area is useful in correct specification of the corresponding parameter in the input file.
For simulations in which the enclosure does not change through time, the viewfactor calculation and corresponding text output will only appear once per enclosure after instantiation.

**Viewfactor Smoothing**

If viewfactor smoothing is specified, information about this step is also output. An example is shown below:

```
 ******************************************************************
 V I E W F A C T O R M A T R I X S M O O T H I N G
 ******************************************************************

   Smoothing of viewfactor matrix for enclosure <space>
   PCG Solver, wt = 2.0
   Max iterations = 500
   Tolerance    = 1e-07

   Enforcing reciprocity by addition...
   Nonzero lower triangular entries = 688 changed to 688
   Nonzero upper triangular entries = 687 changed to 688
   Elapsed time          = 0.00
   Number of passes     = 1
   Number of iterations = 26
   Elapsed time          = 0.00

 Again, the enclosure name is denoted within the < > brackets, with another summary of defined values. This section is useful for debugging problematic enclosures, since viewfactor smoothing can fail when the enclosure is ill-defined.

**Viewfactor Summary**

Once the viewfactors have been computed and smoothing has been applied, Chaparral will output the summary of how well the operations have done. This is shown in the Viewfactor Matrix Summary:

```
 ******************************************************************
 V I E W F A C T O R M A T R I X S U M M A R Y
 ******************************************************************

 Viewfactor matrix summary for enclosure <space>

 Target rowsum  = 62
 Raw rowsum total = 62.000000
 Raw rowsum error min = -1.1479e-07
 Raw rowsum error max = 1.1025e-07
 Raw rowsum error mean = 4.9383e-09 +/- 4.9314e-08
 Smoothed rowsum total = 62.000000
 Smoothed rowsum error min = -7.0897e-08
 Smoothed rowsum error max = 5.9255e-08
 Smoothed rowsum error mean = 2.8875e-10 +/- 2.7300e-14

 Viewfactor matrix is 36.19% dense
```
Total elapsed time = 3.93 (sec)
  (Initialization) = 0.01
  (Calculation) = 3.92
  (Smoothing) = 0.00

Data segment memory size = 0.00Mb

Here one can see that for the space enclosure, the computed rowsum closely matched the target rowsum, with and without smoothing. If one is also concerned about timing, an itemized timing summary is given for the viewfactor matrix setup. This is probably the most useful section to look at if one is concerned about issues with certain enclosures, as well as judging the cost and usefulness of viewfactor smoothing.

Radiosity Solution

After the viewfactor matrix has been populated, Chaparral will use this to compute the radiosity matrix. Output for one iteration of the radiosity matrix solver is shown below:

***************************************************************
R A D I O S I T Y S O L V E S
Step = 214  Time = 2425.53  Time_Step = 6.18025  Iteration = 1
***************************************************************

***************************************************************
R A D I O S I T Y M A T R I X S O L V E R
***************************************************************

Solving radiosity equations with GMRES for enclosureID <space>

Initial residual = 1.553372e+03  tol = 2.000000e-07
  iter:  0  residual = 1.553372e+03
  iter:  1  residual = 2.657621e+04
  iter:  2  residual = 6.553812e+03
  iter:  3  residual = 1.226445e+03
  iter:  4  residual = 3.526103e+02
  iter:  5  residual = 8.437261e+01
  iter:  6  residual = 3.212877e+01
  iter:  7  residual = 8.262992e+00
  iter:  8  residual = 2.397060e+00
  iter:  9  residual = 3.692025e-01
  iter: 10  residual = 1.010501e-01
  iter: 11  residual = 3.528043e-02
  iter: 12  residual = 6.867475e-03
  iter: 13  residual = 1.206750e-03
  iter: 14  residual = 9.768426e-05
  iter: 15  residual = 1.063625e-05
  iter: 16  residual = 1.902986e-06
  iter: 17  residual = 3.433117e-07
  iter: 18  residual = 9.218341e-08

Real residual is  1.48685e-09
Computing final fluxes
This output is somewhat similar to the Aria run-time output, in that the linear iteration count is given along with the linear residual, though the Chaparral output is more detailed, as it gives the linear residual at each linear step instead of a summary. Minimum and maximum values of emissivity and temperature in the enclosure can also help in understanding the behavior of the radiosity solve through time.

27.4 Run Summary

At the end of a completed run, a summary is printed to the log file, which starts with something along the lines of the following:

Procedure AriaProcedure complete May 14 2014 11:19:44
+----+----+----+----+----+----+----+----+----+----+----+----+----+----+----+----+
Execution complete

The different sections of the run summary will be described below.

27.4.1 Timing

The Sierra code as well as Aria contain many timers which encapsulate certain operations of each simulation. The timing summary shows the results of these timers and are organized so that aspects of the model that take more time can be found. A sample timing summary can be seen below and will be further explained:

<table>
<thead>
<tr>
<th>Timer</th>
<th>Count</th>
<th>CPU Time</th>
<th>Wall Time</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sierra</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Procedure AriaProcedure</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Initialize</td>
<td>1</td>
<td>0.017 (&lt;0.01%)</td>
<td>0.017 (&lt;0.01%)</td>
</tr>
<tr>
<td>Execute</td>
<td>1</td>
<td>15:13.010 (99.85%)</td>
<td>15:17.659 (99.85%)</td>
</tr>
<tr>
<td>Mesh input</td>
<td>1</td>
<td>0.168 ( 0.02%)</td>
<td>0.170 ( 0.02%)</td>
</tr>
<tr>
<td>Mesh output</td>
<td>357</td>
<td>0.905 ( 0.10%)</td>
<td>1.196 ( 0.13%)</td>
</tr>
<tr>
<td><strong>Region AriaRegion</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Initialize</td>
<td>1</td>
<td>0.015 (&lt;0.01%)</td>
<td>0.017 (&lt;0.01%)</td>
</tr>
<tr>
<td>Execute</td>
<td>357</td>
<td>15:11.364 (99.67%)</td>
<td>15:15.619 (99.62%)</td>
</tr>
<tr>
<td>Nonlinear Iteration</td>
<td>997</td>
<td>16.482 ( 1.80%)</td>
<td>16.568 ( 1.80%)</td>
</tr>
<tr>
<td>Preprocessing</td>
<td>997</td>
<td>6.811 ( 0.74%)</td>
<td>6.865 ( 0.75%)</td>
</tr>
<tr>
<td>Chemistry Kinetics</td>
<td>997</td>
<td>6.768 ( 0.74%)</td>
<td>6.820 ( 0.74%)</td>
</tr>
<tr>
<td>Category</td>
<td>Count</td>
<td>CPU Time</td>
<td>Wall Time</td>
</tr>
<tr>
<td>-------------------</td>
<td>-------</td>
<td>----------</td>
<td>-----------</td>
</tr>
<tr>
<td>LinSys Assembly</td>
<td>997</td>
<td>6.091 (0.67%)</td>
<td>6.119 (0.67%)</td>
</tr>
<tr>
<td>LinSys Solve</td>
<td>997</td>
<td>2.458 (0.27%)</td>
<td>2.472 (0.27%)</td>
</tr>
<tr>
<td>LinSys Scatter</td>
<td>997</td>
<td>0.276 (0.03%)</td>
<td>0.273 (0.03%)</td>
</tr>
<tr>
<td>Nonlinear Utilities</td>
<td>3988</td>
<td>0.668 (0.07%)</td>
<td>0.669 (0.07%)</td>
</tr>
<tr>
<td>Postprocessing</td>
<td>358</td>
<td>0.034 (&lt;0.01%)</td>
<td>0.046 (&lt;0.01%)</td>
</tr>
<tr>
<td>Utilities</td>
<td>2144</td>
<td>14:51.790 (97.53%)</td>
<td>14:54.651 (97.34%)</td>
</tr>
<tr>
<td>Parallel Sync</td>
<td>5776</td>
<td>0.012 (&lt;0.01%)</td>
<td>0.010 (&lt;0.01%)</td>
</tr>
<tr>
<td>Chaparral Viewfactor</td>
<td>357</td>
<td>14:51.546 (97.50%)</td>
<td>14:54.417 (97.32%)</td>
</tr>
<tr>
<td>Chaparral Radiosity</td>
<td>997</td>
<td>0.629 (0.07%)</td>
<td>0.623 (0.07%)</td>
</tr>
<tr>
<td>Chemistry Utilities</td>
<td>3062</td>
<td>0.085 (&lt;0.01%)</td>
<td>0.079 (&lt;0.01%)</td>
</tr>
<tr>
<td>Contact Search</td>
<td>357</td>
<td>0.003 (&lt;0.01%)</td>
<td>0.000 (&lt;0.01%)</td>
</tr>
<tr>
<td></td>
<td>1074</td>
<td>0.259 (0.03%)</td>
<td>0.372 (0.04%)</td>
</tr>
<tr>
<td>Tmax</td>
<td>358</td>
<td>0.275 (0.03%)</td>
<td>0.277 (0.03%)</td>
</tr>
<tr>
<td>Mesh input</td>
<td>1</td>
<td>0.168 (0.02%)</td>
<td>0.170 (0.02%)</td>
</tr>
<tr>
<td>Mesh output</td>
<td>357</td>
<td>0.904 (0.10%)</td>
<td>1.196 (0.13%)</td>
</tr>
<tr>
<td>Results output</td>
<td>358</td>
<td>0.326 (0.04%)</td>
<td>0.420 (0.05%)</td>
</tr>
<tr>
<td>Restart output</td>
<td>358</td>
<td>0.549 (0.06%)</td>
<td>0.729 (0.08%)</td>
</tr>
<tr>
<td>History output</td>
<td>358</td>
<td>0.015 (&lt;0.01%)</td>
<td>0.027 (&lt;0.01%)</td>
</tr>
<tr>
<td>Heartbeat output</td>
<td>358</td>
<td>0.020 (&lt;0.01%)</td>
<td>0.036 (&lt;0.01%)</td>
</tr>
<tr>
<td>Control</td>
<td>109</td>
<td>0.001 (&lt;0.01%)</td>
<td>0.005 (&lt;0.01%)</td>
</tr>
<tr>
<td>Vis output</td>
<td>358</td>
<td>0.001 (&lt;0.01%)</td>
<td>0.000 (&lt;0.01%)</td>
</tr>
<tr>
<td>Perf: RunSierra</td>
<td>1</td>
<td>1.218 (0.13%)</td>
<td>1.225 (0.13%)</td>
</tr>
<tr>
<td>Perf: RunSierra::parse</td>
<td>1</td>
<td>1.218 (0.13%)</td>
<td>1.225 (0.13%)</td>
</tr>
<tr>
<td>Perf: RunSierra::Domain::execute</td>
<td>1</td>
<td>15:13.197 (99.87%)</td>
<td>15:17.846 (99.87%)</td>
</tr>
</tbody>
</table>

Each entry shows the CPU and Wall Times associated with it, along with a percentage, which is the percentage of the total run time. One can also see where a timer resides with respect to the Procedure, Region, etc. by the level of indentation in the timing summary. For example, one can see by the indentation that Nonlinear Iteration is a part of Execute, which resides in the Region ariaRegion. The sum of timers of all entries on a certain indentation level should be less than or equal to the timer of the parent entry, since the parent timer is a wrapper around the children timers and the corresponding code execution. For example, Preprocessing, LinSys Assembly, LinSys Solve, LinSys Scatter, and Nonlinear Utilities timers are equal to 16.304s of CPU time, which is less than the 16.482s of Nonlinear Iteration.

Here is a general description of what each timer means:

- **Sierra** Complete encapsulation of the simulation
- **Procedure** Procedure-level of the simulation, including Solution Control and Regions
- **Initialize** Initialization of the Procedure
- **Execute** Execution time for the procedure; in a simple one-region simulation, this will be very similar to Region Execute
- **Mesh Input** Time to read in the input Genesis mesh
- **Mesh Output** Time required to output any results, e.g., Results, Restart, History, Heartbeat Output
- **Region** Region scope of the simulation, including physics setup
- **Initialize** Initialization of the Region
- **Execute** Execution time for the Region
- **Nonlinear Iteration** Performance of the nonlinear solves
Preprocessing Setup for the nonlinear iterations

Chemistry Kinetics Setup of chemistry volume source and ODE solver

LinSys Assembly Assembly of the Linear system of matrices

LinSys Solve Solution of the linear system of matrices

LinSys Scatter Division of the linear system of matrices in parallel

Nonlinear Utilities A combination of several utilities (e.g., chemistry, radiation) that occur during the nonlinear step

Postprocessing Execution of user-requested post-processing fields

Utilities Wrapper around utilities required for system of equations

Parallel Sync Communication required between processors for simulations run in parallel

Chaparral Viewfactor Viewfactor calculation for enclosure radiation

Chaparral Radiosity Radiosity calculation for enclosure radiation

Chemistry Utilities Chemistry solve which occurs outside of the nonlinear loop

Contact Search Search required to find contact interactions

Tmax Determination via Encore of Tmax postprocessor

Mesh input Time to read in the Genesis mesh

Mesh output Time to output results

Results output Time to output Results Output block(s)

Restart output Time to output Restart Output block(s)

History output Time to output History Output block(s)

Heartbeat output Time to output Heartbeat Output block(s)

Control Determination of when to output

Vis output Time to output in-situ visualization output

Perf: Sierra performance-related timers

It can be seen that in this simulation, the bulk of the time is spent in Chaparral Viewfactor. This is due to the fact that this simulation involves element death, and the radiation enclosure is updated each time that elements are killed.

27.4.2 Memory

In addition to the run-time memory statistics, there is a memory summary printed at the end of the log file. An example of this is shown below:
The output of this memory summary becomes more varied for a parallel simulation, where the model may not be evenly distributed amongst processors.

Another useful piece of information related to memory reporting is contained in the Performance metric summary:

<table>
<thead>
<tr>
<th>Metric</th>
<th>Largest Processor</th>
<th>Smallest Processor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dynamically Allocated (Heap)</td>
<td>73.8 MB on 0 at 13:33.348</td>
<td>73.8 MB on 0 at 13:33.348</td>
</tr>
<tr>
<td>Largest Free Fragment (Heap)</td>
<td>43.0 MB on 0 at 13:33.348</td>
<td>43.0 MB on 0 at 13:33.348</td>
</tr>
<tr>
<td>Total Memory In Use</td>
<td>398.1 MB on 0</td>
<td>398.1 MB on 0</td>
</tr>
<tr>
<td>Major Page Faults</td>
<td>0 flts on 0</td>
<td>0 flts on 0</td>
</tr>
</tbody>
</table>

27.5 Summary

Though it can be difficult to interpret the sometimes large amounts of data that are output via log files, this information can be useful in ascertaining a simulation’s performance and can also be used to improve the manner in which it is run. Though some of the information presented here is Aria-specific, much of it is also applicable when using other Sierra applications and can aid in use of those codes as well.
Chapter 28

Diagnostic Output

28.1 About Diagnostic Streams

Aria is instrumented with a diagnostic output capability that can be very useful when you’re debugging problems. You can tell Aria to write additional information to the log file by enabling different *print masks*. The list of available print masks is summarized in table 28.1; this list is also supplied in the Aria runtime help which is printed if you add `-h` to your `sierra` command line.

There are two ways to activate a print mask for diagnostic printing. The first is to use the `-arialog` command line option. When you’re using the `sierra` script this is a little odd because you have to pass that option as an option to the script. By example, if you want to enable the sensitivity checker you can do that like this

```
sierra aria ...-arialog sens_check
```

where `...` all of the other command line options you typically use.

The second way to enable a print mask is more versatile and is done through your aria input file. At the highest level of the input file – somewhere in the `BEGIN SIERRA` block – you can add a `DIAGNOSTIC CONTROL` block. This enables you to specify intervals in time or time step number over which print masks should be enabled. As a simple example, let’s say you want to enable the sensitivity checker (see `sens_check` in 28.1) around time step number 47. Here’s an example input file snippet that allows you to do that:

```
Begin Diagnostic Control arialog
   From Step 46 to 48 enable sens_check
End
```

With this, the sensitivity checker will only be active for those three time steps (it’s also valid to say `From Step 47 to 47` ...). The end of this chapter includes a detailed command reference for that input.

Lastly, I’ll just mention here for now that there are additional capabilities for logging, such as per-processor log files (be default only processor zero logs output). If you need more advanced diagnostics contact the Aria developers for help.

28.2 Diagnostic Output Command Reference

28.3 Diagnostic Control

*Scope:* Sierra
<table>
<thead>
<tr>
<th>Print Mask</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>bc</td>
<td>Display boundary condition information</td>
</tr>
<tr>
<td>debug</td>
<td>Display debug diagnostic information</td>
</tr>
<tr>
<td>eq</td>
<td>Display equation information</td>
</tr>
<tr>
<td>expression</td>
<td>Display expression information</td>
</tr>
<tr>
<td>hadapt</td>
<td>Display h-adapt diagnostic information</td>
</tr>
<tr>
<td>finite_check</td>
<td>Display warning messages about any non-finite expression values or sensitivities</td>
</tr>
<tr>
<td>nonlinear</td>
<td>Display nonlinear solver information</td>
</tr>
<tr>
<td>pp</td>
<td>Display postprocessor diagnostic information</td>
</tr>
<tr>
<td>sens_check</td>
<td>Display messages generated by expression sensitivity checker</td>
</tr>
<tr>
<td>fast_sens_check</td>
<td>Display messages generated by sensitivity checker (skipping FAD, element, and kernel expressions)</td>
</tr>
<tr>
<td>species</td>
<td>Display species information</td>
</tr>
<tr>
<td>transfer</td>
<td>Display transfer information</td>
</tr>
<tr>
<td>dualsolve</td>
<td>Display dual solve information</td>
</tr>
<tr>
<td>plugin</td>
<td>Display plugin information</td>
</tr>
<tr>
<td>chaparral</td>
<td>Display chaparral information</td>
</tr>
<tr>
<td>utility</td>
<td>Display equation system utility information</td>
</tr>
</tbody>
</table>

Table 28.1. Aria diagnostic output print masks.

Begin Diagnostic Control Name

Enable Printmask
From Step StartStep To EndStep Enable Printmask
From Time StartTime To EndTime Enable Printmask
On Condition ConditionList Enable Printmask
Set Information Stream Path File_path
End

Summary Specifies the diagnostic writer to set output control features.

Description The diagnostic and informational output can be selectively enabled based on time, step or an application specified condition. During the application’s procedure execution loop, the diagnostic controller evaluates the enclosed line commands in the order specified in the input deck. The diagnostic options specified in the first line command that meets its criteria are applied.

Since control parameters are only applied when the criteria is met, it is important to include an ENABLE line command with the base settings to be applied as a baseline. Refer the the '-h' output for a complete list of diagnostic writers and available values.

fmwkout Specify a comma separated list of: comm Display mpi communication diagnostic information contact Display contact diagnostic information coverage Collect and display traceable function usage coverage dump-load Dump domain after mesh load dump-setup Dump domain after setup error Display error messages field Display field registration information geometry Display geometry diagnostic information input-check Check input deck and mesh load, does not execute material Display material diagnostic information members Display data structure members messages mesh-mod Display mesh modification diagnostic
information no-execute Do not execute no-load Do not load mesh parameters Display parameter diagnostic information plugins Display user function and plugin diagnostic information pretty-print Pretty print input deck and mesh data scontrol Display solver control diagnostic information search Display search diagnostic information solver Display linear and non-linear solver diagnostic information syntax-check Check syntax of input deck, does not load mesh or execute trace Display execution trace trace-stats Display execution time and memory usage during trace transfer Display transfer diagnostic information verbose Dump domain after setup, does not load mesh or execute warning Display warning messages

iossdebug Specify a comma separated list of: coverage Collect and display traceable function usage coverage error Display error messages members Display data structure members messages mesh Display mesh I/O diagnostic information restart Display restart I/O diagnostic information results Display results I/O diagnostic information serialized Display serialized I/O information trace Display execution trace trace-stats Display execution time and memory usage during trace warning Display warning messages

prsrdebug Specify a comma separated list of: coverage Collect and display traceable function usage coverage error Display error messages members Display data structure members messages parser Display parser diagnostic information trace Display execution trace trace-stats Display execution time and memory usage during trace warning Display warning messages

The options which may be enabled varies for each application, diagnostic and information writer. To obtain a list of available options, use the sierra app -i -O -h command.

See Diagnostic Stream for specifying the output destination.

### 28.3.1 Enable

**Scope:** Diagnostic Control

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Printmask</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Specifies the options to enable when no other control option criteria are satisfied.

### 28.3.2 From Step

**Scope:** Diagnostic Control

```
From Step StartStep To EndStep Enable Printmask
```

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>StartStep</td>
<td>integer</td>
<td>undefined</td>
</tr>
<tr>
<td>EndStep</td>
<td>integer</td>
<td>undefined</td>
</tr>
<tr>
<td>Printmask</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Specifies the options to enable when the step is within the specified range.

### 28.3.3 From Time

**Scope:** Diagnostic Control
### 28.3.4 On Condition

**Scope:** Diagnostic Control

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>ConditionList</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Printmask</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary** Specifies the options to enable when the condition mask matches.

### 28.3.5 Set Information Stream Path

**Scope:** Diagnostic Control

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>File_path</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary** File path to information stream information to.

### 28.3.6 Diagnostic Stream

**Scope:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>File_name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary** File path to write diagnostic messages to.

### 28.3.7 Enable

**Scope:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Printmask</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Summary: Specifies the options to enable when no other control option criteria are satisfied.

28.3.8 From Step
Scope:

From Step StartStep To EndStep Enable Printmask

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>StartStep</td>
<td>integer</td>
<td>undefined</td>
</tr>
<tr>
<td>EndStep</td>
<td>integer</td>
<td>undefined</td>
</tr>
<tr>
<td>Printmask</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies the options to enable when the step is within the specified range.

28.3.9 From Time
Scope:

From Time StartTime To EndTime Enable Printmask

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>StartTime</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>EndTime</td>
<td>real</td>
<td>undefined</td>
</tr>
<tr>
<td>Printmask</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies the options to enable when the time is within the specified range.

28.3.10 On Condition
Scope:

On Condition ConditionList Enable Printmask

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>ConditionList</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Printmask</td>
<td>&quot;string&quot;</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies the options to enable when the condition mask matches.

28.3.11 Set Information Stream Path
Scope:

Set Information Stream Path File_path

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>File_path</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: File path to information stream information to.
Chapter 29

Level Set Reference

29.1 Level Set Overview

Aria employs a level set capability supported by the Krino library. Here the basic Krino capability is used to define the level set interface and the problem physics is handled with equations native to Aria.

To use this capability one would define Level_Set for the DOF on a particular volume mesh entity (block) as described in 6.5. Then the level set interface would be defined using the LEVEL SET INTERFACE commands. In tracking the level set interface the Narrow Band Width is the distance over which the level set distance is calculated. Outside this narrow band, the distance is clipped when the redistancing calculation is performed. Advection algorithms in Aria apply over the entire domain, but the redistancing algorithms in Krino support this clipping. Tailoring the Narrow Band Width to one’s problem can help reduce the cost of redistancing and reduce solution artifacts away from the interface.

29.2 Level Set Interface

Scope: Aria Region

```
Begin Level Set Interface InterfaceName
  Composite Name {=|are|is} Variable
  Distance Variable {=|are|is} Variable
  Extension Velocity {=|are|is} Value...
  Initial Offset Distance {=|are|is} Offset
  Narrow Band Element Size Multiplier {=|are|is} Value
  Narrow Band Width {=|are|is} Value
  Perform Initial Redistance {=|are|is} performInitialRedistance
  Redistance Method {=|are|is} RedistanceMethodType
  Reinitialize Every Step
  Begin Analytic Initial Condition BlockName
  End

  Begin Block Interface Initial Condition BlockName
  End

  Begin Compute_Surface_Distance SurfaceName
  End

  Begin Motion Specification Motion

End
```
Summary: Main block for krino level set interface control mechanics.

Description: This is the main block for the level set interface control mechanics. This block specifies a level set interface that is integrated in time using an extension velocity. The plan is that the physics application code will see only this mechanics, although there may be several other supporting mechanics that are plugged in.

29.2.1 Composite Name

Scope: Level Set Interface

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Composite Name</td>
<td>Variable</td>
<td></td>
</tr>
</tbody>
</table>

Summary: Optionally specify a composite level set name which may be used across multiple level sets to define composite phases.

29.2.2 Distance Variable

Scope: Level Set Interface

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Distance Variable</td>
<td>Variable</td>
<td></td>
</tr>
</tbody>
</table>

Summary: Specify the distance variable name.

29.2.3 Extension Velocity

Scope: Level Set Interface

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Extension Velocity</td>
<td>Value</td>
<td></td>
</tr>
</tbody>
</table>

Summary: Specify the extension velocity used to evolve the level set field. Default is 0 0 0.
29.2.4 Initial Offset Distance

Scope: Level Set Interface

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Offset</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify an offset distance to be applied to the initial level set. Allows the user to grow or shrink the initialized level set.

29.2.5 Narrow Band Element Size Multiplier

Scope: Level Set Interface

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify the width of the narrow band width used for the level set field as the given multiplier times the maximum element length scale. If this command is omitted, the distance will be computed throughout the entire domain, but this is not generally scalable in terms of memory or cpu time.

29.2.6 Narrow Band Width

Scope: Level Set Interface

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify the width of the narrow band. This form is deprecated. Use NARROW BAND ELEMENT SIZE MULTIPLIER instead.

29.2.7 Perform Initial Redistance

Scope: Level Set Interface

Summary: Perform redistancing operation just after initial conditions are applied.

29.2.8 Redistance Method

Scope: Level Set Interface

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>RedistanceMethodType</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Parameter | Value | Default
---|---|---
RedistanceMethodType | \{closest\_point|fast\_marching\} | CLOSEST\_POINT

Summary: Specify the algorithm to be used for redistancing.

### 29.2.9 Reinitialize Every Step

**Scope:** Level Set Interface

Summary: Specify that this level set interface should be re-initialized every time step. For example this can be combined with an analytic mesh surface initial condition that is moving due to applied displacements to move the CDFEM interface location with the mesh surface.

### 29.2.10 Narrow Band Width

**Scope:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
</table>
| Narrow Band Width | \{=|are|is\} | Value
| Value | real | undefined |

Summary: Specify the width of the narrow band. This form is deprecated. Use NARROW BAND ELEMENT SIZE MULTIPLIER instead.

### 29.3 Level Set Initial Conditions

As previously stated one can use the Aria initial condition definitions described in 8 by using Level_Set as the DOF.

 Alternatively, one can also use the the level set initial condition support provided by the Krino library. In a multi-region analyses where one has a level set region whose level result is being transferred to another region which controls output of the results, synchronization of the initial state may become an issue. In this event the initial conditions being transferred from the Krino region can be overwritten by the region controlling output. This problem can be dealt with by orchestration of initialization by Solution Control using the command structure included below wherein LS\_Region is the level set region.

If one chooses to use the initial conditions from within Aria instead, the command line for the Initialize command block is not used and the "Use Initialize" command line would not appear in the Solution Control "System" command block.

```plaintext
Begin Solution Control Description
    Use System Main
    
    Begin Initialize The_Init_Block
        Advance LS\_Region
        Event LS\_Region\_REINITIALIZE\_LEVELSETS
    End
```

810
Begin System Main
  Use Initialize The_Init_Block
  
  .

End System Main
  
  .

End Solution Control Description

29.4 Level Set Interface

Scope: Aria Region

<table>
<thead>
<tr>
<th>Begin Level Set Interface InterfaceName</th>
</tr>
</thead>
<tbody>
<tr>
<td>Composite Name {=</td>
</tr>
<tr>
<td>Distance Variable {=</td>
</tr>
<tr>
<td>Extension Velocity {=</td>
</tr>
<tr>
<td>Initial Offset Distance {=</td>
</tr>
<tr>
<td>Narrow Band Element Size Multiplier {=</td>
</tr>
<tr>
<td>Narrow Band Width {=</td>
</tr>
<tr>
<td>Perform Initial Redistance {=</td>
</tr>
<tr>
<td>Redistance Method {=</td>
</tr>
<tr>
<td>Reinitialize Every Step</td>
</tr>
<tr>
<td>Begin Analytic Initial Condition BlockName</td>
</tr>
<tr>
<td>End</td>
</tr>
</tbody>
</table>

| Begin Block Interface Initial Condition BlockName |
| End                                                    |

| Begin Compute_Surface_Distance SurfaceName |
| End                                        |

| Begin Motion Specification Motion |
| End                                |

End

Summary      Main block for krino level set interface control mechanics.
Description  This is the main block for the level set interface control mechanics. This block specifies a level set interface that is integrated in time using an extension velocity. The plan is that the physics application code will see only this mechanics, although there may be several other supporting mechanics that are plugged in.
### 29.4.1 Composite Name

**Scope:** Level Set Interface

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Composite Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Optionally specify a composite level set name which may be used across multiple level sets to define composite phases.

### 29.4.2 Distance Variable

**Scope:** Level Set Interface

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Distance Variable</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specify the distance variable name.

### 29.4.3 Extension Velocity

**Scope:** Level Set Interface

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Extension Velocity</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specify the extension velocity used to evolve the level set field. Default is 0 0 0.

### 29.4.4 Initial Offset Distance

**Scope:** Level Set Interface

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial Offset Distance</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specify an offset distance to be applied to the initial level set. Allows the user to grow or shrink the initialized level set.

### 29.4.5 Narrow Band Element Size Multiplier

**Scope:** Level Set Interface
29.4.6 Narrow Band Width

Scope: Level Set Interface

Summary Specify the width of the narrow band. This form is deprecated. Use NARROW BAND ELEMENT SIZE MULTIPLIER instead.

29.4.7 Perform Initial Redistance

Scope: Level Set Interface

Summary Perform redistancing operation just after initial conditions are applied.

29.4.8 Redistance Method

Scope: Level Set Interface

Summary Specify the algorithm to be used for redistancing.

29.4.9 Reinitialize Every Step

Scope: Level Set Interface

Summary Specify that this level set interface should be re-initialized every time step. For example this can be combined with an analytic mesh surface initial condition that is moving due to applied displacements to move the CDFEM interface location with the mesh surface.
29.4.10 Narrow Band Width

Scope:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify the width of the narrow band. This form is deprecated. Use NARROW BAND ELEMENT SIZE MULTIPLIER instead.
Chapter 30

Local Coordinate System Reference

30.1 Local Coordinate Systems

Aria provides utilities for convenient transformation of model specific information to the computational coordinate frame. Several common local coordinate systems - rectangular, cylindrical, spherical and conical - are pre-defined in Aria.

**Known Issue:** It has been determined that the conical local coordinate system has not been correctly implemented. Users should avoid use of this coordinate system until this has been corrected.

Figure 30.1 shows how each of the afore-mentioned local coordinate systems are constructed using the information supplied in the LOCAL COORDINATE SYSTEM command block. As a general rule, ORIGIN defines the origin of the local coordinate system, VECTOR defines the z-axis of the local coordinate system, and POINT is used to create the x-axis of the local coordinate system. The y-axis is then constructed via a cross-product of the x- and z-axes.

Given ORIGIN \( O \), VECTOR \( V \), and POINT \( P \) for any local coordinate system, internal initialization of geometric data is achieved via the following pseudo-code in 3D:

```c
CoordinateSystem::setLocalCoordinateAxes(Origin O, Vector V, Point P)
{
    e3 = unit_vector(V)  // z axis
    D1 = P - O           // vector from origin to point
    D2 = projected_vector(D1, e3) // project D1 in direction of e3
    D3 = D1 - D2         // orthogonal component of D1 to e3
    e1 = unit_vector(D3) // x axis
    e2 = cross_product(e3, e1) // y axis

    # Saved variables
    origin = O           // origin
    x_axis = e1
    y_axis = e2
    z_axis = e3
}
```

At point \( P \), a local z-axis \( e_3 \) is defined as the unit vector in the direction of vector \( V \). The local x-axis \( e_1 \) is then defined as an orthogonal projection from point \( P \) to \( e_3 \) and finally, the local y-axis \( e_2 \) is defined based on the cross product of \( e_3 \) and \( e_1 \).

Classical transformations consider the evaluation of vector or tensor components from one basis, \( e_i \), in an
alternative basis $e'_i$. By supplying parameters for the pre-defined coordinate system in the LOCAL COORDINATE SYSTEM command block one in effect defines the orientation of the local coordinate system block relative to the computational coordinate directions. Using this information any appropriate transformations can be computed internal to the code. Finally, each local coordinate system can be associated with multiple element blocks using the LOCAL COORDINATE SYSTEM line command.

Local coordinate systems are particularly useful for tensor material properties defined using their local principal values. In this case one must select a "generalized" material model in order to fully utilize the principal component values. Current support of this feature is currently limited to material diffusive coefficients - however, further support is planned.

Given any point $P$ in space at which the global value of a given tensor based material property $[K]$ is to be evaluated, the local principal axes based on the initialization described previously must be computed. This can be represented for the various local coordinate systems by the following pseudo-code:

CartesianCoordinateSystem::getPrincipalAxes(Point P, Vector e1, Vector e2, Vector e3)
{
  e1 = x_axis # principal x axis
  e2 = y_axis # principal y axis
}
e3 = z_axis          # principal z axis
}

CylindricalCoordinateSystem::getPrincipalAxes(Point P, Vector e1, Vector e2, Vector e3)
{
    e3 = z_axis          # principal z axis = same as LCS z axis
    D1 = P - origin      # vector from origin to to point
    D2 = projected_vector(D1, e3) # project D1 in direction of e3
    D3 = D1 - D2         # orthogonal component of D1 to e3
    e1 = unit_vector(D3) # principal x axis in r
    e2 = cross_product(e3, e1) # principal y axis in theta
}

SphericalCoordinateSystem::getPrincipalAxes(Point P, Vector e1, Vector e2, Vector e3)
{
    D1 = P - origin      # vector from origin to to point
    e1 = unit_vector(D1) # principal x axis in r
    e2 = cross_product(z_axis, e1) # principal y axis in theta
    e3 = cross_product(e1, e2) # principal z axis in phi
}

If the principal values \(K_x, K_y, K_z\) are provided, these correspond to principal directions \(e_1, e_2\) and \(e_3\) respectively. Given \([K]\) defined locally in the local coordinate system, the transformation matrix \([M]\) that rotates the orthogonal triplet \(\{e_1, e_2, e_3\}\) into the global coordinate space \(\{x, y, z\}\) is computed and the material property in global coordinates is defined as

\[
[K_g] = [M]^T [K] [M]
\] (30.1)

30.2 Local Coordinate System
Scope: Sierra

Begin Local Coordinate System Coordinate_System_Name
    Origin {=|are|is} Value...
    Point {=|are|is} Value...
    Type {=|are|is} CoordType
    Vector {=|are|is} Value...
End

Summary Defines the local coordinate system parameters.

Description This Command Block allows the definition of a coordinate system that can be used to transform values for output or field calculations. For any specific local coordinate system three pieces of information are required for completely defining a coordinate system are the origin, a coordinate axis vector and a point lying on one of the other axes. Using this information a transformation from the local coordinate system to the computational coordinate system can be determined.
30.2.1 Origin
Scope: Local Coordinate System

Origin {=*|are|is} Value...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Local coordinate system origin.

Description Specifies the point as the origin of a local coordinate system. The number of coordinates may be either 2 or 3 depending on the spatial dimension. By default, a value of $z=0.0$ is used for 2D.

30.2.2 Point
Scope: Local Coordinate System

Point {=*|are|is} Value...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Local coordinate alternative point.

Description Declare the point that lies on an alternate axis and is in the plane subtended by the specified VECTOR and the vector defined by the point and the origin.

30.2.3 Type
Scope: Local Coordinate System

Type {=*|are|is} CoordType

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>CoordType</td>
<td>{cartesian</td>
<td>conical</td>
</tr>
</tbody>
</table>

Summary Local coordinate system type.

Description This line command selects the coordinate system type.

30.2.4 Vector
Scope: Local Coordinate System

Vector {=*|are|is} Value...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Summary  Local coordinate axis vector.

Description  Declare the vector that defines one of the coordinate axes. The specific usage depends on the selected coordinate system type.

30.2.5 Local Coordinate System

Scope:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Local Coordinate System {=} {are} {is} \textit{Mesh Entities}</td>
<td>\textit{Mesh Entities}</td>
<td>string</td>
</tr>
</tbody>
</table>

Summary  Associate coordinate system with mesh entity.

Description  Specify the local coordinate system to be used in conjunction with given element blocks.
Chapter 31

Bulk Volume Reference

31.1 Bulk Volume Element

For some engineering problems the variation of a field variable over the problem domain can be roughly represented by some average behavior. In these cases it may be more appropriate to understand the averaged behavior rather than to resolve the behavior in fine detail. When numerically modeling these types of problems, it is convenient to devise lumped parameter models that simulate this averaged behavior. One such modeling strategy, a bulk volume model, is employed to capture bulk fluid behavior without regard to spatial resolution. Herein the bulk volume model is described within the context of a Galerkin finite element model where the terms bulk volume element and bulk node are often used synonymously.

Following standard finite element method procedures, balance equations are written for a physical region surrounding the bulk volume where the equations possibly include contributions from unknowns associated with the bulk volume. Similarly a general conservation equation is written for a bulk volume system with flux contributions provided from bounding surfaces of the finite element method discretization. Under the assumption that the volume is known in a quasi-static sense, the spatial dependence is eliminated from equation terms containing temporal derivatives. The conservation equation for the bulk volume system is defined in the same manner as for other element blocks except that it may be specified from a command block that specifically describes the bulk volume system. This balance relation provides closure for the discretized system of equations.

It is important to note that the bulk volume element interacts with the meshed model through boundary conditions and thus serves as an intervening media whose characterization has yet to be determined. Hence the methodology does not apply to problems in which the nature of interaction is known a priori. One example for which the interaction is known a priori would be convective heat transfer between two parallel surfaces of different fixed temperatures. Here heat transfer occurs directly between the two surfaces and need not be partitioned. In this case use of the bulk volume element would wrongfully introduce an additional resistance to flow of heat between the two surfaces.

Since the bulk volume degree-of-freedom is defined in terms of a time derivative, its transient evolution follows naturally. For steady problems not requiring solution converged parameter updates it is possible to solve for the bulk volume degree-of-freedom directly. Otherwise the steady-state solution for bulk volume degree-of-freedom must be obtained using a false transient simulation.

Usage of the bulk volume capability is outlined below.

Begin Sierra myJob

Begin Aria Material a_bulk_volume

properties for bulk volume

End
Begin Procedure My_Aria_Procedure

Begin Solution Control Description

  transient solution control commands

End

Begin Aria Region My_Region

Begin bulk volume bulk_volume_name
  material = a_bulk_volume
  EQ bulk_volume_name for bulk_volume_dof with P0 using mass

  bulk volume specific definitions

End

other Region level commands

End

End

End Sierra myJob

31.1.1 Boundary Conditions

At present there are two supported methods for applying boundary conditions that couple equations on a bulk volume element to volumetric equations. For heat transfer problems where the energy equation is being solved for temperature on both the volume and the bulk volume element then either a convective or a radiative boundary condition block can be coupled to the bulk volume element. For either type of boundary condition the "USE BULK ELEMENT" line command should be added to the input block, further details can be found in Chapter 33.

For other equations boundary conditions may be applied by using a line command of the form:

BC BULKNODE_FLUX FOR volume_equation ON surface_name = model_name BULK_NODE=bulk_node_name

In some cases it is desirable to couple equations being solved for different degrees of freedom on the volume and the bulk volume element. For example coupling a porous media equation for the gas phase temperature on a volume block to an overall temperature on the bulk volume element, or an equation solved for enthalpy to an equation solved for temperature. If the same degree of freedom is not present on the bulk node and the volume then the following mappings are checked for:

- Same degree of freedom (e.g. temperature, pressure, mass fraction) but without a material phase specification on the bulk node when one is present on the volume block.

- Temperature degrees of freedom on the volume will check for enthalpy in either the same material phase or no material phase on the bulk node (and vice versa for an enthalpy degree of freedom on the volume). Additional mappings can be supported where they make sense, please email sierra-help@sandia.gov if you have such a use case.
If either no valid mapping or multiple valid mappings are found the simulation will terminate during initialization.

### 31.2 Bulk Fluid Element

**Scope:** Equation System

```plaintext
Begin Bulk Fluid Element Name

Bulk Coordinates {=|are|is} Parameters...
Bulk Element Pressure {=|are|is} VolType [Parameters]...
Bulk Element Volume {=|are|is} VolType [Parameters]...

Bulk Eq EquationName For AssociatedDoF [ {of|species} SpeciesName |{in|material_phase} MaterialPhaseName ]Using ElementType With Terms...
Bulk Source For EquationName {=|are|is} SrcType [Parameters]...
Calculate Volume From Enclosing Surface
Initial Pressure {=|are|is} p
Initial Temperature {=|are|is} t
Material {=|are|is} MatName
Reinitialize Dof Name {=|are|is} value
Temperature Is Celsius
Update Bulk Volume Pressure [ Atmospheric {=|are|is} Value ]
Use Block BlockName

End
```

**Summary**
Defines a bulk fluid element that can be used by a convective flux boundary condition.

**Description**
Generally speaking, the reference temperature used in Newton’s Law of Cooling and is a known quantity because the fluid with which it is associated is modeled as an infinite reservoir. However, if the size of this reservoir is finite, then its temperature can be affected by the energy transfer across the surface in question. This situation can be modeled by the bulk fluid element, wherein the energy of the reservoir is determined by a finite volume conservation equation.

For a bulk fluid element with a name specified as `name`, the thermodynamic properties such as density, volume and temperature may be accessed via global variables. Density is accessed using `name_RHO`, volume by `name_V`, temperature at the current state by `name_T` and temperature at the old state by `name_TOld`.

### 31.2.1 Bulk Coordinates

**Scope:** Bulk Fluid Element

```plaintext
Bulk Coordinates {=|are|is} Parameters...

<table>
<thead>
<tr>
<th>Parameter Parameters</th>
<th>Value real...</th>
<th>Default undefined</th>
</tr>
</thead>
</table>

**Summary**
Specifies the physical coordinates for the bulk node.
31.2.2 Bulk Element Pressure
Scope: Bulk Fluid Element

Bulk Element Pressure \(=\) ARE|IS VolType [Parameters]...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>VolType</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Parameters</td>
<td>[string]...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies the pressure of the bulk element with type Pressure Type (CONSTANT, USER) for bulk element defined by the specifications of the parameter list.

31.2.3 Bulk Element Volume
Scope: Bulk Fluid Element

Bulk Element Volume \(=\) ARE|IS VolType [Parameters]...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>VolType</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Parameters</td>
<td>[string]...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies the volume of the bulk element with type VolType (CONSTANT, USER) for bulk element defined by the specifications of the parameter list.

31.2.4 Bulk Eq
Scope: Bulk Fluid Element

Bulk Eq EquationName For AssociatedDoF [ {of|species} SpeciesName |{in|material_phase} MaterialPhaseName ]Using ElementType With Terms...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>EquationName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>AssociatedDoF</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>SpeciesName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>MaterialPhaseName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>ElementType</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Terms</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies an equation to solve on the bulk node. User provides the degree of freedom associated with the equation, the element type, and the terms in the equation that are active.

31.2.5 Bulk Source For
Scope: Bulk Fluid Element

Bulk Source For EquationName \(=\) ARE|IS SrcType [Parameters]...
### 31.2.6 Calculate Volume From Enclosing Surface

**Scope:** Bulk Fluid Element

**Summary**
Indicates a source term of type SrcType (CONSTANT, USER) for the specified equation on the bulk element defined by the specifications of parameter list.

### 31.2.7 Initial Pressure

**Scope:** Bulk Fluid Element

**Summary**
Specifies that the volume of the bulk element is to be calculated at each timestep from an enclosing surface. The surface must be closed (although Aria currently does not currently check for this) and is defined by the extent of the flux boundary condition which uses this bulk element.

### 31.2.8 Initial Temperature

**Scope:** Bulk Fluid Element

**Summary**
Specifies the initial pressure of the bulk node.

### 31.2.9 Material

**Scope:** Bulk Fluid Element

**Summary**
Specifications that the material properties of the bulk element are defined in the material block named matName.
31.2.10  Reinitialize Dof
Scope:  Bulk Fluid Element

**Reinitialize Dof Name {=|are|is} value**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Requests reinitialization of a bulk node DOF. This will allow override of the file supplied bulk node DOF value when using an initial condition from file. Here the initial value supplied from file will be replaced by a specified value.

31.2.11  Temperature Is Celsius
Scope:  Bulk Fluid Element

Summary  Specifies the initial temperature of the bulk node is given in degrees Celsius. Internal to the code this is relevant only when performing updates of the bulk element pressure via an ideal gas law which requires an absolute temperature.

31.2.12  Update Bulk Volume Pressure
Scope:  Bulk Fluid Element

Summary  If a bulk volume is filled with an ideal gas then the pressure can be calculated at each timestep provided that the initial temperature and pressure are given. Here it is assumed that the associated volume is closed and that the volume has been defined as constant or is being computed. If the bulk volume is being computed then one must also include the CALCULATE VOLUME FROM ENCLOSING SURFACE command line.

31.2.13  Use Block
Scope:  Bulk Fluid Element

**Use Block BlockName**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>BlockName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Specifies that the given element block should be used for the bulk node. The caveat here is that there must be only one element, one node and they must all satisfy the proper mesh object topologies. This option is used only if an element block for the bulk element is already present in the mesh. If the bulk element does not exist on the input mesh, the element block will be created and will exist on the output mesh.
31.3 Closed Surface Volume

Scope: Equation System

Begin Closed Surface Volume ModelName
    Add Surface SurfaceList...
    Utility Group {=|are|is} Frequency
End

Summary: This command block defines the parameters for computation of the volume of a closed surface. The computed volume is available as a global variable with name `modelName`.

Description: Defines a means for computing the volume of a closed surface and associate it with name `modelName`.

31.3.1 Add Surface

Scope: Closed Surface Volume

Add Surface SurfaceList...

Parameter | Value | Default
--- | --- | ---
SurfaceList | string... | undefined

Summary: Adds surfaces by name, (surface_id), to a mesh extent list.

Description: This line command is used to add surfaces to a mesh extent list. In Exodus II, surfaces are specified as side sets, that have a global integer identifier. For example, side set 12 would be added by this line command using the surface name `surface_12`. Note that in SIERRA, each element of an array of strings must be separated by whitespace.

31.3.2 Utility Group

Scope: Closed Surface Volume

Utility Group {=|are|is} Frequency

Parameter | Value | Default
--- | --- | ---
Frequency | {begin_nonlinear_solve|end_nonlinear_solve | manually_run|post_accept_solution | post_iterate|post_linear_solve| post_nonlinear_solve|pre_iterate| pre_linear_solve|pre_nonlinear_solve | run_initially|run_once| run_post_linear_system_initialization| run_post_mesh_mod} | undefined

Summary: Sets the point in the execution to evaluate utility.
31.4 Restarting With Bulk Volume Element

The bulk volume element capability is supported for restarted simulations 3.10. It is worth noting that while bulk volume element is generally not part of the original mesh discretization, the element is being created internal to the code and is also being added to the results file as well as the restart file if defined. In some cases one may wish to initialize a simulation from a previous restart file that does not contain the bulk volume element. In this case provision must be made to ignore the fact that the initialization does not include the bulk volume element so that the bulk volume element can be created anew.

Input for restart jobs subsequent to the bulk volume element being added to a model should not contain the BULK NODES IGNORE RESTART command line.

31.4.1 Bulk Nodes Ignore Restart
Scope: 

Summary Causes bulk nodes to ignore the restart file.

31.5 Bulk Node Coupling
Scope: Equation System

Begin Bulk Node Coupling Name

Additional Parameter PropertyName {=|are|is} ModelName [Parameters]...
Bulk Nodes {=|are|is} Names_1 Names_2...
Couple DofName Model {=|are|is} ModelName [Parameters]...
End

Summary Defines a coupling between equations on two bulk nodes.

Description This block command allows a user to couple equations on different bulk nodes to one another. For example, the density of gas on two bulk nodes can be coupled using a flow coefficient based on the relative pressure of the two bulk nodes. The same degree of freedom must be present on both bulk nodes, and both have to be in the same equation system in order for the coupling to be possible.

31.5.1 Additional Parameter
Scope: Bulk Node Coupling

Additional Parameter PropertyName {=|are|is} ModelName [Parameters]...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>PropertyName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>ModelName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Parameters</td>
<td>[string]...</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Declare an additional parameter for the coupling models present. For example, the "K_FACTOR_FLOW" model for density or species coupling requires that a "K_FACTOR" be provided using this line command.

### 31.5.2 Bulk Nodes

**Scope:** Bulk Node Coupling

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Names</td>
<td>string_1, string_2...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** Specify the name of the bulk nodes involved in the coupling.

### 31.5.3 Couple

**Scope:** Bulk Node Coupling

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>DofName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>ModelName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Parameters</td>
<td>[string]...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary:** Specify what degree of freedom to couple between the bulk nodes, and what model to use for the coupling.
Chapter 32

Toggle Reference

32.1 Feature Toggling

For transient analysis Aria provides a mechanism for enabling and disabling certain aspects of a model like boundary conditions, contact conditions, source contributions and element block physics during selected time periods. This option is useful when modeling a time sequence of discontinuous events and the capability will be subsequently referred to as "toggling". Toggling as applied to simple electric switches means that one either opens or closes the switch (i.e. the switch is on or off). Similarly, from an analysis perspective one simply defines a specific model feature to be either active or inactive. Toggling does not imply the ability to swap in another physics model or to apply different model parameters.

Generally speaking the use of toggling is justified when a change in model physics will result in a change to the structure of the system matrix or to the structure of linear system contributions. For example, change of a flux boundary condition to a Dirichlet boundary condition requires a change to the structure of the matrix contribution since unknown values are now specified and need not be computed. Similarly, deactivation of a element block will remove degrees-of-freedom (rows and columns) from the Jacobian matrix thus reducing the number of system unknowns. Note that a change of flux boundary condition input parameters (e.g. the convective coefficient) will modify the system matrix coefficients but not affect the structure of the associated linear system matrix contributions so feature toggling is not justified in this case. Instead parameter changes in time should be accounted for through the parameter model definition.

In order to invoke this option one needs to consider its possible effect on results. Toggling of a model feature makes its usage take on a different character than expected for standard use. In particular, invocation of toggling for a simulation feature requires that the analyst prescribe the active/inactive state of the feature for the time periods of interest. Syntactically speaking the definition of feature toggling consists of two parts, the toggle block and selection of its usage.

The active/inactive state history of a feature is defined using a TOGGLE BLOCK command block. Within the Aria input file the TOGGLE BLOCK command block appears outside of the application scope, i.e. outside of the Procedure definition. Here the time period aspect of this specification is handled by mapping the desired state to a time block of the simulation. On the other hand specific usage of the toggle feature will appear within the Region scope.

For boundary conditions, contact conditions and source contributions the association between a particular model feature is prescribed by referencing the named TOGGLE BLOCK in the definition of a model feature. For native Aria model feature toggling the association with a named TOGGLE BLOCK is made by supplying an argument \( toggle = toggle\_block\_name \) to the feature (IC, BC or Source term) command line. To define toggling with Calore style input command blocks 33.1 the association between a model feature is prescribed by including a USE TOGGLE BLOCK command line within that command block.

For element block toggling (toggling of all element block physics), the association between the block and a named TOGGLE BLOCK is established via a standalone USE TOGGLE BLOCK command line within the Region scope. For toggling of the element block associated with a bulk fluid element 31.1 we note that
the corresponding element block name is BLOCK__FOR__bulk_element_name.

Toggling is supported for flux boundary conditions, contact conditions and source terms by simply zeroing out their contributions. Thus the number of DOF in the linear system will not change. Toggling of Dirichlet boundary conditions and element blocks is currently supported by removing the element DOF from the linear system.

Oftentimes one may wish to toggle off a surface boundary condition and toggle on a variant of the same boundary condition. As an example, one might replace a convective flux boundary condition with constant reference temperature with one having a reference temperature defined by a tabular function. Since the original boundary condition still exists (but it’s toggled off) it will conflict with the new boundary condition. Cases such as this will require that the second boundary condition be applied to a different surface (differently named) that overlies the surface used in the original boundary condition in order to resolve this conflict.

Toggling has been proven to be unsupported for material properties or enclosure definitions.

Usage of toggling is demonstrated in the outline below where we note that the time period within the TOGGLE BLOCK corresponds to one of the time periods defined within the Solution Control 14 command block.

Begin Sierra myJob
.
Begin Toggle Block first_toggle
  period = a_time_period_2
  state = active
End
.
Begin Toggle Block second_toggle
  period = a_time_period_3
  state = inactive
End
.
Begin Procedure My_Aria_Procedure
.
Begin Solution Control Description
  Use system main
  Begin system main
    Simulation start time = 0.0
    Simulation termination time = 3600.0
    Begin transient a_time_period_1
      advance My_Region
    End
    Begin transient a_time_period_2
      advance My_Region
    End
    Begin transient a_time_period_3
      advance My_Region
    End
  End
Begin parameters for transient a_time_period_2
  start time = 0.0
  termination time = 2.0
Begin parameters for Aria region My_Region
.

End parameters for Aria region My_REGION
End parameters for transient a_time_period_2

Begin parameters for transient a_time_period_2
start time = 2.0
termination time = 5.0
Begin parameters for Aria region My_REGION

End parameters for Aria region My_REGION
End parameters for transient a_time_period_2

End
.
Begin Aria Region My_REGION
.
Use toggle block first_toggle for block_8
.
Begin command block ablock
.
Use toggle block second_toggle
End
.
End
.
End
.
End Sierra myJob

Note that if one wishes to replace a boundary condition (the physics model) then one can do so by first adding a duplicate surface mesh entity and adding a new toggle block corresponding to usage of the new physics model.

It is important to note that within a TOGGLE BLOCK only one event (active or inactive) can be specified for the selected time period(s). Time periods other than the ones specified within the TOGGLE BLOCK command block will assume the opposite character (active/inactive). With reference to the previous outline this behavior is demonstrated by simple examples.

For the following toggle block

Begin Toggle Block my_toggle
period = a_time_period_2
state = active
End

a feature is active for a_time_period_2 and inactive for a_time_period_1 and a_time_period_3. Similarly the same behavior could be obtained using

Begin Toggle Block my_toggle
period = a_time_period_1 a_time_period_3
state = inactive
End
32.2 Toggle Block

Scope: Sierra

Begin Toggle Block ModelName

Freeze Element Block Solution State
Period {=|are|is} Period_list...
State {=|are|is} ToggleState

End

Summary
This command block defines the period(s), state and initial conditions for model feature toggling.

Description
Defines a generic entity toggling specification and associate it with name modelName. The same modelName can be used for various model features such as element blocks, boundary conditions, contact and volumetric sources.

32.2.1 Freeze Element Block Solution State

Scope: Toggle Block

Summary
Specify that the solution state remain the same when an element block has been toggled on.

Description
Specifies that the initial condition is not reset on any associated element block that has been toggled. The default behavior is to have the initial conditions reset on element blocks that have been toggled back on.

32.2.2 Period

Scope: Toggle Block

Period {=|are|is} Period_list...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Period_list</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Specify the periods over which this toggling model is to be used.

Description
Defines the time periods over which this toggling model is to be used. Note that more than one PERIOD can be appear in the command block.
32.2.3 State
Scope: Toggle Block

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>ToggleState</td>
<td>{active</td>
<td>inactive}</td>
</tr>
</tbody>
</table>

Summary Specify the state for this toggling model over the periods it is to be used.

Description Defines the toggle state for the time periods over which this toggling model is to be used. Note that more than one STATE can be appear in the command block.

32.2.4 Use Toggle Block
Scope:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>ToggleName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specification for toggling entities in the computational model based on Toggle Block parameters. When used at the region level, the list of element blocks to be toggled must be provided. Otherwise a listing of entities is not needed as the Toggle Block will be associated with the command line or the enclosing command block.
Chapter 33

Thermal Analysis Reference

33.1 General Thermal Analysis

Aria supports the solution of the energy equation for temperature as described an earlier section 5.3, following the paradigm of native Aria input. Here a user can also utilize any of the standard Aria input commands previously described in the initial conditions Chapter 8, boundary conditions Chapter 9 and volumetric source terms Chapter 11.

Previous users of the Calore code will find that many of the general input file commands for initial conditions, boundary conditions and volume sources are in fact recognized by Aria. However, there exist major differences between the two code inputs with regard to the material property definitions. Users transitioning from the Calore to Aria should review the chapter on Aria Material properties 4. While most of the property specifications differ only slightly, the most striking difference between the two command blocks is that the Aria material input requires an additional command line definition of a heat conduction model 4.15.

The Calore and Aria code code inputs also differ slightly in the nonlinear solution strategy. Generally speaking, a solution step in Aria cannot be advanced until the linear system is fully converged whereas in Calore the solution need not be converged for the step to advance. Calore and Aria also differ in the method of residual norm evaluation. Here Calore uses a residual norm scaled by the DOF whereas Aria uses an unscaled residual norm. Provisions are made in Aria to mimic the Calore code in the aforementioned behavior by use of additional input instructions, ?? and ??, described in the nonlinear solution specification chapter ??.

Aria also provides support of user subroutines for initial condition, volumetric source, boundary condition or material property. When a user subroutine is named within a standard thermal analysis command block it must be referred to as a Calore_User_Sub. Usage of input file data within these user subroutines is facilitated by use of named Data Blocks that can be made available to the user subroutines. These Data Blocks fill the role formerly occupied by Real and Int command lines in the Calore.

As previously mentioned, one may setup the solution of a thermal problem using either Calore style input commands or native Aria input commands. It is worth noting that when using the native Aria commands for heat flux, the sign convention for heat flux in the boundary condition commands for constant flux 9.2.1 and user_function flux 9.2.32 differs from that in the Calore style of heat flux in that they are of opposite sign.

Aria supports a number of non-native (Calore style) command line blocks to facilitate definition of the thermal problem to be solved. These command blocks are defined in the sections which follow. Here input specification on portions of the FEM mesh can be simplified by aggregation of the parts into Mesh Groups 6.26.
33.2 Checking Conservation

Aria has a builtin energy conservation check that can be activated for the energy equation. This is accomplished by adding the following line to whatever block you specify the energy equation in (either an Equation System block or the Region block):

Check Conservation of Energy

When this is active, after each time step the accumulated internal energy is compared with the energy added from all boundaries and sources and the level of conservation is printed in the log file.

This is a conservation check targeted towards conduction problems. The balance calculations do not currently include advection or divergence sources and will not be accurate for compressible fluids or domains with advection at the boundaries.

33.3 Initial Condition

**Scope:** Aria Region

```plaintext
Begin Initial Condition BlockName
    Add Surface PartList...
    Add Volume PartList...
    All Volumes
    Node Subroutine {=|are|is} Name
    Temperature {=|are|is} t0
    Use Data Block Name
    Use File Variable File_variable_name For Aria_variable_name [{of|species} SpeciesName |{in|material_phase} MaterialPhaseName ][ At Time Value ]
End
```

**Summary**
Allows the specification of a user-defined initial conditions on any combination of volumes, surfaces and nodes.

**Description**
There are three methods for setting the initial conditions of the thermal model. The temperature specification must be of exactly one type: e.g., constant, defined in a user subroutine, or read from a solution database file. If a user subroutine is used, real and integer data may be declared that will be local in scope to this instance of the boundary condition. This data may be accessed using a user query function.

33.3.1 Add Surface

**Scope:** Initial Condition

```plaintext
Add Surface PartList...
```

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>PartList</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>
33.3.2 Add Volume
Scope: Initial Condition

Add Volume PartList...

Parameter  Value  Default
PartList   string... undefined

Summary  Block names to which this initial condition applies. May specify a space-delimited list.

33.3.3 All Volumes
Scope: Initial Condition

Summary  Apply this command block to all volumes in the mesh file.

33.3.4 Node Subroutine
Scope: Initial Condition

Node Subroutine {=|are|is} Name

Parameter  Value  Default
Name       string  undefined

Summary  Specifies that the named user-defined subroutine be used. At most one subroutine name may be specified for a given quantity. This subroutine must conform to the "node signature" argument list type.

33.3.5 Temperature
Scope: Initial Condition

Temperature {=|are|is} t0

Parameter  Value  Default
 t0         real    undefined

Summary  Set the initial temperature to a constant value.

33.3.6 Use Data Block
Scope: Initial Condition

Use Data Block Name

839
33.3.7 Use File Variable
Scope: Initial Condition

Use File Variable File_variable_name For Aria_variable_name [ {of|species} SpeciesName |{in|material_phase} MaterialPhaseName ][ At Time Value ]

Parameter | Value | Default
---|---|---
File_variable_name | string | undefined
Aria_variable_name | string | undefined
SpeciesName | string | undefined
MaterialPhaseName | string | undefined

Summary Assign file nodal variable to valid registered nodal variable for initial condition.

Description This line command specifies that the named Aria variable be initialized from a variable contained in the mesh file that is assigned to the region. For example, this line command may be used to initialize a transient calculation from a previously performed steady-state calculation in the following way. Perform the steady-state calculation and output the temperature. Then copy that output file and use it as a mesh file for the transient calculation. Finally, use this line command to tie the variable named "file_variable_name" in the mesh file to the aria variable named "aria_variable_name". Here "aria_variable_name" must be set to nonlinear_solution->temperature.

The closest time step to the time value found the IO database will be used for the assignment. For an initial condition taken from a steady state results file, omit the optional time command. The optional command is necessary if the initial condition is coming from a file with multiple timesteps.

NOTES: 1) Current framework services do not allow restricting this initialization by mesh subset. e.g., if you want to initialize the temperature, then all nodes in the mesh will be initialized. 2) Today, Aria only supports initialization of temperature field. This restriction may be removed in the future so that chemistry variables or user-defined variables may also be initialized.

33.4 Temperature Boundary Condition
Scope: Equation System

Begin Temperature Boundary Condition BC name
Add Surface SurfaceList...
Field {=|are|is} VariableName
Integer Data Values...
Node Subroutine {=|are|is} Name
Real Data Values...
Temperature \{=|are|is\} Value
Temperature Fortran Subroutine \{=|are|is\} Name
Temperature Node Variable \{=|are|is\} Name
Temperature Scale Factor \{=|are|is\} Magnitude
Temperature Time Function \{=|are|is\} FunctionName
Use Data Block Name
Use Death Name
Use Toggle Block ToggleName [ \{@|at|for|in|on|over\} ElementBlockList... ]
Value \{=|are|is\} VariableValue

Summary   Specified temperature boundary condition.
Description For this boundary condition, the conservation equations are discarded, and the temperature
is completely determined by this boundary condition. The temperature specification must be
of exactly one type: e.g., constant, time-dependent, or defined in a user subroutine. If a user
subroutine is used, real and integer data may be declared that will be local in scope to this
instance of the boundary condition. This data may be accessed using a user query function.

33.4.1 Add Surface
Scope:   Temperature Boundary Condition

Add Surface SurfaceList...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>SurfaceList</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary   Adds surfaces, by name, to a boundary condition’s extent.
Description This line command is used to add surfaces to the extent of a boundary condition. In Exodus
II, surfaces are specified as side sets, that have a global integer identifier. For example, side
set 12 would be added by this line command using the surface name surface_12. Note that
in SIERRA, each element of an array of strings must be separated by whitespace.

33.4.2 Field
Scope:   Temperature Boundary Condition

Field \{=|are|is\} VariableName

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>VariableName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary   This command can be used with Dirichlet conditions for any solution unknown used in Aria.
The FIELD command argument defines the name of the variable to be specified with this
boundary condition.
33.4.3 Integer Data
Scope: Temperature Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>integer...</td>
<td>integer...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary List of integer data values to be used by the FORTRAN user subroutine. Copies of these values are provided to the subroutine hence changes to these values within the subroutine are not saved.

33.4.4 Node Subroutine
Scope: Temperature Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>string</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specifies that the named user-defined subroutine be used. At most one subroutine name may be specified for a given quantity. This subroutine must conform to the "node signature" argument list type.

33.4.5 Real Data
Scope: Temperature Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>real...</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary List of real data values to be used by the FORTRAN user subroutine. Copies of these values are provided to the subroutine hence changes to these values within the subroutine are not saved.

33.4.6 Temperature
Scope: Temperature Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>real</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specify the constant value of the temperature. Using this command is equivalent to using both the FIELD = temperature and VALUE = real_value command lines.
33.4.7 Temperature Fortran Subroutine

**Scope:** Temperature Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature Fortran Subroutine {=</td>
<td>are</td>
<td>is} Name</td>
</tr>
</tbody>
</table>

**Summary**
Specify the name of a FORTRAN user subroutine that will be used to calculate the nodal temperature boundary condition.

33.4.8 Temperature Node Variable

**Scope:** Temperature Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature Node Variable {=</td>
<td>are</td>
<td>is} Name</td>
</tr>
</tbody>
</table>

**Summary**
Specify the name of the node variable to use for the temperature associated with this boundary condition.

**Description**
The indicated node variable must be a legal Aria variable. This variable is defined from the input file in the user variable command block or with the USER FIELD command. Typically, this variable would be calculated in another SIERRA region, e.g., Fuego, and transferred to Aria.

33.4.9 Temperature Scale Factor

**Scope:** Temperature Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature Scale Factor {=</td>
<td>are</td>
<td>is} Magnitude</td>
</tr>
</tbody>
</table>

**Summary**
Specifies the magnitude by which the distribution factors will be multiplied in order to determine the boundary condition. If this line command is used, node distribution factors must be defined in the input mesh for all nodesets which are associated with this boundary condition.

**Description**
If this line command is used, node distribution factors must be defined in the input mesh for all nodesets which are associated with this boundary condition. There is currently an inconsistency with respect to temperature boundary conditions and distribution factors: It is not legal to specify sideset distribution factors with the temperature boundary condition: nodesets must be used. Distribution factors also do not work with h-adaptivity, as their values are not currently interpolated to the new nodes. In any case, for temperature boundary conditions, in order for this interpolation to occur, then a sideset would have to be used, which is not supported at this time.
33.4.10  Temperature Time Function

Scope: Temperature Boundary Condition

Temperature Time Function \{=|are|is\} FunctionName

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>FunctionName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Name of the time-dependent function that specifies the temperature.

33.4.11  Use Data Block

Scope: Temperature Boundary Condition

Use Data Block Name

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Reference to predefined data to be used by the user subroutine. These values may be changed by the user subroutine.

33.4.12  Use Death

Scope: Temperature Boundary Condition

Use Death Name

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Associates this boundary condition with element faces that are created as a result of element death due to the specified element death command block. Note that this line command makes the owning boundary condition have a dynamic extent that changes as elements die.

Description  This line command specifies that the faces that are created as a result of the element death criterion in the named element death command block are added to the extent of this boundary condition. A boundary condition may specify an extent via the "add surface" line command, or the "use death" line command. In this way it is possible to have a boundary condition that is initially empty and then grows dynamically with the surface created as a result of the element death. At least one of these methods must be used.

A boundary condition may be associated with more than one death command block by including this line command more than once. However, to avoid issues of precedence, there is a rule that a given element may not die for more than one reason.

33.4.13  Use Toggle Block

Scope: Temperature Boundary Condition

Use Toggle Block ToggleName [ {@|at|for|in|on|over} ElementBlockList... ]
Parameter | Value | Default
---|---|---
**ToggleName** | **string** | **undefined**

**Summary**
Specification for toggling entities in the computational model based on Toggle Block parameters. When used at the region level, the list of element blocks to be toggled must be provided. Otherwise a listing of entities is not needed as the Toggle Block will be associated with the command line or the enclosing command block.

### 33.4.14 Value

**Scope:** Temperature Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>VariableValue</strong></td>
<td><strong>string</strong></td>
<td><strong>undefined</strong></td>
</tr>
</tbody>
</table>

**Summary**
This command is used in conjunction with the FIELD command. A constant value of the variable given in the FIELD command is specified with this boundary condition.

### 33.5 Heat Flux Boundary Condition

**Scope:** Equation System

```
Begin Heat Flux Boundary Condition Name
    Add Surface SurfaceList...
    Element Subroutine {=}is Name
    Equation {=}is EquationName
    Field Scaling {=}is Scale_factor
    Flux {=}is Value
    Flux Fortran Subroutine {=}is Name
    Flux Node Variable {=}is Name [ Multiplier {=}is Value ]
    Flux Scale Factor {=}is Value
    Flux Temperature Function {=}is Namep
    Flux Time Function {=}is Name
    Flux Vector Node Variable {=}is Name [ Multiplier {=}is Value ]
    Ignore Flux Coverage
    Integer Data Values...
    Integrated Flux Output VariableName
    Integrated Power Output VariableName
    Real Data Values...
    Scaling With Global Variable {=}is name
    Use Data Block Name
    Use Toggle Block ToggleName [ {at|for|in|on|over} ElementBlockList... ]
```
Summary This command block specifies that a known heat flux is to be applied normal to the given surface.

Description The flux can vary in arbitrary way, e.g. it may be a specified constant, a function of temperature, time etc. You may only specify one kind of flux e.g it is not legal to specify a constant value and a time dependent function. If you want to add two such fluxes on a single surface, then either write a user subroutine or use two different command blocks. If a function name is given then the function must be defined in a function definition command block.

33.5.1 Add Surface
Scope: Heat Flux Boundary Condition

Add Surface $SurfaceList...$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$SurfaceList$</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Adds surfaces, by name, to a boundary condition’s extent.

Description This line command is used to add surfaces to the extent of a boundary condition. In Exodus II, surfaces are specified as side sets, that have a global integer identifier. For example, side set 12 would be added by this line command using the surface name surface_12. Note that in SIERRA, each element of an array of strings must be separated by whitespace.

33.5.2 Element Subroutine
Scope: Heat Flux Boundary Condition

Element Subroutine $=|are|is$ $Name$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$Name$</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specifies that the named user-defined subroutine be used. At most one subroutine name may be specified for a given quantity. This subroutine must conform to the "element signature" argument list type.

33.5.3 Equation
Scope: Heat Flux Boundary Condition

Equation $=|are|is$ $EquationName$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$EquationName$</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>
This command can be used to apply the Heat Flux conditions for an equation other than the ENERGY equation.

### 33.5.4 Field Scaling

**Scope:** Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Scale_factor</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: This command is only relevant when the FLUX NODE VARIABLE or FLUX VECTOR NODE VARIABLE command line option is present. The command enables scaling of the Field by a single value for the surfaces within the Heat Flux command block. For vector Fields all components will be scaled by the provided value.

### 33.5.5 Flux

**Scope:** Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: This line command specifies a constant value of heat flux.

### 33.5.6 Flux Fortran Subroutine

**Scope:** Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify the name of a FORTRAN user subroutine that will be used to calculate the flux boundary condition.

### 33.5.7 Flux Node Variable

**Scope:** Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Specify the name of the node variable to use for the flux that is associated with this boundary condition. Here the node variable is assumed to be the product of the surface unit normal vector with the nodal heat flux. Optionally, the value can be scaled by product of Multiplier value and FIELD SCALING.

The indicated node variable must be a legal Aria variable. In most cases this variable is defined by the user in the user variable definition command block. The node variable is interpolated to the quadrature points during the evaluation of flux term contributions. If need be, both the sign and magnitude of the node variable can be modified by using the FIELD SCALING command line. Typically, the node variable would be calculated in another SIERRA region, e.g., Fuego, and transferred to Aria.

### 33.5.8 Flux Scale Factor

**Scope:** Heat Flux Boundary Condition

| Flux Scale Factor {=|are|is} Value |
|-----------------------------------|
| **Parameter** | **Value** | **Default** |
| Value | real | undefined |

**Summary**

Specifies the magnitude by which the distribution factors will be multiplied in order to determine the boundary condition. If this line command is used, sideset distribution factors must be defined in the input mesh for all surfaces which are associated with this boundary condition.

**Description**

The flux on each face is computed by interpolating the distribution factors to the Gauss points and then multiplying the result by the given magnitude. Distribution factors do not currently work with h–adaptivity, since their values are not currently interpolated to the new nodes.

### 33.5.9 Flux Temperature Function

**Scope:** Heat Flux Boundary Condition

| Flux Temperature Function {=|are|is} Namep |
|----------------------------------------|
| **Parameter** | **Value** | **Default** |
| Namep | string | undefined |

**Summary**

Specifies the name of a temperature dependent heat flux function that is used to calculate the normal flux.

### 33.5.10 Flux Time Function

**Scope:** Heat Flux Boundary Condition

| Flux Time Function {=|are|is} Name |
|-----------------------------------|
| **Parameter** | **Value** | **Default** |
| Name | string | undefined |
Summary Specifies the name of a time dependent heat flux function that is used to calculate the normal flux.

### 33.5.11 Flux Vector Node Variable

**Scope:** Heat Flux Boundary Condition

#### Flux Vector Node Variable `{=|are|is} Name [ Multiplier `{=|are|is} Value ]`

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary** Specify the name of the vector node variable to use for the flux that is associated with this boundary condition. Here the flux contribution for a surface will be evaluated by forming a dot product of the unit normal vector with the vector node variable. Optionally, the vector values can be scaled by product of Multiplier value and FIELD SCALING.

**Description** The indicated node variable must be a legal Aria vector variable. In most cases this variable is defined by the user in the user variable definition command block. The node variable is interpolated to the quadrature points during the evaluation of flux term contributions. If need be, both the sign and magnitude of the node variable can be modified by using the FIELD SCALING command line. Typically, the vector node variable would be calculated in another SIERRA region, e.g., Fuego, and transferred to Aria.

### 33.5.12 Ignore Flux Coverage

**Scope:** Heat Flux Boundary Condition

**Summary** This command causes the code to ignore the flux coverage Field when contact is present. Thus the flux will be applied even if the BC is being set on a contact surface.

### 33.5.13 Integer Data

**Scope:** Heat Flux Boundary Condition

**Summary** List of integer data values to be used by the FORTRAN user subroutine. Copies of these values are provided to the subroutine hence changes to these values within the subroutine are not saved.

### 33.5.14 Integrated Flux Output

**Scope:** Heat Flux Boundary Condition

**Summary**

Integrated Flux Output *VariableName*
Summary: Calculate the average flux associated with this flux boundary condition.

Description: This line command specifies that, as a postprocess, the normal flux associated with this boundary condition be integrated over the surface to obtain the total power. This power is then divided by the total area of the surface to obtain the average flux on the surface, and stored in a global variable named "variableName". This global variable may then be output to history files, or accessed in user subroutines, etc.

### 33.5.15 Integrated Power Output

**Scope:** Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>VariableName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Calculate the total power associated with this flux boundary condition.

Description: This line command specifies that, as a postprocess, the normal flux associated with this boundary condition be integrated over the surface to obtain the total power which is then stored into a global variable named "variableName". This global variable may then be output to history files, or accessed in user subroutines, etc.

### 33.5.16 Real Data

**Scope:** Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Values</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: List of real data values to be used by the FORTRAN user subroutine. Copies of these values are provided to the subroutine hence changes to these values within the subroutine are not saved.

### 33.5.17 Scaling With Global Variable

**Scope:** Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Summary
This command is only relevant when the FLUX NODE VARIABLE or FLUX VECTOR NODE VARIABLE command line option is present. The command enables scaling of the Field by a global variable for the surfaces within the Heat Flux command block. For vector Fields all components will be scaled by the named scalar global variable. The scalar global variable itself must be defined at the Region scope as type Real with length 1.

33.5.18 Use Data Block
Scope: Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Reference to predefined data to be used by the user subroutine. These values may be changed by the user subroutine.

33.5.19 Use Toggle Block
Scope: Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>ToggleName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Specification for toggling entities in the computational model based on Toggle Block parameters. When used at the region level, the list of element blocks to be toggled must be provided. Otherwise a listing of entities is not needed as the Toggle Block will be associated with the command line or the enclosing command block.

33.6 Laser Heat Flux Boundary Condition
Scope: Equation System

Begin Laser Heat Flux Boundary Condition Name

Absorption Coefficient {=} string value
Add Surface SurfaceList...
Angular Velocity {=} string value
Beam Diameter {=} string laser_diameter
Beam Radius {=} string value
Circular Path Center {=} string Point...
Circular Path Start Vector {=} string Value...
Effective Beam Radius {=} string value
Element Subroutine {=} string Name
Flux {=} string Value
Flux Fortran Subroutine Name
Flux Type LaserDistributionType
Integer Data Values...
Integrated Flux Output VariableName
Integrated Power Output VariableName
Max Degrees value
Path Function Function
Path Radius value
Real Data Values...
Rotation Axis Vector Value...
Source Direction Vector Value...
Source Start Location Point...
Source Velocity Vector Value...
Start Time Value
Stop Time Value
Use Data Block Name
Use Toggle Block ToggleName [ @ ElementBlockList... ]
End

Summary  This command block specifies that a laser heat flux is to be applied on the given surface.

Description  The laser heat flux depends upon the laser beam characteristics and will vary with time and space in arbitrary manner as specified by the path description and its position relative to the surface normal.

### 33.6.1  Absorption Coefficient

**Scope:** Laser Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Absorption Coefficient</td>
<td>value</td>
<td>real</td>
</tr>
<tr>
<td></td>
<td></td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Specifies the surface absorption coefficient.

### 33.6.2  Add Surface

**Scope:** Laser Heat Flux Boundary Condition

Add Surface SurfaceList...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>SurfaceList</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Summary | Adds surfaces, by name, to a boundary condition’s extent.
---|---
Description | This line command is used to add surfaces to the extent of a boundary condition. In Exodus II, surfaces are specified as side sets, that have a global integer identifier. For example, side set 12 would be added by this line command using the surface name surface_12. Note that in SIERRA, each element of an array of strings must be separated by whitespace.

### 33.6.3 Angular Velocity

**Scope:** Laser Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Angular Velocity</td>
<td>value</td>
<td>real</td>
</tr>
</tbody>
</table>

Summary | Specifies angular velocity of laser movement in revolutions per unit of time.

### 33.6.4 Beam Diameter

**Scope:** Laser Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beam Diameter</td>
<td>laser_diameter</td>
<td>real</td>
</tr>
</tbody>
</table>

Summary | Specifies laser spot size.

### 33.6.5 Beam Radius

**Scope:** Laser Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beam Radius</td>
<td>value</td>
<td>real</td>
</tr>
</tbody>
</table>

Summary | Specifies radius of laser beam.

### 33.6.6 Circular Path Center

**Scope:** Laser Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Circular Path Center</td>
<td>Point...</td>
<td>real...</td>
</tr>
</tbody>
</table>

Summary | Specifies the center of circular path for laser source travel.

853
### 33.6.7 Circular Path Start Vector

**Scope:** Laser Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Circular Path Start Vector</td>
<td>=</td>
<td>are</td>
</tr>
<tr>
<td><strong>Value</strong></td>
<td><strong>real</strong>...</td>
<td><strong>undefined</strong></td>
</tr>
</tbody>
</table>

**Summary** Specifies the starting location on circular path, a radius vector relative to CIRCULAR PATH CENTER.

### 33.6.8 Effective Beam Radius

**Scope:** Laser Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Effective Beam Radius</td>
<td>=</td>
<td>are</td>
</tr>
<tr>
<td><strong>value</strong></td>
<td><strong>real</strong></td>
<td><strong>undefined</strong></td>
</tr>
</tbody>
</table>

**Summary** Specifies effective radius of laser beam.

### 33.6.9 Element Subroutine

**Scope:** Laser Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Element Subroutine</td>
<td>=</td>
<td>are</td>
</tr>
<tr>
<td><strong>Name</strong></td>
<td><strong>string</strong></td>
<td><strong>undefined</strong></td>
</tr>
</tbody>
</table>

**Summary** Specifies that the named user-defined subroutine be used. At most one subroutine name may be specified for a given quantity. This subroutine must conform to the "element signature" argument list type.

### 33.6.10 Flux

**Scope:** Laser Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flux</td>
<td>=</td>
<td>are</td>
</tr>
<tr>
<td><strong>Value</strong></td>
<td><strong>real</strong></td>
<td><strong>undefined</strong></td>
</tr>
</tbody>
</table>

**Summary** This line command specifies a constant value of heat flux.

### 33.6.11 Flux Fortran Subroutine

**Scope:** Laser Heat Flux Boundary Condition
Flux Fortran Subroutine 

**Parameter** 

<table>
<thead>
<tr>
<th>Name</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary** 
Specify the name of a FORTRAN user subroutine that will be used to calculate the flux boundary condition.

### 33.6.12 Flux Type

**Scope:** Laser Heat Flux Boundary Condition

**Summary** 
Selects the laser flux distribution model applied to a portion of the underlying surface as defined by a geometric path function. Flux types produce either a Gaussian flux with maximum intensity at the center of the laser spot or constant flux.

### 33.6.13 Integer Data

**Scope:** Laser Heat Flux Boundary Condition

**Summary** 
List of integer data values to be used by the FORTRAN user subroutine. Copies of these values are provided to the subroutine hence changes to these values within the subroutine are not saved.

### 33.6.14 Integrated Flux Output

**Scope:** Laser Heat Flux Boundary Condition

**Summary** 
Calculate the average flux associated with this flux boundary condition.

**Description** 
This line command specifies that, as a postprocess, the normal flux associated with this boundary condition be integrated over the surface to obtain the total power. This power is then divided by the total area of the surface to obtain the average flux on the surface, and stored in a global variable named "variableName". This global variable may then be output to history files, or accessed in user subroutines, etc.

### 33.6.15 Integrated Power Output

**Scope:** Laser Heat Flux Boundary Condition
Integrated Power Output \textit{VariableName}

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{VariableName}</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Calculate the total power associated with this flux boundary condition.

Description  This line command specifies that, as a postprocess, the normal flux associated with this boundary condition be integrated over the surface to obtain the total power which is then stored into a global variable named \textquote{variableName}. This global variable may then be output to history files, or accessed in user subroutines, etc.

33.6.16 Max Degrees

Scope: Laser Heat Flux Boundary Condition

\texttt{Max Degrees \{=|are|is\} value}

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Specifies maximum degrees of laser rotational motion.

33.6.17 Path Function

Scope: Laser Heat Flux Boundary Condition

\texttt{Path Function \{=|are|is\} Function}

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Specifies a time and position function for which a laser source is applied in the named volume.

33.6.18 Path Radius

Scope: Laser Heat Flux Boundary Condition

\texttt{Path Radius \{=|are|is\} value}

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Specifies radius for circular laser path.

33.6.19 Real Data

Scope: Laser Heat Flux Boundary Condition

\texttt{Real Data Values...}
33.6.20 Rotation Axis Vector

Scope: Laser Heat Flux Boundary Condition

Rotation Axis Vector {=|are|is} Value...

Summary: Specifies the axis of rotation for circular source travel, relative to CIRCULAR PATH CENTER.

33.6.21 Source Direction Vector

Scope: Laser Heat Flux Boundary Condition

Source Direction Vector {=|are|is} Value...

Summary: Specifies the vector orientation of the laser source.

33.6.22 Source Start Location

Scope: Laser Heat Flux Boundary Condition

Source Start Location {=|are|is} Point...

Summary: Specifies the physical coordinate location of the laser source.

33.6.23 Source Velocity Vector

Scope: Laser Heat Flux Boundary Condition

Source Velocity Vector {=|are|is} Value...

Summary: Specifies the velocity vector of laser source motion.
### 33.6.24 Start Time

**Scope:** Laser Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Start Time</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specifies the time at which the laser beam is activated.

### 33.6.25 Stop Time

**Scope:** Laser Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stop Time</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specifies the time at which the laser beam is deactivated.

### 33.6.26 Use Data Block

**Scope:** Laser Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Use Data Block</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Reference to predefined data to be used by the user subroutine. These values may be changed by the user subroutine.

### 33.6.27 Use Toggle Block

**Scope:** Laser Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Use Toggle Block</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specification for toggling entities in the computational model based on Toggle Block parameters. When used at the region level, the list of element blocks to be toggled must be provided. Otherwise a listing of entities is not needed as the Toggle Block will be associated with the command line or the enclosing command block.
33.7 Convection Heat Transfer

A broad class of thermal applications involve heat transfer between a solid volume and a surrounding fluid. In numerical simulations these interactions are modeled using various surface boundary conditions depending upon the detail to which the fluid flow is resolved. If both heat transfer in the solid model and fluid flow are explicitly modeled then it is often appropriate to solve the conjugate heat transfer problem. However, in many cases the expense of this calculation may not be justified and one reverts to alternative methods of modeling these effects.

In many situations it may be appropriate to simply characterize a bulk behavior of the fluid and employ a convective flux boundary condition of the form

\[ q = h(T)(T_s - T_b) \]

where \( q \) is the heat flux per unit surface area, \( h(T) \) is the surface heat transfer coefficient or convection coefficient, \( T_s \) is the surface temperature and \( T_b \) is the bulk fluid temperature. Many of the engineering models based upon experimental data for given configurations include variations of the values that enter into the expression above. In particular the values of \( h(T) \) and \( T_b \) are often linked to representative fluid flow conditions.

For quick estimates of heat transfer, textbooks provide a tabulated range of surface heat transfer coefficients for general conditions. Furthermore, heat transfer handbooks provide a collection of heat transfer correlations from which convective coefficients can be computed. These correlations include empirical constants and are functions dimensionless numbers which characterize the bulk fluid flow and the bulk fluid temperature.

The Sierra Thermal module includes a number of different ways for specifying \( h(T) \) and \( T_b \). Additionally module provides a catalog of heat transfer correlations 34.1.

33.8 Convective Flux Boundary Condition

Scope:  Equation System

```
Begin Convective Flux Boundary Condition Name
  Add Surface SurfaceList...
  Average Temperature Variable {=|are|is} NName
  Convective Coefficient {=|are|is} Value
  Convective Coefficient Fortran Subroutine {=|are|is} Name
  Convective Coefficient Node Variable {=|are|is} Name
  Convective Coefficient Scale Factor {=|are|is} Magnitude
  Convective Coefficient Subroutine {=|are|is} Name
  Convective Coefficient Temperature Difference Function {=|are|is} NName
  Convective Coefficient Temperature Function {=|are|is} NName
  Convective Coefficient Time Function {=|are|is} Name
  Equation {=|are|is} EquationName
  Ignore Flux Coverage
  Integer Data Values...
  Integrated Flux Output VariableName
```
Integrated Power Output \textit{VariableName}

Real Data \textit{Values...}

Ref Temp Convective Coefficient Subroutine \{=|are|is\} \textit{Name} [ \textit{UsingRefTemp} ]

Reference Temperature \{=|are|is\} \textit{Value}

Reference Temperature Fortran Subroutine \{=|are|is\} \textit{Name}

Reference Temperature Global Variable \{=|are|is\} \textit{GlobalVariableName}

Reference Temperature Node Variable \{=|are|is\} \textit{Name}

Reference Temperature Subroutine \{=|are|is\} \textit{Name}

Reference Temperature Temperature Function \{=|are|is\} \textit{Name}

Reference Temperature Time Function \{=|are|is\} \textit{FunctionName}

Scaled Convective Coefficient Subroutine \{=|are|is\} \textit{Name} \textit{FieldName} \{=|are|is\} \textit{Field}

Scaled Ref Temp Convective Coefficient Subroutine \{=|are|is\} \textit{Name} \textit{FieldName} \{=|are|is\} \textit{Field}

\textit{Uq Flux Multiplier} \{=|are|is\} \textit{Value}

Use Advective Bar \textit{Name} [ \textit{BulkNodes} ]

Use Bulk Element \textit{Name}

Use Correlation Convection Model \textit{Name}

Use Data Block \textit{Name}

Use Enclosure \textit{Name}

Use Toggle Block \textit{TagName} [ \{@|at|for|in|on|over\} \textit{ElementBlockList...} ]

User Field Mask \{=|are|is\} \textit{Name} [ \textit{Threshold} \{=|are|is\} \textit{Value} ]

User Field Scaling \{=|are|is\} \textit{Name}

\textbf{End}

\textbf{Summary} This command block specifies heat transfer on a boundary surface that can be modeled using Newton’s law of cooling.

\textbf{Description} Newton’s law of cooling specifies that the heat flux normal to a surface is proportional to the difference between the unknown temperature of the surface and some reference temperature of the fluid in which the surface is immersed: \( q_n = h(T - T_r) \). The convection coefficient, \( h \), and the reference temperature, \( T_r \), may be specified in several ways as explained in detail below. In particular, note that the reference temperature is usually a known quantity, because the fluid with which it is associated is modeled as an infinite reservoir. However, if the size of this reservoir is finite, then its temperature can be affected by the energy transfer across the surface in question. This situation can be modeled by the bulk fluid element, wherein the energy of the reservoir is determined by a finite volume conservation equation.

You must specify exactly one convection coefficient, and either exactly one reference temperature or exactly one bulk fluid element name.
33.8.1 Add Surface
Scope: Convective Flux Boundary Condition

Add Surface *SurfaceList...*

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>SurfaceList</em></td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Adds surfaces, by name, to a boundary condition’s extent.

Description: This line command is used to add surfaces to the extent of a boundary condition. In Exodus II, surfaces are specified as side sets, that have a global integer identifier. For example, side set 12 would be added by this line command using the surface name surface_12. Note that in SIERRA, each element of an array of strings must be separated by whitespace.

33.8.2 Average Temperature Variable
Scope: Convective Flux Boundary Condition

Average Temperature Variable *=|are|is* *NName*

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>NName</em></td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify the name of a global variable to store the average temperature of the specified surfaces for this convective flux command block. This is used if the convective coefficient temperature difference function is set else it is ignored.

33.8.3 Convective Coefficient
Scope: Convective Flux Boundary Condition

Convective Coefficient *=|are|is* *Value*

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Value</em></td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify a constant convective coefficient for this boundary condition.

33.8.4 Convective Coefficient Fortran Subroutine
Scope: Convective Flux Boundary Condition

Convective Coefficient Fortran Subroutine *=|are|is* *Name*

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Name</em></td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify the name of a FORTRAN user subroutine that will be used to calculate the convective coefficient for this boundary condition.
33.8.5 Convective Coefficient Node Variable

Scope: Convective Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify the name of the node variable to use for the convective coefficient that is associated with this boundary condition.

Description: The indicated node variable must be a legal Aria variable. Typically, this variable is defined by the user in the user definition command block. The node variable is interpolated to the integration points during the integration of the flux term. Typically, this variable would be calculated in another SIERRA region, e.g., Fuego, and transferred to Aria.

33.8.6 Convective Coefficient Scale Factor

Scope: Convective Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Magnitude</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify that the convective coefficient is to be computed using sideset distribution factors that must be defined in the mesh database.

Description: The coefficient on each face is computed by interpolating the distribution factors to the Gauss points and then multiplying the result by the given magnitude. Distribution factors do not currently work with h–adaptivity, since their values are not currently interpolated to the new nodes.

33.8.7 Convective Coefficient Subroutine

Scope: Convective Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify the name of a user subroutine that is to be used to calculate the convective coefficient for this boundary condition.

If the user subroutine employs a user specified reference temperature model to compute the heat transfer coefficient then optional arguments USING_REF_TEMP must also appear at the end of the command line.
### 33.8.8 Convective Coefficient Temperature Difference Function

**Scope:** Convective Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{NName}</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specify the name of a temperature difference-dependent function that is to be used to calculate the convective coefficient for this boundary condition. If this line command is specified, then a line command for the average temperature must also be specified.

### 33.8.9 Convective Coefficient Temperature Function

**Scope:** Convective Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{NName}</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specify the name of a temperature-dependent function that is to be used to calculate the convective coefficient for this boundary condition.

### 33.8.10 Convective Coefficient Time Function

**Scope:** Convective Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{Name}</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specify the name of a time-dependent function that is used to calculate the convective coefficient for this boundary condition.

### 33.8.11 Equation

**Scope:** Convective Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{EquationName}</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
This command can be used to apply the Convective Flux conditions for an equation other than the ENERGY equation.
33.8.12 Ignore Flux Coverage
Scope: Convective Flux Boundary Condition

Summary This command causes the code to ignore the flux coverage Field when contact is present. Thus the flux will be applied even if the BC is being set on a contact surface.

33.8.13 Integer Data
Scope: Convective Flux Boundary Condition

<table>
<thead>
<tr>
<th>Integer Data Values...</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter</td>
</tr>
<tr>
<td>Values</td>
</tr>
</tbody>
</table>

Summary List of integer data values to be used by the FORTRAN user subroutine. Copies of these values are provided to the subroutine hence changes to these values within the subroutine are not saved.

33.8.14 Integrated Flux Output
Scope: Convective Flux Boundary Condition

<table>
<thead>
<tr>
<th>Integrated Flux Output VariableName</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter</td>
</tr>
<tr>
<td>VariableName</td>
</tr>
</tbody>
</table>

Summary Calculate the average flux associated with this flux boundary condition.

Description This line command specifies that, as a postprocess, the normal flux associated with this boundary condition be integrated over the surface to obtain the total power. This power is then divided by the total area of the surface to obtain the average flux on the surface, and stored in a global variable named "VariableName". This global variable may then be output to history files, or accessed in user subroutines, etc.

33.8.15 Integrated Power Output
Scope: Convective Flux Boundary Condition

<table>
<thead>
<tr>
<th>Integrated Power Output VariableName</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter</td>
</tr>
<tr>
<td>VariableName</td>
</tr>
</tbody>
</table>

Summary Calculate the total power associated with this flux boundary condition.

Description This line command specifies that, as a postprocess, the normal flux associated with this boundary condition be integrated over the surface to obtain the total power which is then...
stored into a global variable named "variableName". This global variable may then be output to history files, or accessed in user subroutines, etc.

### 33.8.16 Real Data

**Scope:** Convective Flux Boundary Condition

**Real Data Values...**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Values</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

List of real data values to be used by the FORTRAN user subroutine. Copies of these values are provided to the subroutine hence changes to these values within the subroutine are not saved.

### 33.8.17 Ref Temp Convective Coefficient Subroutine

**Scope:** Convective Flux Boundary Condition

**Ref Temp Convective Coefficient Subroutine {=|are|is} Name [ UsingRefTemp ]**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Specify the name of a user subroutine that is to be used to calculate the convective coefficient for this boundary condition. Use of this command line will enable the reference temperature to be supplied to the subroutine interface.

### 33.8.18 Reference Temperature

**Scope:** Convective Flux Boundary Condition

**Reference Temperature {=|are|is} Value**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Specify a constant reference temperature for this boundary condition.

### 33.8.19 Reference Temperature Fortran Subroutine

**Scope:** Convective Flux Boundary Condition

**Reference Temperature Fortran Subroutine {=|are|is} Name**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Specifies the name of a FORTRAN user-defined subroutine that will be used to calculate the reference temperature associated with this boundary condition.
33.8.20 Reference Temperature Global Variable

Scope: Convective Flux Boundary Condition

Reference Temperature Global Variable \{=|are|is\} GlobalVariableName

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>GlobalVariableName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specify a global variable to be used for reference temperature.

33.8.21 Reference Temperature Node Variable

Scope: Convective Flux Boundary Condition

Reference Temperature Node Variable \{=|are|is\} Name

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specify the name of the node variable to use for the reference temperature that is associated with this boundary condition.

Description The indicated node variable must be a legal Aria variable. Typically, this variable is defined by the user in the user definition command block. The node variable is interpolated to the integration points during the integration of the flux term. Typically, this variable would be calculated in another SIERRA region, e.g., Fuego, and transferred to Aria.

33.8.22 Reference Temperature Subroutine

Scope: Convective Flux Boundary Condition

Reference Temperature Subroutine \{=|are|is\} Name

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specifies the name of a user-defined subroutine that is to be used to calculate the reference temperature associated with this boundary condition.

33.8.23 Reference Temperature Temperature Function

Scope: Convective Flux Boundary Condition

Reference Temperature Temperature Function \{=|are|is\} Name

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specifies the name of the temperature-dependent function that is to be used to calculate the reference temperature associated with this boundary condition.
33.8.24 Reference Temperature Time Function
Scope: Convective Flux Boundary Condition

Reference Temperature Time Function \(\text{are|is} \text{ FunctionName}\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>FunctionName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify the name of a time-dependent function for the reference temperature for this boundary condition.

33.8.25 Scaled Convective Coefficient Subroutine
Scope: Convective Flux Boundary Condition

Scaled Convective Coefficient Subroutine \(\text{are|is} \text{ Name FieldName =|are|is} \text{ Field}\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>FieldName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Field</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify the name of a user subroutine that is to be used to calculate the scaled convective coefficient for this boundary condition. The interpolated scaling Field is supplied directly to the user subroutine.

33.8.26 Scaled Ref Temp Convective Coefficient Subroutine
Scope: Convective Flux Boundary Condition

Scaled Ref Temp Convective Coefficient Subroutine \(\text{are|is} \text{ Name FieldName =|are|is} \text{ Field}\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>FieldName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Field</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify the name of a user subroutine that is to be used to calculate the scaled convective coefficient for this boundary condition. Use of this command will enable the reference temperature to be supplied to the user subroutine. Additionally interpolated values of the scaling Field will be delivered to the subroutine.

33.8.27 Uq Flux Multiplier
Scope: Convective Flux Boundary Condition

Uq Flux Multiplier \(\text{are|is} \text{ Value}\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>real</td>
<td>1.0</td>
</tr>
</tbody>
</table>
Summary Specify constant scaling of the convective flux.

Description Intended use of this scaling parameter is primarily for evaluation of model sensitivities.

33.8.28 Use Advective Bar
Scope: Convective Flux Boundary Condition

<table>
<thead>
<tr>
<th>Use Advective Bar Name [ BulkNodes ]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter</td>
</tr>
<tr>
<td>Name</td>
</tr>
</tbody>
</table>

Summary Use the named advective bar to model the reference temperature.

Description This line command specifies the name of an advective bar that has been defined using the advective bar command block. The temperature of the advective bar is used as the reference temperature, \( T_r \), for the convective heat transfer and this is computed based on a geometric coupling algorithm. Note that it is illegal to specify both a reference temperature and an advective bar.

33.8.29 Use Bulk Element
Scope: Convective Flux Boundary Condition

<table>
<thead>
<tr>
<th>Use Bulk Element Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter</td>
</tr>
<tr>
<td>Name</td>
</tr>
</tbody>
</table>

Summary Use the named bulk element to model the reference temperature.

Description This line command specifies the name of a bulk fluid element that has been defined using the bulk fluid command block. The temperature of the bulk fluid element is used as the reference temperature, \( T_r \), for the convective heat transfer. Note that it is illegal to specify both a reference temperature and a bulk element.

33.8.30 Use Correlation Convection Model
Scope: Convective Flux Boundary Condition

<table>
<thead>
<tr>
<th>Use Correlation Convection Model Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter</td>
</tr>
<tr>
<td>Name</td>
</tr>
</tbody>
</table>

Summary Specifies correlation model for convection coefficient
33.8.31 Use Data Block
Scope: Convective Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Reference to predefined data to be used by the user subroutine. These values may be changed by the user subroutine.

33.8.32 Use Enclosure
Scope: Convective Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Use the named enclosure to define the surface list and possibly model the reference temperature using the MBL bulk node of the enclosure.

Description: This line command specifies the name of an enclosure that has been defined using the enclosure definition command block. The list of surfaces for this convective BC is taken from the enclosure surfaces and it is illegal to specify both an enclosure and a surface list. If the enclosure has the Mean Beam Length (MBL) model activated, then this convective BC is linked to the bulk fluid element associated with the MBL model. The temperature of the bulk fluid element is used as the reference temperature, $T_r$, for the convective heat transfer. Note that it is illegal to specify both a reference temperature and an enclosure with an MBL bulk element.

33.8.33 Use Toggle Block
Scope: Convective Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>ToggleName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specification for toggling entities in the computational model based on Toggle Block parameters. When used at the region level, the list of element blocks to be toggled must be provided. Otherwise a listing of entities is not needed as the Toggle Block will be associated with the command line or the enclosing command block.

33.8.34 User Field Mask
Scope: Convective Flux Boundary Condition
33.8.35 User Field Mask

**Summary**
Specify that the convective boundary condition is to be masked by nonzero values of a user defined Field.

**Description**
The convective boundary condition is applied in a conventional manner but then masked by nonzero values of an interpolated user defined Field. In most cases the user Field is transferred to Aria but it could also be computed. Default threshold for the masked value is 0.0, i.e. mask value equals 1.0 for values greater than zero.

**33.8.35 User Field Scaling**

**Scope:** Convective Flux Boundary Condition

**Summary**
Specify that the convective boundary condition is to be scaled by a user defined Field.

**Description**
The convective boundary condition is applied in a conventional manner but then scaled by an interpolated user defined Field. In most cases the user Field is transferred to Aria and acts as a weighting of the flux condition.

### 33.9 Aerodynamic Convection Heat Transfer

High-speed flows about bodies are often characterized by large temperature gradients and large variations of the flow properties through the boundary layer. The problem of heat transfer about the body is often addressed using numerical techniques of CFD to resolve the thermal transport through the boundary layer. Because CFD resources may not be readily applied at the design stage researchers often apply simplifying assumptions to arrive at alternative models for solving the problem of heat transfer about bodies exposed to high-speed flow. One common modeling approach is to define a convection coefficient and reference temperature that depend upon flight conditions. The aero heat flux boundary condition provides a means for supplying details of the flight conditions to arrive at a representative convection coefficient and reference temperature for two heat flux models, one derived by [37] and one that varies only with altitude.

The model by Eckert suggests that a convective heat transfer approach ignoring boundary layer temperature gradients may be used if the properties are evaluated at an alternative reference temperature representative of the averaged flow conditions. Syntax for typical usage of this model is shown below.

```plaintext
Begin AERO HEAT FLUX BOUNDARY CONDITION body_heating
  START TIME = 0.0
  STOP TIME = 20.0
  freestream temperature = 233.15
  freestream pressure   = 5.066e3
End
```
mach number = 3.0
fluid viscosity temperature function = air_viscosity
fluid specific heat temperature function = air_specheat
fluid thermal conductivity temperature function = air_cond
fluid gamma = 1.4
adiabatic wall temperature from recovery factor
eckert convective coefficient
COORDINATE OFFSET AXIS = x
cone length = 0.0
add surface surface_1
integrated power output total_heating
End

Another aerodynamic heat flux model defines a convective coefficient that relies upon the variation of density with altitude. Syntax for typical usage of this model is shown below.

Begin AERO HEAT FLUX BOUNDARY CONDITION body_heating
START TIME = 0.0
STOP TIME = 20.0
Density Ratio Convective Coefficient
Freestream Density Altitude Function = density_func
Altitude time function = altitude_time_func
Reference Density = 1.15
Reference HTC = 25.0
Density Ratio exponent = 0.6
reference temperature time function = ref_temp_func
add surface surface_1
integrated power output total_heating
End

A more complete description of syntax for the aero heating models is described in the section which follows.

33.10 Aero Heat Flux Boundary Condition
Scope: Equation System

Begin Aero Heat Flux Boundary Condition Name
Add Surface SurfaceList...
Adiabatic Wall Temperature {=|are|is} Value
Adiabatic Wall Temperature Altitude Function {=|are|is} Function
Adiabatic Wall Temperature From Recovery Factor
Adiabatic Wall Temperature Time Function {=|are|is} Function
Altitude {=|are|is} Value
Altitude Time Function {=|are|is} Function
Cone Length {=|are|is} Value
Coordinate Offset {=|are|is} Value
Coordinate Offset Axis {=|are|is} axis
Density Ratio Convective Coefficient
Density Ratio Exponent \(=\) Value
Eckert Convective Coefficient
Fluid Gamma \(=\) Value
Fluid Properties Temperature Function \(=\) Function
Fluid Specific Heat Temperature Function \(=\) Function
Fluid Thermal Conductivity Temperature Function \(=\) Function
Fluid Viscosity Temperature Function \(=\) Function
Freestream Density Altitude Function \(=\) Function
Freestream Pressure \(=\) Value
Freestream Pressure Altitude Function \(=\) Function
Freestream Pressure Time Function \(=\) Function
Freestream Temperature \(=\) Value
Freestream Temperature Altitude Function \(=\) Function
Freestream Temperature Time Function \(=\) Function
Integrated Flux Output \(\text{VariableName}\)
Integrated Power Output \(\text{VariableName}\)
Mach Number \(=\) Value
Mach Number Time Function \(=\) Function
Reference Density \(=\) Value
Reference Htc \(=\) Value
Reference Temperature \(=\) Value
Reference Temperature Time Function \(=\) FunctionName
Start Time \(=\) Value
Stop Time \(=\) Value
Uq Flux Multiplier \(=\) Value
Use Toggle Block \(\text{ToggleName}\) [ \{@|at|for|in|on|over\} \text{ElementBlockList}... ]

Summary  This command block defines a distributed convective heat flux about a body based upon aerodynamic flight conditions applied on the given surface.

Description  The aerodynamic heat flux depends upon characterization of flight conditions prescribed using data tables and spatial location on the body of interest.

### 33.10.1 Add Surface

**Scope:**  Aero Heat Flux Boundary Condition

Add Surface \(\text{SurfaceList}...\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\text{SurfaceList})</td>
<td>\text{string}...</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Summary Adds surfaces, by name, to a boundary condition’s extent.

Description This line command is used to add surfaces to the extent of a boundary condition. In Exodus II, surfaces are specified as side sets, that have a global integer identifier. For example, side set 12 would be added by this line command using the surface name surface_12. Note that in SIERRA, each element of an array of strings must be separated by whitespace.

33.10.2 Adiabatic Wall Temperature
Scope: Aero Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter Value</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adiabatic Wall Temperature (=</td>
<td>\text{are}</td>
<td>\text{is}) Value</td>
</tr>
</tbody>
</table>

Summary Specifies a constant adiabatic wall temperature. Usage: ECKERT CONVECTIVE COEFFICIENT.
FREESTREAM TEMPERATURE and fluid properties consistent with this ADIABATIC WALL TEMPERATURE must also be provided.

33.10.3 Adiabatic Wall Temperature Altitude Function
Scope: Aero Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter Function</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adiabatic Wall Temperature Altitude Function (=</td>
<td>\text{are}</td>
<td>\text{is}) Function</td>
</tr>
</tbody>
</table>

Summary Specifies the adiabatic wall temperature versus altitude function for climate condition. Usage: ECKERT CONVECTIVE COEFFICIENT.
FREESTREAM TEMPERATURE and fluid properties consistent with this ADIABATIC WALL TEMPERATURE must also be provided.

33.10.4 Adiabatic Wall Temperature From Recovery Factor
Scope: Aero Heat Flux Boundary Condition

Summary Specifies that the adiabatic wall temperature be computed using the recovery factor, the stagnation temperature and the freestream temperature. Usage: ECKERT CONVECTIVE COEFFICIENT.

Description The adiabatic wall temperature \(T_{aw}\) is computed as

\[
T_{aw} = rT_o + (1 - r)T_\infty
\]

where \(r\) is the recovery factor, \(T_o\) is the stagnation temperature and \(T_\infty\) is the freestream temperature. The user must provide specification of MACH NUMBER, FREESTREAM
TEMPERATURE, FREESTREAM PRESSURE and FLUID_GAMMA models. Stagnation temperature and flow velocity \( v \) will be internally computed from the MACH NUMBER. The recovery factor is given in terms of the freestream Prandtl number \( \frac{C_p \mu}{K} \) as

\[
r = \begin{cases} 
  \frac{P}{r^{1/2}} & \text{laminar flow} \\
  \frac{P}{r^{1/3}} & \text{turbulent flow} \quad Re > 5 \times 10^5 
\end{cases}
\]

Hence one must provide functions for specific heat, dynamic viscosity and thermal conductivity as a function of temperature. The flow regime varies spatially in the COORDINATE OFFSET direction as determined using the local Reynolds number

\[
Re_x = \frac{\rho v x}{\mu}.
\]

Here the pressure \( P \) will vary with altitude hence density is evaluated using the ideal gas law

\[
\rho = \frac{P}{RT_{\infty}}.
\]

33.10.5 Adiabatic Wall Temperature Time Function

**Scope:** Aero Heat Flux Boundary Condition

| Adiabatic Wall Temperature Time Function \{=|are|is\} Function |
|-----------------------------------------------------------|
| **Parameter** | **Value** | **Default** |
| Function | string | undefined |

**Summary**

Specifies the name of adiabatic wall temperature versus time function. Usage: ECKERT CONVECTIVE COEFFICIENT.

FREESTREAM TEMPERATURE and fluid properties consistent with this ADIABATIC WALL TEMPERATURE must also be provided.

33.10.6 Altitude

**Scope:** Aero Heat Flux Boundary Condition

| Altitude \{=|are|is\} Value |
|-----------------------------|
| **Parameter** | **Value** | **Default** |
| Value | real | 0.0 |

**Summary**

Specifies a constant altitude.

Used in conjunction with FREESTREAM TEMPERATURE ALTITUDE FUNCTION, FREESTREAM TEMPERATURE FUNCTION and ECKERT CONVECTIVE COEFFICIENT. Alternative usage with FREESTREAM DENSITY ALTITUDE FUNCTION and DENSITY RATIO CONVECTIVE COEFFICIENT.

33.10.7 Altitude Time Function

**Scope:** Aero Heat Flux Boundary Condition

| Altitude Time Function \{=|are|is\} Function |
### Summary

Specifies the altitude versus time function for a given flight. The tabulated time is defined relative to the START TIME.

Used in conjunction with FREESTREAM TEMPERATURE ALTITUDE FUNCTION, FREESTREAM TEMPERATURE FUNCTION and ECKERT CONVECTIVE COEFFICIENT. Alternative usage with FREESTREAM DENSITY ALTITUDE FUNCTION and DENSITY RATIO CONVECTIVE COEFFICIENT.

#### 33.10.8 Cone Length

**Scope:** Aero Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cone Length</td>
<td>Value</td>
<td></td>
</tr>
<tr>
<td>Parameter</td>
<td>Value</td>
<td>Default</td>
</tr>
<tr>
<td>Value</td>
<td>real</td>
<td>0.0</td>
</tr>
</tbody>
</table>

Summary

Specifies the length at which the body profile transitions to a constant diameter. Usage: ECKERT CONVECTIVE COEFFICIENT.

#### 33.10.9 Coordinate Offset

**Scope:** Aero Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coordinate Offset</td>
<td>Value</td>
<td></td>
</tr>
<tr>
<td>Parameter</td>
<td>Value</td>
<td>Default</td>
</tr>
<tr>
<td>Value</td>
<td>real</td>
<td>0.0</td>
</tr>
</tbody>
</table>

Summary

Specifies the coordinate offset from which the spatial position will be defined in the ECKERT CONVECTIVE COEFFICIENT calculations \( x = X - \text{offset} \) where \( X \) is the Cartesian coordinate prescribed using COORDINATE OFFSET AXIS.

#### 33.10.10 Coordinate Offset Axis

**Scope:** Aero Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coordinate Offset Axis</td>
<td>axis</td>
<td></td>
</tr>
<tr>
<td>Parameter</td>
<td>Value</td>
<td>Default</td>
</tr>
<tr>
<td>axis</td>
<td>string</td>
<td>none</td>
</tr>
</tbody>
</table>

Summary

Specifies the Cartesian coordinate axis (X, Y or Z) from which spatial position will be defined when using the ECKERT CONVECTIVE COEFFICIENT. The surface heat transfer coefficient will be computed spatially relative to zero on this axis.

For models in which the geometry is offset from a zero reference one must define the COORDINATE OFFSET.
33.10.11 Density Ratio Convective Coefficient  
**Scope:** Aero Heat Flux Boundary Condition  

**Summary**  
Specifies a density ratio heat transfer coefficient model. Requires that FREESTREAM DENSITY ALTITUDE FUNCTION, REFERENCE DENSITY, REFERENCE HTC, DENSITY RATIO EXPONENT, START TIME and STOP TIME specifications be supplied.  

**Description**  
The heat transfer coefficient that varies as  
\[ h = h_o \left( \frac{\rho(A)}{\rho_o} \right)^n \]  
where \( h_o \) is the reference heat transfer coefficient, \( \rho(A) \) is the altitude dependent density, \( \rho_o \) is the reference density and \( n \) is the density ratio exponent. Here the model parameters \( h_o, \rho_o \) and \( n \) are calibrated based upon measured temperature data.  

33.10.12 Density Ratio Exponent  
**Scope:** Aero Heat Flux Boundary Condition  

**Table:**  

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Density Ratio Exponent</td>
<td>{=</td>
<td>are</td>
</tr>
<tr>
<td>Value</td>
<td>real</td>
<td>0.0</td>
</tr>
</tbody>
</table>

**Summary**  
Specifies the density ratio exponent value for the density ratio heat transfer coefficient model. Usage: DENSITY RATIO CONVECTIVE COEFFICIENT.  

33.10.13 Eckert Convective Coefficient  
**Scope:** Aero Heat Flux Boundary Condition  

**Summary**  
Specifies use of the Eckert heat transfer coefficient model for high-speed flow. Requires that freestream FLUID PROPERTY TEMPERATURE FUNCTIONs, START TIME, STOP TIME, FREESTREAM PRESSURE and ADIABATIC WALL TEMPERATURE specifications be supplied.  

**Description**  
A constant property heat transfer coefficient approach in which properties are evaluated at an alternative temperature  
\[ T^* = T_\infty + 0.5(T - T_\infty) + 0.22(T_{aw} - T_\infty) \]  
where \( T_\infty \) is the freestream temperature, \( T \) is the surface temperature, \( T_{aw} \) is the adiabatic wall temperature. \( T_{aw} \) will vary spatially depending upon the local Reynolds number  
\[ Re_x = \frac{\rho^* v x}{\mu} \]  
where \( \mu \) is evaluated at \( T^* \). The FREESTREAM PRESSURE varies with altitude hence the density \( \rho^* \) is evaluated using the ideal gas law  
\[ \rho^* = \frac{P}{RT^*} \].
The Stanton number represents the ratio of heat transferred into the fluid flow to the thermal capacity of the fluid. By developing the relationship between fluid friction and heat transfer frictional resistance, the heat transfer can be expressed in terms of the Stanton number to arrive at an approximate heat transfer coefficient for various portions of the flow regime

\[ St^*Pr^{2/3} = \begin{cases} 
0.332 \left( \frac{Re_x^*}{f_M} \right)^{-1/2} & Re_x^* < 5 \times 10^5 \text{ laminar} \\
0.0296 \left( \frac{Re_x^*}{f_M} \right)^{-1/5} & 5 \times 10^5 < Re_x^* < 10^7 \text{ turbulent} \\
0.185 [\log_{10}(Re_x^*/f_M)]^{-2.584} & 10^7 < Re_x^* < 10^9 \text{ turbulent} \\
0.0063 & Re_x^* > 10^9 \text{ turbulent} 
\end{cases} \]

The parameter \( f_M \) is the Mangler transformation for cone geometry

\[ f_M = \begin{cases} 
3 & \text{cone geometry laminar} \\
2 & \text{cone geometry turbulent} \\
1 & \text{flat plate geometry laminar} 
\end{cases} \]

While Eckert’s approach considers an average heat transfer coefficient over portions of a surface, here we consider a local heat transfer coefficient, \( h(x) \) based upon the above relations that can be applied directly to the surface discretization

\[ h(x) = \rho^*C_p^*U_\infty St_x^* \]

### 33.10.14 Fluid Gamma

**Scope:** Aero Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Fluid Gamma {=}</th>
<th>are</th>
<th>is</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Parameter</strong></td>
<td>Value</td>
<td>Default</td>
<td></td>
</tr>
<tr>
<td>Value</td>
<td>real</td>
<td>1.4</td>
<td></td>
</tr>
</tbody>
</table>

**Summary**

Specifies a constant specific heat ratio of the fluid medium. Usage: ECKERT CONVEXTIVE COEFFICIENT.

Only constant values of \( \gamma \) are allowed.

### 33.10.15 Fluid Properties Temperature Function

**Scope:** Aero Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Fluid Properties Temperature Function {=}</th>
<th>are</th>
<th>is</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Parameter</strong></td>
<td>Value</td>
<td>Default</td>
<td></td>
</tr>
<tr>
<td>Function</td>
<td>string</td>
<td>undefined</td>
<td></td>
</tr>
</tbody>
</table>

**Summary**

Specifies a multi-column function containing fluid properties density, specific heat, conductivity and dynamic viscosity versus temperature with respective property columns named CP, K and MU. Usage: ECKERT CONVEXTIVE COEFFICIENT.

### 33.10.16 Fluid Specific Heat Temperature Function

**Scope:** Aero Heat Flux Boundary Condition
### Fluid Specific Heat Temperature Function

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td></td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specifies the name of a user tabular function for specific heat as a function of temperature. Usage: ECKERT CONVECTIVE COEFFICIENT.

### 33.10.17 Fluid Thermal Conductivity Temperature Function

**Scope:** Aero Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td></td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specifies the name of a user tabular function for thermal conductivity as a function of temperature. Usage: ECKERT CONVECTIVE COEFFICIENT.

### 33.10.18 Fluid Viscosity Temperature Function

**Scope:** Aero Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td></td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specifies the name of a user tabular function for dynamic viscosity as a function of temperature. Usage: ECKERT CONVECTIVE COEFFICIENT.

### 33.10.19 Freestream Density Altitude Function

**Scope:** Aero Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td></td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specifies the freestream density versus altitude function for climate condition. Usage: DENSITY RATIO CONVECTIVE COEFFICIENT.

Requires that one supply an ALTITUDE model.

### 33.10.20 Freestream Pressure

**Scope:** Aero Heat Flux Boundary Condition

Freestream Pressure {=} Value

878
Parameter | Value | Default  
---|---|---
Value | real | 0.0

**Summary**
Specifies a constant freestream pressure.

### 33.10.21 Freestream Pressure Altitude Function

**Scope:** Aero Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specifies the freestream pressure versus altitude function for climate condition.

### 33.10.22 Freestream Pressure Time Function

**Scope:** Aero Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specifies the name of freestream pressure versus time function.

### 33.10.23 Freestream Temperature

**Scope:** Aero Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>real</td>
<td>0.0</td>
</tr>
</tbody>
</table>

**Summary**
Specifies a constant freestream temperature. Usage: ECKERT CONVECTIVE COEFFICIENT.

### 33.10.24 Freestream Temperature Altitude Function

**Scope:** Aero Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specifies the freestream temperature versus altitude function for climate condition. Usage: ECKERT CONVECTIVE COEFFICIENT.
33.10.25  Freestream Temperature Time Function

Scope:  Aero Heat Flux Boundary Condition

Freestream Temperature Time Function {=} Function

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Specifies the name of freestream temperature versus time function. Usage: ECKERT CONVECTIVE COEFFICIENT.

33.10.26  Integrated Flux Output

Scope:  Aero Heat Flux Boundary Condition

Integrated Flux Output VariableName

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>VariableName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Calculate the average flux associated with this flux boundary condition.

Description  This line command specifies that, as a postprocess, the normal flux associated with this boundary condition be integrated over the surface to obtain the total power. This power is then divided by the total area of the surface to obtain the average flux on the surface, and stored in a global variable named "variableName". This global variable may then be output to history files, or accessed in user subroutines, etc.

33.10.27  Integrated Power Output

Scope:  Aero Heat Flux Boundary Condition

Integrated Power Output VariableName

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>VariableName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Calculate the total power associated with this flux boundary condition.

Description  This line command specifies that, as a postprocess, the normal flux associated with this boundary condition be integrated over the surface to obtain the total power which is then stored into a global variable named "variableName". This global variable may then be output to history files, or accessed in user subroutines, etc.

33.10.28  Mach Number

Scope:  Aero Heat Flux Boundary Condition

Mach Number {=} Value
Summary: Specifies a constant Mach number. Usage: ECKERT CONVECTIVE COEFFICIENT.

### 33.10.29 Mach Number Time Function

**Scope:** Aero Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Mach Number Time Function \{=|are|is\} Function**

Summary: Specifies the Mach number versus time function for a given flight. The tabulated time is defined relative to the START TIME. Usage: ECKERT CONVECTIVE COEFFICIENT.

### 33.10.30 Reference Density

**Scope:** Aero Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Reference Density \{=|are|is\} Value**

Summary: Specifies the reference density value for the density ratio heat transfer coefficient model. Usage: DENSITY RATIO CONVECTIVE COEFFICIENT.

### 33.10.31 Reference Htc

**Scope:** Aero Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Reference Htc \{=|are|is\} Value**

Summary: Specifies the reference heat transfer coefficient value for the density ratio heat transfer coefficient model. Usage: DENSITY RATIO CONVECTIVE COEFFICIENT.

### 33.10.32 Reference Temperature

**Scope:** Aero Heat Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Reference Temperature \{=|are|is\} Value**

Summary: Specify a constant reference temperature for this boundary condition.
33.10.33 Reference Temperature Time Function
Scope: Aero Heat Flux Boundary Condition

| Reference Temperature Time Function {\texttt{is}|	exttt{are}} \texttt{FunctionName} |
|---------------------------------|------------------|-----------------|
| Parameter                       | Value            | Default         |
| \texttt{FunctionName}           | \texttt{string}  | \texttt{undefined} |

Summary Specify the name of a time-dependent function for the reference temperature for this boundary condition.

33.10.34 Start Time
Scope: Aero Heat Flux Boundary Condition

| Start Time {\texttt{is}|	exttt{are}} \texttt{Value} |
|---------------------------------|-----------------|-----------------|
| Parameter                       | Value           | Default         |
| \texttt{Value}                  | \texttt{real}   | \texttt{0.0}    |

Summary Specifies the analysis time at which the flight begins and the AERO heat flux will be applied.

33.10.35 Stop Time
Scope: Aero Heat Flux Boundary Condition

| Stop Time {\texttt{is}|	exttt{are}} \texttt{Value} |
|---------------------------------|-----------------|-----------------|
| Parameter                       | Value           | Default         |
| \texttt{Value}                  | \texttt{real}   | \texttt{none}   |

Summary Specifies the analysis time at which the flight ends and the AERO heat flux BC becomes inactive.

33.10.36 Uq Flux Multiplier
Scope: Aero Heat Flux Boundary Condition

| Uq Flux Multiplier {\texttt{is}|	exttt{are}} \texttt{Value} |
|---------------------------------|-----------------|-----------------|
| Parameter                       | Value           | Default         |
| \texttt{Value}                  | \texttt{real}   | \texttt{1.0}    |

Summary Specify constant scaling of the convective flux.

Description Intended use of this scaling parameter is primarily for evaluation of model sensitivities.
33.10.37 Use Toggle Block

Scope: Aero Heat Flux Boundary Condition

Use Toggle Block ToggleName [ {at|for|in|on|over} ElementBlockList... ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>ToggleName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specification for toggling entities in the computational model based on Toggle Block parameters. When used at the region level, the list of element blocks to be toggled must be provided. Otherwise, a listing of entities is not needed as the Toggle Block will be associated with the command line or the enclosing command block.

33.11 Advective Bar

Beta Capability: The advective bar feature has not been fully tested and optimized for production use. Use this capability with caution.

For problems in which convection heat transfer occurs at the surface the analyst must supply some judgment about the fluid flow that gives rise to convection. While this information usually by performing a coupled conjugate heat transfer analysis including details of the fluid flow. In most cases the cost of such an analysis is prohibitive so the analyst resorts to alternative lower order models. One such model, the advective bar, can be applied to cases in which the convective heat transfer is driven by a flow internal to the body of interest as in Figure 33.1. Here a simplified one-dimensional model of an advected flow (advective bar) is directly coupled with the three-dimensional thermal model. The advantage gained by using the simplified velocity representation is that the velocity can be used to define convective heat transfer coefficients appropriate to the problem 34.1.

![Flow](image)

Figure 33.1. Interior Body Flow

33.12 Advective Bar Network

Scope: Equation System

Begin Advective Bar Network Name
Add Volume Parameters...

- Annulus Diameter Ratio Function
- Annulus Diameter Ratio Threshold
- Coordinate Reference
- Entrance Effect Starting Node Id
- Flow Cross Sectional Area Function
- Fluid Density
- Fluid Density Function
- Fluid Viscosity
- Hydraulic Diameter Function
- Initial Temperature
- Mass Flow Rate
- Mass Flow Rate Function
- Maximum Pressure Solve Iterations
- Maximum Search Tolerance
- Pressure Bc
- Pressure Solve Tolerance
- Search Interval
- Solve Pressure
- Surface Roughness
- Velocity Bc
- Visualization
- Wetted Perimeter Function

Summary

Defines an advective bar that can be used by a convective flux boundary condition.

Description

Newton’s law of cooling specifies that the heat flux normal to a surface is proportional to the difference between the unknown temperature of the surface and some reference temperature of the fluid in which the surface is immersed: \( q_n = h(T - T_r) \). The convection coefficient, \( h \), and the reference temperature, \( T_r \), may be specified in several ways. Of particular interest is the situation where the fluid flow can be reduced to a simple 1D flow. In this case, heat transfer between the surface and the fluid is modeled with an advective bar model where each segment on the bar and each facet on the surface assumes a constant heat flux based on a constant \( h, T \) and \( T_r \).

33.12.1 Add Volume

Scope: Advective Bar Network

Add Volume Parameters...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameters</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary

Adds bar element volumes, by name, to a boundary condition’s extent.
### 33.12.2 Annulus Diameter Ratio Function

**Scope:** Advective Bar Network

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary** Specifies the name of a user tabular function for annulus diameter ratio.

---

### 33.12.3 Annulus Diameter Ratio Threshold

**Scope:** Advective Bar Network

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Threshold</td>
<td>real</td>
<td>0.05</td>
</tr>
</tbody>
</table>

**Summary** Specifies the minimum value for annulus diameter ratio. This is used with both automatic calculation as well as user tabular function.

---

### 33.12.4 Coordinate Reference

**Scope:** Advective Bar Network

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nodelist</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>xyzs</td>
<td>{s</td>
<td>x</td>
</tr>
</tbody>
</table>

**Summary** Specifies the reference nodelist and direction for bar coordinates.

---

### 33.12.5 Entrance Effect Starting Node Id

**Scope:** Advective Bar Network

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>NodeID</td>
<td>integer</td>
<td>0</td>
</tr>
</tbody>
</table>

**Summary** Specifies the node in the advective bar that corresponds to the position where entrance effects begin.

**Description** Some of the available heat transfer coefficient correlations (at present the three Gnielinski correlations, types 72-74) support an entrance effect correction factor that will adjust the local heat transfer coefficient based on how far a point is from the inlet of the pipe or annulus. By default the inlet of the pipe is assumed to be at the first node of the advective bar. This...
option lets the user specify a different node ID to use as the pipe entrance for calculating entrance effects. This node ID must be part of the advective bar network.

### 33.12.6 Flow Cross Sectional Area Function

**Scope:** Advective Bar Network

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Specifies the name of a user tabular function for cross-sectional flow area.

### 33.12.7 Fluid Density

**Scope:** Advective Bar Network

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Specifies bar fluid density.

### 33.12.8 Fluid Density Function

**Scope:** Advective Bar Network

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Specifies the name of a user tabular function of temperature for density.

### 33.12.9 Fluid Viscosity

**Scope:** Advective Bar Network

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Specifies bar fluid absolute viscosity.
33.12.10 Hydraulic Diameter Function
Scope: Advective Bar Network

Hydraulic Diameter Function {=} string Function

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specifies the name of a user tabular function for hydraulic diameter.

33.12.11 Initial Temperature
Scope: Advective Bar Network

Initial Temperature {=} real $t$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$t$</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specifies the initial temperature of the bulk node.

33.12.12 Mass Flow Rate
Scope: Advective Bar Network

Mass Flow Rate {=} real $\text{Nodelist}$ Value

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nodelist</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specifies the mass flow rate at a node.

33.12.13 Mass Flow Rate Function
Scope: Advective Bar Network

Mass Flow Rate Function {=} string $\text{Nodelist}$ Function

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nodelist</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Function</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specifies the mass flow rate as a function of time at a node with a user function.

33.12.14 Maximum Pressure Solve Iterations
Scope: Advective Bar Network

Maximum Pressure Solve Iterations {=} iterations

887
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Iterations</em></td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies maximum number of bar pressure solve iterations.

33.12.15 Maximum Search Tolerance

Scope: Advective Bar Network

Maximum Search Tolerance \(=|\text{are}|\text{is} \) *Tolerance*

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Tolerance</em></td>
<td>real</td>
<td>0.5</td>
</tr>
</tbody>
</table>

Summary: Specifies the maximum search tolerance for bar/facet geometric coupling search and this must be between 0 and 1. For an idealized case of a curved bar with a constant curvature \((R)\) and a facet that is at a distance \((d)\) from the bar, the analytical solution tolerance is given by

\[
tol = \frac{d}{2R}
\]

Note that for a straight bar with infinite radius of curvature, this gives a tolerance of 0.

33.12.16 Pressure Bc

Scope: Advective Bar Network

Pressure Bc \(@|\text{at}|\text{for}|\text{in}|\text{on}|\text{over} \) *Nodelist* \(=|\text{are}|\text{is} \) *Pres*

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Nodelist</em></td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td><em>Pres</em></td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies bar inlet pressure.

33.12.17 Pressure Solve Tolerance

Scope: Advective Bar Network

Pressure Solve Tolerance \(=|\text{are}|\text{is} \) *Tolerance*

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Tolerance</em></td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies bar pressure solution convergence tolerance.

33.12.18 Search Interval

Scope: Advective Bar Network

Search Interval \(=|\text{are}|\text{is} \) *Interval*
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Interval</td>
<td>integer</td>
<td>5</td>
</tr>
</tbody>
</table>

**Summary**  Specifies the number of search intervals for bar/facet geometric coupling search. This is used in conjunction with the maximum search tolerance to generate search tolerances from 0.0 to the maximum search tolerance specified. Must be greater than 0.

### 33.12.19 Solve Pressure

**Scope:** Advective Bar Network

**Summary**  Invoke pressure solution. This capability is not currently enabled for split network solutions.

### 33.12.20 Surface Roughness

**Scope:** Advective Bar Network

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Roughness</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  Specifies roughness coefficient of surface.

### 33.12.21 Velocity Bc

**Scope:** Advective Bar Network

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nodelist</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Velocity</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  Specifies bar inlet velocity.

### 33.12.22 Visualization

**Scope:** Advective Bar Network

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>OnOff</td>
<td>{off, on}</td>
<td>Off</td>
</tr>
</tbody>
</table>

**Summary**  Specifies Exodus output toggling of the created subsetted surfaces which are directly affected by the bulk nodes on the advective bar.
33.12.23 Wetted Perimeter Function
Scope: Advective Bar Network

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies the name of a user tabular function for wetted perimeter.

33.13 Radiative Flux Boundary Condition
Scope: Equation System

Begin Radiative Flux Boundary Condition Name

Add Surface SurfaceList...
Emissivity Function Value On SurfaceName
Emissivity Fortran Subroutine Function Name
Emissivity Function FunctionName On SurfaceName
Emissivity Subroutine MySub On SurfaceName
Emissivity Time Function FunctionName On SurfaceName
Equation EquationName
Form Factor Fortran Subroutine Name
Ignore Flux Coverage
Integer Data Values...
Integrated Flux Output VariableName
Integrated Power Output VariableName
Irradiation Node Variable Name
Radiation Form Factor Function Value
Radiation Form Factor Subroutine Name
Radiation Form Factor Temperature Function Name
Radiation Form Factor Time Function Name
Real Data Values...
Reference Temperature Function Value
Reference Temperature Fortran Subroutine Name
Reference Temperature Subroutine Name
Reference Temperature Temperature Function Name
Reference Temperature Time Function FunctionName
Use Bulk Element Name
Use Data Block Name
Use Toggle Block ToggleName ElementBlockList...

End
Summary: Radiative heat flux boundary condition. You must specify either a reference temperature or provide a nodal field for irradiation.

Description: This boundary condition models the radiative heat transfer from a specified surface to a surrounding surface. The temperature of the surrounding surface is assumed to have enough thermal mass that its temperature is unaffected by changes in computed temperature, and is therefore known. The heat flux normal to the surface is given by

\[ q_n = \varepsilon F (\sigma T^4 - G) \]

where the incident radiative flux, or irradiation, \( G \), may either be specified directly as a nodal field, or may be specified through a reference temperature, \( G = \sigma T_r^4 \). The Stefan-Boltzmann constant, \( \sigma \), is defined in a Global Constants command block, \( \varepsilon \) is the surface emissivity, and \( F \) is the form factor.

With the exception of \( \sigma \), each of these parameters may vary in one of several ways, as defined below. The emissivity, if left unspecified, is determined locally from the material associated with the element that underlies each face on the surface. In the case of faces that are created as a result of element death, the emissivity is always determined by the underlying element.

33.13.1 Add Surface

Scope: Radiative Flux Boundary Condition

```
Add Surface SurfaceList...
```

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>SurfaceList</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Adds surfaces, by name, to a boundary condition’s extent.

Description: This line command is used to add surfaces to the extent of a boundary condition. In Exodus II, surfaces are specified as side sets, that have a global integer identifier. For example, side set 12 would be added by this line command using the surface name surface_12. Note that in SIERRA, each element of an array of strings must be separated by whitespace.

33.13.2 Emissivity

Scope: Radiative Flux Boundary Condition

```
Emissivity {=}are|is} Value [ On SurfaceName ]
```

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Sets a constant value of emissivity for a defined surface. If the optional parameters are not included, then this is the default emissivity for the enclosure. Otherwise it is only applied to the indicated surface.

**Note that if a surface is called out more than once, the emissivity definition it is overwritten: last one in wins.
33.13.3 Emissivity Fortran Subroutine

**Scope:** Radiative Flux Boundary Condition

**Emissivity Fortran Subroutine {=|are|is} Name**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specify the name of a FORTRAN user subroutine that will be used to calculate the convective coefficient for this boundary condition.

33.13.4 Emissivity Function

**Scope:** Radiative Flux Boundary Condition

**Emissivity Function {=|are|is} FunctionName [ On SurfaceName ]**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>FunctionName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Sets a emissivity function for a defined surface. If the optional parameters are not included, then this is the default emissivity for the enclosure. Otherwise it is only applied to the indicated surface.

**Note that if a surface is called out more than once, the emissivity definition it is overwritten: last one in wins.**

33.13.5 Emissivity Subroutine

**Scope:** Radiative Flux Boundary Condition

**Emissivity Subroutine {=|are|is} MySub [ On SurfaceName ]**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>MySub</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Sets a emissivity user subroutine for a defined surface. If the optional parameters are not included, then this is the default emissivity for the enclosure. Otherwise it is only applied to the indicated surface.

Also, the software supports using locally scoped user data for most user subroutines, but I haven’t figured out a syntax for it here yet. So it is not yet supported. If you need to get data into this subroutine, use the region’s "REAL DATA" and "INTEGER DATA" line commands.

**Note that if a surface is called out more than once, the emissivity definition it is overwritten: last one in wins.**

33.13.6 Emissivity Time Function

**Scope:** Radiative Flux Boundary Condition
Emissivity Time Function \(=|are|is\) FunctionName [ On SurfaceName ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>FunctionName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Sets a emissivity function of time for a defined surface. If the optional parameters are not included, then this is the default emissivity for the enclosure. Otherwise it is only applied to the indicated surface.

**Note that if a surface is called out more than once, the emissivity definition it is overwritten: last one in wins.**

33.13.7 Equation
Scope: Radiative Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>EquationName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
This command can be used to apply the Radiative Flux conditions for an equation other than the ENERGY equation.

33.13.8 Form Factor Fortran Subroutine
Scope: Radiative Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Specify the name of a FORTRAN user subroutine that will be used to calculate the form factor for this boundary condition.

33.13.9 Ignore Flux Coverage
Scope: Radiative Flux Boundary Condition

Summary
This command causes the code to ignore the flux coverage Field when contact is present. Thus the flux will be applied even if the BC is being set on a contact surface.

33.13.10 Integer Data
Scope: Radiative Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Values</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

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Summary

List of integer data values to be used by the FORTRAN user subroutine. Copies of these values are provided to the subroutine hence changes to these values within the subroutine are not saved.

33.13.11 Integrated Flux Output
Scope: Radiative Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>VariableName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary

Calculate the average flux associated with this flux boundary condition.

Description

This line command specifies that, as a postprocess, the normal flux associated with this boundary condition be integrated over the surface to obtain the total power. This power is then divided by the total area of the surface to obtain the average flux on the surface, and stored in a global variable named "variableName". This global variable may then be output to history files, or accessed in user subroutines, etc.

33.13.12 Integrated Power Output
Scope: Radiative Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>VariableName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary

Calculate the total power associated with this flux boundary condition.

Description

This line command specifies that, as a postprocess, the normal flux associated with this boundary condition be integrated over the surface to obtain the total power which is then stored into a global variable named "variableName". This global variable may then be output to history files, or accessed in user subroutines, etc.

33.13.13 Irradiation Node Variable
Scope: Radiative Flux Boundary Condition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary

Specifies the name of the node variable in which the incident radiative flux associated with this boundary condition is stored.
Description  The indicated node variable must be a valid user variable. Typically, this variable is defined by the user within the Region scope. The node variable is interpolated to the integration points during the integration of the flux term. Typically, this variable would be calculated in another SIERRA region, e.g., Fuego, and populated by transfer in the appropriate Region.

33.13.14 Radiation Form Factor
Scope: Radiative Flux Boundary Condition

<table>
<thead>
<tr>
<th>Radiation Form Factor {=</th>
<th>are</th>
<th>is} Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter Value</td>
<td>Value</td>
<td>Default</td>
</tr>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Specifies a constant value for the radiation form factor for this boundary condition.

Description  For two surfaces, the BC surface (1) and far-field surface (2), ε denoting emissivity, A denoting area, and \( F_{12} \) is the view factor.

\[
F = \begin{cases} 
1 & \text{if } A_2 \gg A_1 \\
F_{12}/[(1 - \varepsilon_1)F_{12} + \varepsilon_1] & \text{for } \varepsilon_1 = 1 \text{ and } A_2 \not\gg A_1 \\
F_{12} & \text{if } \varepsilon_1 = \varepsilon_2 = 1
\end{cases}
\]

33.13.15 Radiation Form Factor Subroutine
Scope: Radiative Flux Boundary Condition

<table>
<thead>
<tr>
<th>Radiation Form Factor Subroutine {=</th>
<th>are</th>
<th>is} Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter Name</td>
<td>Value</td>
<td>Default</td>
</tr>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Specifies the name of a user subroutine that is used to calculate the radiation form factor associated with this boundary condition.

33.13.16 Radiation Form Factor Temperature Function
Scope: Radiative Flux Boundary Condition

<table>
<thead>
<tr>
<th>Radiation Form Factor Temperature Function {=</th>
<th>are</th>
<th>is} Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter Name</td>
<td>Value</td>
<td>Default</td>
</tr>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Specifies the name of a temperature-dependent function that is used to calculate the radiation form factor associated with this boundary condition.

33.13.17 Radiation Form Factor Time Function
Scope: Radiative Flux Boundary Condition
**33.13.18 Real Data**

**Scope:** Radiative Flux Boundary Condition

**Real Data Values...**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Values</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

List of real data values to be used by the FORTRAN user subroutine. Copies of these values are provided to the subroutine hence changes to these values within the subroutine are not saved.

---

**33.13.19 Reference Temperature**

**Scope:** Radiative Flux Boundary Condition

**Reference Temperature {=|are|is} Value**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Specify a constant reference temperature for this boundary condition.

---

**33.13.20 Reference Temperature Fortran Subroutine**

**Scope:** Radiative Flux Boundary Condition

**Reference Temperature Fortran Subroutine {=|are|is} Name**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Specifies the name of a FORTRAN user-defined subroutine that will be used to calculate the reference temperature associated with this boundary condition.

---

**33.13.21 Reference Temperature Subroutine**

**Scope:** Radiative Flux Boundary Condition

**Reference Temperature Subroutine {=|are|is} Name**
33.13.22 Reference Temperature Temperature Function

Scope: Radiative Flux Boundary Condition

Reference Temperature Temperature Function \{=|are|is\} Name

Summary: Specifies the name of the temperature-dependent function that is to be used to calculate the reference temperature associated with this boundary condition.

33.13.23 Reference Temperature Time Function

Scope: Radiative Flux Boundary Condition

Reference Temperature Time Function \{=|are|is\} FunctionName

Summary: Specify the name of a time-dependent function for the reference temperature for this boundary condition.

33.13.24 Use Bulk Element

Scope: Radiative Flux Boundary Condition

Use Bulk Element Name

Summary: Use the named bulk element to model the reference temperature.

Description: This line command specifies the name of a bulk fluid element that has been defined using the bulk fluid command block. The temperature of the bulk fluid element is used as the reference temperature, $T_r$, for the convective heat transfer. Note that it is illegal to specify both a reference temperature and a bulk element.

33.13.25 Use Data Block

Scope: Radiative Flux Boundary Condition
### 33.13.26 Use Toggle Block

**Scope:** Radiative Flux Boundary Condition

**Use Toggle Block**

```
Use Toggle Block ToggleName [ @{at|for|in|on|over} ElementBlockList... ]
```

**Parameter**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>ToggleName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Specification for toggling entities in the computational model based on Toggle Block parameters. When used at the region level, the list of element blocks to be toggled must be provided. Otherwise a listing of entities is not needed as the Toggle Block will be associated with the command line or the enclosing command block.

### 33.14 User Variable

**Scope:** Equation System

**Begin User Variable**

```
Begin User Variable Name
    Add Part part...
    Get Global From Region other_region RemoteVariable Index
    Global Operator {=}|are|is} UserVarOperator
    Initial Value {=}|are|is} Value...
    Read Variable {=}|are|is} Variable [ At Time TimeValue ]
    Reset On Assign
    Type {=}|are|is} UserVarMeshObject UserVarType [ Length {=}|are|is} Length ]
    Use With Restart
End
```

**Summary**

Defines user-named registered variables for this region. The name of the variable is specified by the line parameter ‘name’.

**Description**

This command block allows the user to define their own variables, which have the same status as pre-existing variables. For example, they may be output in the same way as other Aria variables, and may participate in SIERRA restart services. This feature is useful when a user has a need to store data on each mesh object (e.g. nodes) or wishes to evaluate a scalar quantity that requires parallel reduction across processors (e.g. a sum). Typically, a user variable would be set by a SIERRA framework transfer service from another region, or via user subroutines.
33.14.1 Add Part

Scope: User Variable

Add Part *part*...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>part</em></td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Adds surface, block or mesh group, by name, to a user variable's extent.

Description: This line command is used to add surfaces or blocks to the extent of a user variable. In Exodus II, surfaces are specified as side sets, that have a global integer identifier. For example, side set 12 would be added by this line command using the surface name surface_12. Similarly, element blocks also have a global identifier. Note that in SIERRA, each element of an array of strings must be separated by whitespace.

33.14.2 Get Global From Region

Scope: User Variable

Get Global From Region *other_region* *RemoteVariable* *Index*

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>other_region</em></td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td><em>RemoteVariable</em></td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td><em>Index</em></td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies the user global variable value will be assigned the value of the named RemoteVariable retrieved from another application Region. Index is an offset into the RemoteVariable.

33.14.3 Global Operator

Scope: User Variable

Global Operator {*|=|are|is}* *UserVarOperator*

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>UserVarOperator</em></td>
<td>{max</td>
<td>min</td>
</tr>
</tbody>
</table>

Summary: REQUIRED for GLOBAL variables. Since global variables are often used in user subroutines a need arises in parallel simulations to resolve a representative single GLOBAL variable value across processors. The choice of operator determines how the single global value is to be resolved from each of the individual processor local values of the variable.

For SUM the local global variable values on each processor are summed to obtain a single value of the global variable.

For MIN the local global variable values on each processor are compared and the minimum value over all processors is assigned to the global variable.

For MAX the local global variable values on each processor are compared and the maximum value over all processors is assigned to the global variable.

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Parallel synchronization of the single global variables will occur both at the beginning of a nonlinear solution iteration and at the completion of a nonlinear step. Once the synchronization has occurred the local global variable is reset to its initial value.

### 33.14.4 Initial Value

**Scope:** User Variable

**Initial Value** `{=|are|is} Value...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Optional. Specifies the initial value of the user variable. The number of initial values specified with this line command must match the length of the variable specified in the "TYPE" command. If this command is present the specified initial value will automatically be applied to the user variable.

### 33.14.5 Read Variable

**Scope:** User Variable

**Read Variable** `{=|are|is} Variable [ At Time TimeValue ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Variable</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Requests that the variable will be assigned from a time plane of the input ExodusII database. If an AT TIME is not specified, the time plane defaults to the last time plane in the database.

### 33.14.6 Reset On

**Scope:** User Variable

**Summary**
Specifies when the user global variable will be reset. By default global variable will be reset after the nonlinear step has converged.

### 33.14.7 Type

**Scope:** User Variable

**Type** `{=|are|is} UserVarMeshObject UserVarType [ Length `{=|are|is} Length ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>UserVarMeshObject</td>
<td>{edge</td>
<td>element</td>
</tr>
<tr>
<td>UserVarType</td>
<td>{integer</td>
<td>real</td>
</tr>
</tbody>
</table>

**Summary**
This line command is required. The type of user variable is either real or integer, and is either a single, global value, or is associated with each mesh object of the specified kind. Global
variables must have a 'GLOBAL OPERATOR' command associated with them to define its synchronization. The default length is 1. The default initial value for global variables is 0.

### 33.14.8 Use With Restart

**Scope:** User Variable

**Summary**

Optional. The default behavior is for user-defined variables to not be preserved across a restart. The presence of this line command specifies that the variable is to be written to and restored from the restart file.

### 33.15 Volume Heating

**Scope:** Equation System

```plaintext
Begin Volume Heating Name
    Add Volume VolumeList...
    Chemeq {=|are|is} Model
    Element Subroutine {=|are|is} Name
    Element Variable {=|are|is} Name
    Field Scaling {=|are|is} Scale_factor
    Integer Data Values...
    Integrated Power Output VariableName
    Nodal Variable {=|are|is} Name
    Real Data Values...
    Scaling With Global Variable {=|are|is} name
    Source Fortran Subroutine {=|are|is} Name
    Temperature Function {=|are|is} Function
    Time Function {=|are|is} Function
    Use Data Block Name
    Use Toggle Block ToggleName [ {@|at|for|in|on|over} ElementBlockList... ]
    Use Verdi
    Value {=|are|is} vhs
    Verdi Qdot Scaling {=|are|is} Scale_factor For Keyval...
End
```

**Summary**

Defines a volumetric heat source. Exactly one heating type specification is allowed.

**Description**

The volumetric heating must be applied to at least one volume. If a user subroutine is specified, then real and integer data may be declared that will be local in scope to this instance of the volume heating.
33.15.1 Add Volume
Scope: Volume Heating

Add Volume VolumeList...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>VolumeList</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies the mesh volume to which this code feature is to be applied. More than one volume name may be specified, as a space-delimited list.

33.15.2 Chemeq
Scope: Volume Heating

Chemeq {=|are|is} Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies that volume heating is due to chemical reaction. Here MODEL pertains to the paired CHEMEQ MODEL and CHEMEQ SOLVER command blocks that define the chemical system and how it can evolve. Chemical heating source contributions are further controlled through parameters in the CHEMEQ SOLVER command block.

33.15.3 Element Subroutine
Scope: Volume Heating

Element Subroutine {=|are|is} Name

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies that the named user-defined subroutine be used. At most one subroutine name may be specified for a given quantity. This subroutine must conform to the "element signature" argument list type.

33.15.4 Element Variable
Scope: Volume Heating

Element Variable {=|are|is} Name

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies the name of an element variable in which the volumetric heat source is stored. The units of this term are power per unit volume. Currently, this must be constant in space over each finite element.
33.15.5 Field Scaling
Scope: Volume Heating

Field Scaling \(\text{=} \text{are}\text{is} \) Scale\_factor

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Scale_factor</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
This command is only relevant when the NODAL VARIABLE or ELEMENT VARIABLE command line option is present. The command enables scaling of the element Field for the element blocks within this Volume Heat command block.

33.15.6 Integer Data
Scope: Volume Heating

Integer Data Values...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Values</td>
<td>integer...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
List of integer data values to be used by the FORTRAN user subroutine. Copies of these values are provided to the subroutine hence changes to these values within the subroutine are not saved.

33.15.7 Integrated Power Output
Scope: Volume Heating

Integrated Power Output VariableName

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>VariableName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Calculate the total power associated with this flux boundary condition.

Description
This line command specifies that, as a postprocess, the normal flux associated with this boundary condition be integrated over the surface to obtain the total power which is then stored into a global variable named "variableName". This global variable may then be output to history files, or accessed in user subroutines, etc.

33.15.8 Nodal Variable
Scope: Volume Heating

Nodal Variable \(\text{=} \text{are}\text{is} \) Name

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Summary   Specifies the name of a nodal variable in which the volumetric heat source is stored. The units of this term are power per unit volume.

Description   The nodal variable could be user defined, which could originate from a transfer in the case of coupled mechanics, or from some user subroutine. The nodal values are interpolated to the Gauss points and then integrated over each element.

33.15.9   Real Data
Scope:   Volume Heating

<table>
<thead>
<tr>
<th>Real Data Values...</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter</td>
</tr>
<tr>
<td>Values</td>
</tr>
</tbody>
</table>

Summary   List of real data values to be used by the FORTRAN user subroutine. Copies of these values are provided to the subroutine hence changes to these values within the subroutine are not saved.

33.15.10   Scaling With Global Variable
Scope:   Volume Heating

| Scaling With Global Variable {=|are|is} name |
|---------------------|
| Parameter   | Value   | Default   |
| name        | string  | undefined |

Summary   This command is only relevant when the NODE VARIABLE or ELEMENT VARIABLE command line option is present. The command enables scaling of the source field by a global variable for the surfaces within the Volume Heat command block. The scalar global variable itself must be defined at the Region scope as type Real with length 1.

33.15.11   Source Fortran Subroutine
Scope:   Volume Heating

| Source Fortran Subroutine {=|are|is} Name |
|---------------------|
| Parameter   | Value   | Default   |
| Name        | string  | undefined |

Summary   Specify the name of a FORTRAN user subroutine that will be used to calculate the volumetric source.

33.15.12   Temperature Function
Scope:   Volume Heating
### Temperature Function \{=|are|is\} Function

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specifies a temperature dependent function name for a volumetric heat source to be applied in the named volume.

#### 33.15.13 Time Function

**Scope:** Volume Heating

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specifies a time function dependent name for a volumetric heat source to be applied in the named volume.

#### 33.15.14 Use Data Block

**Scope:** Volume Heating

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Reference to predefined data to be used by the user subroutine. These values may be changed by the user subroutine.

#### 33.15.15 Use Toggle Block

**Scope:** Volume Heating

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>ToggleName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specification for toggling entities in the computational model based on Toggle Block parameters. When used at the region level, the list of element blocks to be toggled must be provided. Otherwise a listing of entities is not needed as the Toggle Block will be associated with the command line or the enclosing command block.

#### 33.15.16 Use Verdi

**Scope:** Volume Heating

905
Summary: Specifies that the external Goodyear code VERDI should be used to calculate the volumetric heating source term. The value is assumed to be constant in space over each finite element. WARNINGS: There are some caveats: the code will abort (1) with multi-processor runs, since Verdi is inherently serial; (2) with 3-d simulations, since Verdi expects a 2D heat problem to be solved; (3) whenever mesh adaptivity is in use, since Verdi uses three different meshes in its work that must remain very closely related; and (4) whenever Verdi is called from an executable other than eagle (including Calore), since only eagle links against Verdi.

### 33.15.17 Value

**Scope:** Volume Heating

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \text{vhs} )</td>
<td>\text{real}</td>
<td>\text{undefined}</td>
</tr>
</tbody>
</table>

**Summary:** Specifies a constant value for a volumetric heat source to be applied in the named volume.

### 33.15.18 Verdi Qdot Scaling

**Scope:** Volume Heating

**Verdi Qdot Scaling \( \text{Scale\_factor} \text{ For Keyval...} \)**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \text{Scale_factor} )</td>
<td>\text{real}</td>
<td>\text{undefined}</td>
</tr>
<tr>
<td>( \text{Keyval} )</td>
<td>\text{string...}</td>
<td>\text{undefined}</td>
</tr>
</tbody>
</table>

**Summary:** This command is only relevant when the USE VERDI command line option is present. The command enables scaling of the element Qdot scale factor for a collection of element blocks. This command can appear within the command block several times with different scaling factors for each set of elements. Qdot scaling for element blocks not appearing in a VERDI QDOT SCALING command line is set to unity.

### 33.16 Laser Heating

**Scope:** Equation System

**Begin Laser Heating** \( \text{Name} \)

- **Activation Temperature \( \text{Value} \)**
- **Add Volume** \( \text{VolumeList...} \)
- **Aft Semi Axis \( \text{aft\_semiaxis} \)**
- **Beam Diameter \( \text{laser\_diameter} \)**
- **Depth Direction \( \text{Function} \)**
- **Depth Semi Axis \( \text{depth\_semiaxis} \)**
- **Distribution \( \text{LaserSourceDistributionType} \)**

906
Efficiency = Efficiency
Integrated Power Output = VariableName
Path Function = Function
Power = laser_power
Power Encore Function = Function
Power Time Function = Function
Source Type = LaserSourceType
Spatial Influence Factor = Value
Speed = EquationName
Start Time = Value
Stop Time = Value
Travel Semi Axis = travel_semiaxis
Use Toggle Block [ ElementBlockList... ]
Width Semi Axis = width_semiaxis

Summary
Defines a moving laser volumetric heat source. Exactly one heating type specification is allowed.

Description
The laser volumetric heating must be applied to at least one volume. If a user subroutine is specified, then real and integer data may be declared that will be local in scope to this instance of the volume heating.

33.16.1 Activation Temperature
Scope: Laser Heating

Activation Temperature = Value

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Activation Temperature</td>
<td>real</td>
<td>Real_Max</td>
</tr>
</tbody>
</table>

Summary
Defines temperature at which an element may be activated.

Description
Elements within a geometric region bounded using SPATIAL INFLUENCE FACTOR will be activated when this temperature is exceeded. Used in conjunction with ACTIVATION_HEMISPHERE or ACTIVATION_SPHERES in addition to the ACTIVATION_USER_FUNCTION thermal conductivity model.

33.16.2 Add Volume
Scope: Laser Heating

Add Volume VolumeList...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>VolumeList</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Summary  Specifies the mesh volume to which this code feature is to be applied. More than one volume name may be specified, as a space-delimited list.

33.16.3  Aft Semi Axis
Scope:  Laser Heating

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>aft_semiaxis</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Defines the aft direction semiaxis for the double ellipsoidal and double ellipsoidal laser source model.

33.16.4  Beam Diameter
Scope:  Laser Heating

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>laser_diameter</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Specifies laser spot size.

33.16.5  Depth Direction
Scope:  Laser Heating

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Specifies the depth coordinate direction, X, Y, or Z, the directional axis in which laser is applied.

33.16.6  Depth Semi Axis
Scope:  Laser Heating

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>depth_semiaxis</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Defines the depth direction semiaxis for the ellipsoidal and double ellipsoidal source model.
### 33.16.7 Distribution

**Scope:** Laser Heating

**Summary**

Specifies the spatial distribution of applied laser energy.

**Description**

Defines how the laser energy is deposited onto the source geometry. Generally speaking the spatial distribution is GAUSSIAN centered on the source point but for fiber bundles one may opt for the top hat or UNIFORM distribution.

**Parameter** | **Value** | **Default**
--- | --- | ---
*LaserSourceDistributionType* | \{gaussian|uniform\} | GAUSSIAN

---

### 33.16.8 Efficiency

**Scope:** Laser Heating

**Summary**

Defines the transmission efficiency of the available laser power.

**Parameter** | **Value** | **Default**
--- | --- | ---
*Efficiency* | real | undefined

---

### 33.16.9 Integrated Power Output

**Scope:** Laser Heating

**Summary**

Calculate the total power associated with this flux boundary condition.

**Description**

This line command specifies that, as a postprocess, the normal flux associated with this boundary condition be integrated over the surface to obtain the total power which is then stored into a global variable named "variableName". This global variable may then be output to history files, or accessed in user subroutines, etc.

**Parameter** | **Value** | **Default**
--- | --- | ---
*VariableName* | string | undefined

---

### 33.16.10 Path Function

**Scope:** Laser Heating

**Summary**

Specifies a time and position function for which a laser source is applied in the named volume.
33.16.11 Power

Scope: Laser Heating

Power \(\text{is} laser\_power\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>laser_power</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies strength of the laser source.

33.16.12 Power Encore Function

Scope: Laser Heating

Power Encore Function \(\text{is} Function\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies that power be applied with the named Encore function in the named volume.

33.16.13 Power Time Function

Scope: Laser Heating

Power Time Function \(\text{is} Function\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies that power be applied as a function of time in the named volume.

33.16.14 Source Type

Scope: Laser Heating

Source Type \(\text{is} LaserSource\_Type\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>LaserSource_Type</td>
<td>activation_hemisphere</td>
<td>activation_sphere</td>
</tr>
</tbody>
</table>

Summary: Selects the laser source model applied to a portion of the underlying volume as defined by a geometric function. When used with a Gaussian DISTRIBUTION all source types produce maximum intensity at the center of the laser spot. Aside from the ACTIVATION SPHERE model these sources are currently limited to paths aligned with the x,y,z coordinate directions.
### 33.16.15 Spatial Influence Factor

**Scope:** Laser Heating

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spatial Influence Factor</td>
<td>=</td>
<td>real 0.0</td>
</tr>
</tbody>
</table>

**Summary**

Defines a distance from beam focus for which an element may be activated.

**Description**

A multiplier which expands the bounding source geometry. When used with ACTIVATION TEMPERATURE and ACTIVATION_USER_FUNCTION thermal conductivity elements within bounded region become candidates for temperature activation. Used in conjunction with ACTIVATION_HEMISPHERE or ACTIVATION_SPHERE source models.

### 33.16.16 Speed

**Scope:** Laser Heating

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Speed</td>
<td>=</td>
<td>string undefined</td>
</tr>
</tbody>
</table>

**Summary**

Specifies the rate at which the laser beam moves across the body.

**Description**

The speed at which the beam moves in traversing the specified geometric path. When a geometric path function is provided the speed is internally computed based upon the START TIME and STOP TIME.

### 33.16.17 Start Time

**Scope:** Laser Heating

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Start Time</td>
<td>=</td>
<td>real undefined</td>
</tr>
</tbody>
</table>

**Summary**

Specifies the time at which the laser beam is activated.

### 33.16.18 Stop Time

**Scope:** Laser Heating

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stop Time</td>
<td>=</td>
<td>real undefined</td>
</tr>
</tbody>
</table>

**Summary**

Specifies the time at which the laser beam is deactivated.
33.16.19  Travel Semi Axis
Scope:  Laser Heating

Travel Semi Axis {=} are is travel_semiaxis

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>travel_semiaxis</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Defines the travel direction semiaxis for the ellipsoidal and double ellipsoidal source model.

33.16.20  Use Toggle Block
Scope:  Laser Heating

Use Toggle Block ToggleName [ {@|at|for|in|on|over} ElementBlockList... ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>ToggleName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Specification for toggling entities in the computational model based on Toggle Block parameters. When used at the region level, the list of element blocks to be toggled must be provided. Otherwise a listing of entities is not needed as the Toggle Block will be associated with the command line or the enclosing command block.

33.16.21  Width Semi Axis
Scope:  Laser Heating

Width Semi Axis {=} are is width_semiaxis

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>width_semiaxis</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Defines the width direction semiaxis for the ellipsoidal and double ellipsoidal laser source model.

33.17  Closed Surface Volume
Scope:  Equation System

Begin Closed Surface Volume ModelName
   Add Surface SurfaceList...
   Utility Group {=} are is Frequency
End

Summary  This command block defines the parameters for computation of the volume of a closed surface. The computed volume is available as a global variable with name modelName.
Description
Defines a means for computing the volume of a closed surface and associate it with name modelName.

33.17.1 Add Surface
Scope: Closed Surface Volume

Add Surface SurfaceList...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>SurfaceList</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
Adds surfaces by name, (surface_id), to a mesh extent list.

Description
This line command is used to add surfaces to a mesh extent list. In Exodus II, surfaces are specified as side sets, that have a global integer identifier. For example, side set 12 would be added by this line command using the surface name surface_12. Note that in SIERRA, each element of an array of strings must be separated by whitespace.

33.17.2 Utility Group
Scope: Closed Surface Volume

Utility Group {=|are|is} Frequency

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Frequency</td>
<td>{begin_nonlinear_solve</td>
</tr>
</tbody>
</table>

Default undefined

Summary
Sets the point in the execution to evaluate utility

33.18 Real and Integer Data Access - Data Block

Aria provides a means by which a user can define problem dependent real and integer data to be used by user subroutines. General commands for usage of this capability are described in the chapter on Data Block 7. For users of the Calore code, these data blocks occupy the role formerly served by REAL DATA and INTEGER DATA commands in that code while providing some additional flexibility. Data Blocks can used both within Calore-style boundary condition, initial condition, volume heat command blocks as well as in Calore-style user subroutines.

Data Block information can be accessed by Calore-style subroutines in two ways. The information can be directly accessed through user query functions referencing the Data Block by name within a user subroutine or referencing indirectly by implicit association with a command block, again through user queries.
In either event, data extraction within the user subroutine is accomplished in the same manner but using different interfaces. Accessing Data Block values from a user subroutine involves a simple three-step extraction sequence

1. Define a variable of data type associated with the values one wishes to extract from the Data Block, RealArray1d, RealArray2d, IntegerArray1d or IntegerArray2d.

2. Define a string with the name of the values to be extracted.

3. Perform a user_query request to retrieve the data from the Data Block.

Usage of this information within a user subroutine is best described using simple examples that use single values from the data block. What remains here is to describe the various user_query requests of data block information.

### 33.18.1 Data Block Referenced From Command Block

In order to reference a Data Block from a command block the Data Block must first be bound to the scope of the command block by including the `USE DATA BLOCK` directive within the scope of the command block. The choice of scope will dictate the type of user query function used to access the Data Block information.

**Data Block Referenced From Region Command Block**

When the Data Block is defined within the Region scope this information can be made available to the user subroutine system by supplying the `USE DATA BLOCK` directive at the Region scope. From the user subroutine, this data can then be retrieved for usage without knowledge of the data block name. Let the input file contain a data block definition within the Region scope:

```plaintext
Begin Procedure my_procedure
  .
  begin data block flux_data
    real flux = -0.05
  end data block flux_data
  .
  Begin Aria Region myregion
    .
    use data block flux_data
    .
    Begin heat flux boundary condition freddy
      add surface surface_4
      element subroutine = aucal_heat_flux
    End
    .
  End Aria Region

End Procedure
```

Within the user subroutine source code the values can then be retrieved from the Data Block by following the extraction sequence targeting Region data.
RealArray1d real_array;
sierra::String flux_label("flux");

user_query.getUserRealRegionData(real_array, flux_label);

Real flux_value = real_array(0);

Here we note the presence of Region in the user_query and the user subroutine becomes

```cpp
int aucal_heat_flux(
    UserQuery & user_query,
    ElementIds element_ids,
    Int num_elements,
    Int num_integration_points,
    Int spatial_dimension,
    CoordinateElementIntegrationPoints coordinates,
    TemperatureElementIntegrationPoints temperature,
    ScalarElementIntegrationPoints scalar)
{
    RealArray1d real_array;
    sierra::String flux_label("flux");

    user_query.getUserRealRegionData(real_array, flux_label);

    Real flux_value = real_array(0);

    for (int q = 0; q < num_integration_points; ++q)
    {
        for (int i = 0; i < num_elements; ++i)
        {
            scalar(i, q) = flux_value;
        }
    }
    return 0;
}
```

Here we note that the Data Block name, flux_data, appeared only at the Region scope of the input.

Data Block Referenced From BC, IC or Volume Source Command Block

Data Block information can be made accessible to boundary condition, initial condition or volumetric source user subroutines defined within command blocks. This is accomplished by simply supplying the USE DATA BLOCK directive at the command block scope. Returning to the example, the USE DATA BLOCK enables the association of flux_data with the aucal_heat_flux user subroutine.

Begin Procedure my_procedure
```
    begin data block flux_data
        real flux = -0.05
    end data block flux_data
```
Within the user subroutine source code the values can then be retrieved from the Data Block by following the extraction sequence

```c++
RealArray1d real_array;
sierra::String flux_label("flux");
user_query.getUserRealInstanceData(real_array, flux_label);
Real flux_value = real_array(0);
```

Here we note the presence of `Instance` in the `user_query` and the user subroutine becomes

```c++
int aucal_heat_flux(
    UserQuery & user_query,
    ElementIds element_ids,
    Int num_elements,
    Int num_integration_points,
    Int spatial_dimension,
    CoordinateElementIntegrationPoints coordinates,
    TemperatureElementIntegrationPoints temperature,
    ScalarElementIntegrationPoints scalar)
{
    RealArray1d real_array;
    sierra::String flux_label("flux");
    user_query.getUserRealInstanceData(real_array, flux_label);
    Real flux_value = real_array(0);
    for (int q = 0; q < num_integration_points; ++q)
    {
        for (int i = 0; i < num_elements; ++i)
        {
            scalar(i, q) = flux_value;
        }
    }
    return 0;
}
```
Data Block Referenced From Material Command Block

The Data Block information can be made automatically available for material property evaluations employing user subroutines.

```plaintext
Begin Procedure my_procedure
  begin data block k_data
    real my_K = 4.0
  end data block k_data

  begin Aria material my_material
    use data block flux_data
    thermal conductivity = calore_user_sub name = aucal_elem_cond type=element
  end Aria material my_material

  Begin Aria Region myregion

  End Aria Region

End Procedure
```

Within the user subroutine source code the values can then be retrieved from the Data Block by following the extraction sequence

```plaintext
RealArray1d real_array;
sierra::String k_label("my_k");

user_query.getUserRealMaterialData(real_array, k_label);

Real k_value = real_array(0);
```

Here we note the presence of `Material` in the `user_query` and the user subroutine becomes

```plaintext
int aucal_elem_cond(
    UserQuery & user_query,
    ElementIds element_ids,
    Int num_elements,
    Int num_integration_points,
    Int spatial_dimension,
    CoordinateElementIntegrationPoints coordinates,
    TemperatureElementIntegrationPoints temperature,
    ScalarElementIntegrationPoints cond)
{

RealArray1d real_array;
sierra::String k_label("my_k");

user_query.getUserRealMaterialData(real_array, k_label);

Real k_value = real_array(0);

for (int q = 0; q < num_integration_points; ++q)
{
    for (int i = 0; i < num_elements; ++i)
    {
        cond(i, q) = k_value(i);
    }
}
return 0;

The previous examples demonstrated association of Data Block information with the scope of a command block in order to avoid explicit specification of the Data Block name. However, alternative access methods also allow one to retrieve Data Block information more directly by explicitly supplying the Data Block name in the user_query operation rather than by supplying the *USE DATA BLOCK* directive at a given scope. A complete list of the various query operations is included in the section on For the examples previously described this style of queries translate to

```
user_query.getUserRealData(real_array, flux_label,"flux_data");
and
user_query.getUserRealData(real_array, k_label,"k_data");
```

For the simple examples provided the type of user_query is a personal choice. However, an overall strategy for usage of Data Block information within user subroutines often dictates the type of user_query employed. Considerations in this strategy often include the number of subroutines that will be supported and how the subroutines are managed. As an example one could configure all input files in the same way and selectively load subroutines that employ the same interface. Again we emphasize that the different methods of accessing Data Block information differ only by the interface used to obtain the data definition.

### 33.18.2 Data Block Referenced By Line Command

For commands not defined within a Command Block the Data Block can still be used to specify parameters for that line command using the *USE DATA BLOCK* directive. The data block parameters are placed in the line command as if it was a command block.

Creating the data block in the region using the data block command block:

```
begin data block bc_flux
    real h = 300.0
    integer t_ref = 1
end
```

its data can be accessed using a command line like

```
BC Flux for Energy on no_slip_boundary = Nat_Conv use data block bc_flux
```
which is equivalent to

BC Flux for Energy on no_slip_boundary = Nat_Conv H=300.0  T_ref=1

This architecture allows the data to be consistently used with boundary conditions for several equations.

33.19 User Subroutines and Variables

Aria offers support of C language Calore-style user subroutines and FORTRAN user subroutines through fixed-interface signatures. Callback data query requests from the subroutines are also supported, again through fixed-interface function signatures. It is important to note that Calore-style subroutines provide some level of variable-type safety and bounds checking that is not supported with FORTRAN subroutines.

Some of the subroutine interface callback function calls enable writing to user defined node, element or face Fields, which can be used internal to the code or output to the results file. Generally speaking, values made available to a subroutine through its calling interface are quadrature point values. Internally the application code can either process all quadrature points at one time or all at once. Hence in order to save user fields that are populated within the subroutine one should understand exactly how the subroutine is being processed before relying upon data saved from within the subroutine.

Usage of both C language Calore-style 33.20 and FORTRAN subroutines 33.24 within Aria are described in the sections which follow.

33.20 C Calore-Style User Subroutines and Variables

Aria supports the use of C language user subroutines. Historically, the Calore heat transfer code supported only FORTRAN subroutines. In later releases of Calore, support of FORTRAN subroutines has continued but now C language style user functions are supported as well. With only minor syntactical changes these C language user functions support the full functionality of FORTRAN subroutines. When written in a general manner, the C Calore-style user functions can be used in either Calore or Aria. The overall nature of the Calore style user function is described below and is followed by a simple example of its use with the DATA_BLOCK utility. In what follows, the terms "signature", "argument list", and "interface" are synonyms for the list of variables and arrays that are passed into a user subroutine. Furthermore, the term Calore style user subroutine will be loosely associated with the term Calore style user functions.

An Aria Calore-style user subroutine must be a text file of suffix .C. In contrast to the static link approach formerly used in the Calore code, here the file contents are made available to the application as a dynamically linked function using the same procedure as for Aria User Plugin functions 18. For example, a file named Aucal_density.C the file will be referenced from the Aria input file using the line command

load user plugin file ./Aucal_Density.so

Note that, since Aucal_density.C is a filename, its input file name is case-sensitive, and considerations must be made concerning its location path. The user is free to name the file and subroutines in any way, provided the name is consistent with compiler and operating system limitations. For example, special characters such as '*' and ',' should be avoided. Finally actual use of the function by the application is triggered by association of the "model name", calore_user_sub and the user subroutine name in particular command line, i.e. initial condition, source or boundary condition.
Multiple user subroutines for a given parameter, e.g. conductivity, may be defined. If user conductivity for two materials, say gold and silver was needed then one could write a subroutine for each of the two conductivities. Each subroutine must have a different name in order to be properly referenced, and each must conform to the required call signature. This approach eliminates the need for "if-then-else if" subroutine constructs defining specific behavior for a particular material, volume or surface.

Regardless of the information included in the call signature, the need will arise to access data that is not currently provided by the subroutine arguments. Altering the call signature to fulfill that need would require extensive changes to Calore style user subroutines. Furthermore, backward compatibility between code versions would be lost, since all subroutines currently using an older interface would no longer function properly with a newer interface.

For reasons previously stated, the design philosophy behind the user subroutine capability in both Calore and Aria is that the call signature will kept to a minimum. An advantage of this approach is that it provides a clean, simple interface. A disadvantage is that it limits the information that a user may access. This inherent disadvantage is addressed by employing query functions for use within the subroutines. Thus if one wishes to access additional information from the user subroutine, it becomes a relatively straightforward task to implement a new query function to provides the required data. In this way, no changes are required of the subroutine signature, and backward compatibility is maintained. Users having specialized needs of data access should consider implementations of native Aria user plugin functions instead.

An attempt has been made to ease the transition from Calore user subroutines to Calore style user subroutines in Aria by providing common C interfaces to the two codes. Two versions of the subroutine signatures are provided here, the first more reminiscent of a FORTRAN interface and the other being a C language interface.

In many applications several user subroutines are being used in a single simulation. In this regard it is worth mentioning that with both the FORTRAN and C-style usage one must register the various subroutines/functions with the application before they can be recognized and accessed by the application. This registration establishes an association between the a code interface, function signature and the name by which the function will be referenced from the input file command lines. For the Aria C-style subroutines the registration can be done in one of two ways, with individual registration lines for each subroutine or by using a single function to register one or more subroutines.

Subroutine registration amounts to adding a line to the subroutine source file with a line declaring the signature corresponding to the subroutine containing its registered name and a pointer to the user subroutine. As an example, if a user subroutine file named Aucal_Density.C contained a function named my_density and one wished to reference the function as bulk_density one could register the subroutine as:

```c
RegisterFunction(ElemUserSubC, my_density, "bulk_density");
```

Alternatively one could also use a more obscure form of the registration:

```c
sierra::Aria::AcalElemUserSubC<my_density>::Register distr_register("bulk_density");
```

Again we note that both registration methods contain the application interface signature, the function pointer and the name one wishes to associate the function with from the input file. This registration causes the subroutine to be recognized within the function name registry. Since the compiled function name is not available prior to compilation, the registration line should appear after the subroutine function in the subroutine source code.

Access to the subroutine from within the application is enabled using the following command line:

```
load user plugin file ./Aucal_Density.so
```
Subroutine registration using a single function can be performed simply by wrapping the individual registrations previously described in a function. Using the subroutine name previously discussed the corresponding registration using a function call could be obtained using the following module

```c
extern "C"
void
bulk_prop_subs()
{
    RegisterFunction(ElemUserSubC, my_density, "bulk_density");
}
```

As in the case of single line subroutine registration compiled function names are not available prior to compilation, thus the registration function must appear in the subroutine source code after the subroutine function being registered.

Access to the subroutine from within the application is enabled using the following command line:

```text
load user plugin file ./Aucal_Density.so using function bulk_prop_subs
```

One can also avoid using a named registration module by using the default function name `dl_register`,

```c
extern "C"
void
dl_register()
{
    RegisterFunction(ElemUserSubC, my_density, "bulk_density");
}
```

where once again access to the subroutine from within the application is enabled without using a function name

```text
load user plugin file ./Aucal_Density.so
```

Finally in order to access the subroutine from the input file according to its usage from the Aria material command block one would reference the subroutine as

```text
Begin Aria Material bulk
    .
    .
    density = Calore_User_Sub name=bulk_density type = element
    .
End Aria Material bulk
```

### 33.21 FORTRAN-like C Interfaces

While Calore style user subroutines could be defined to supply many different parameters, only five different signatures are provided in Aria. These signatures characterize the argument lists by which the subroutine interacts with the code. The various subroutine signatures are denoted as:
• node signature 33.21.1
• element signature 33.21.2
• scaled element signature 33.21.3
• reference temperature dependent element signature 33.21.4
• scaled, reference temperature dependent element signature 33.21.5
• tensor thermal conductivity signature 33.21.6
• transformed tensor thermal conductivity signature 33.21.7
• reaction rate signature 33.21.8
• auxiliary variable signature 33.21.9.

The node subroutine signature is generally used for Dirichlet boundary conditions. The element signature is used to define material properties, heat flux and heat transfer coefficients. The reference temperature dependent signatures are used primarily for defining heat transfer coefficients. The last two signatures are specific to the CHEMEO chemistry package. Scaled signatures are used to combine the basic user subroutine functionality combined with Field variable scaling. The different subroutine call signatures are outlined below. With the exception of the node signature, these signatures will be used for subroutines that operate on quadrature point values.

It is worth noting that the purpose of the user subroutine is simply to provide an evaluation of needed values based upon the data supplied through the subroutine interface. In many cases the data supplied to the user subroutine is more that what is necessary for the targeted evaluation. Here the user is free to perform the evaluation using only the information needed to perform the target evaluation. Note that only the last argument of the interface is associated with values returned to the application code, the remaining arguments represent non-modifiable data local to the element, face or node being processed.

### 33.21.1 Node Signature

For example, to specify a temperature Dirichlet boundary condition, the node signature is

```c
/**
* Prototype for the nodal signature.
*
* user_query: query function
* node_id: array of global ids of the nodes
* num_nodes: number of nodes in the array
* spatial_dimension: number of nodes in the array
* coordinate_array: array of coordinates of the current nodes
* node_data_array: array of value to be calculated
*/
Int Aca1NodeUserSubC(
    UserQuery & user_query,
    Int node_id,
    Int num_nodes,
    Int spatial_dimension,
    CoordinateNodes coordinate_array,
    DataNodes node_data_array);
```
The node subroutine for a Dirichlet boundary condition could be invoked using a command line:

BC Dirichlet for Temperature on surface_4 = Calore_User_Sub name=dirich_bc type=node

A simple example using the nodal signature subroutine is as follows:

```c
#include <utility>
#include "math.h"
#include "Aria_Calore_User_Sub_Support.h"

int dirichsub(
    UserQuery & user_query,
    Int nodeid,
    Int nnode,
    Int spatial_dimension,
    CoordinateNodes coords,
    DataNodes temperature)
{
    for (int i = 0; i < nnode; ++i) {
        Real ynode = coords(1,i);
        if ( ynode < 0.5 ) {
            temperature(i) = 1.5;
        } else {
            temperature(i) = 0.1;
        }
    }
    return 0;
}

RegisterFunction(NodeUserSubC, dirichsub, "dirich_bc");
```

### 33.2.1.2 Element Signature

For density, specific heat, volumetric heat source, surface heat flux and most surface heat transfer coefficients defined via user subroutine one will use an element based call signature. Additionally, for problems with isotropic thermal conductivity one can supply the single value of conductivity using the element signature. In either case the subroutine will be used to assign quadrature point evaluations. That is, the coordinate_array values can be used to compute node_data_array values The call signature for scalar element-based user subroutines is

```c
/**
 * Prototype for the scalar element signature.
 * user_query: query function
 * elem_ids: pointer to the global id of the current element
 * num_elements: pointer to the number of elements in the workset
 * nint: pointer to the number of gauss points
 * coordinate_array: pointer to the coordinates of the current node (3, nelem, nint)
 * temperature_array: pointer to the temperature (nelem, nint)
 */
```
The scalar element subroutine signature is supported for all element-centric evaluations. An example of specifying the element user subroutine to be used for density is as follows:

```
Begin Aria Material sticky_icky
  density = calore_user_sub name = aucal_density type=element
  specific heat = constant cp = 385.0
  thermal conductivity = constant k = 400.0
  heat conduction = basic
End Aria Material sticky_icky
```

A simple example using the scalar element signature subroutine is as follows:

```c
#include <utility>
#include "math.h"
#include "Aria_Calore_User_Sub_Support.h"

int aucal_density(
  UserQuery & user_query,
  ElementIds elemid,
  Int nelem,
  Int nint,
  Int spatial_dimension,
  CoordinateElementIntegrationPoints coords,
  TemperatureElementIntegrationPoints temp,
  ScalarElementIntegrationPoints rho)
{
  // Load the density array
  for (int j = 0; j < nint; ++j) {
    for (int i = 0; i < nelem; ++i) {
      rho(i,j) = 0.1 + temp(i,j);
    }
  }
  return 0;
}
```

RegisterFunction(ElemUserSubC, aucal_density, "aucal_density");
The same methodology used in the previous example is valid when defining isotropic thermal conductivity that depends upon other local state information. Here all directions are material principal directions so thermal conductivity will be a single value for each quadrature points of an element.

Note that in this example the user subroutine name is specified within quotation marks. As per 2.5.1 the quotation marks are only required when the name by which the subroutine is referenced by from the input file begins with a number. Otherwise quotation marks are not needed in the specification of user subroutine name.

```
BEGIN ARIA MATERIAL my_mat
   density = constant rho = 0.1
   specific heat = constant cp = 385.0
   thermal conductivity = calore_user_sub name="2Dcond" type = element
   heat conduction = basic
END ARIA MATERIAL my_mat
```

For isotropic conductivity an example of the associated user subroutine and its function registration would be:

```c
#include <utility>
#include "math.h"
#include "Aria_Calore_User_Sub_Support.h"

int mycond(
    UserQuery & user_query,
    ElementIds element_ids,
    Int num_elements,
    Int num_integration_points,
    Int spatial_dimension,
    CoordinateElementIntegrationPoints coordinates,
    TemperatureElementIntegrationPoints temperature,
    ScalarElementIntegrationPoints cond)
{
    for (int q = 0; q < num_integration_points; ++q)
    {
        for (int i = 0; i < num_elements; ++i)
        {
            cond(i, q) = 400.0;
        }
    }
    return 0;
}

RegisterFunction(ElemUserSubC, mycond, "2Dcond");
```

### 33.21.3 Scaled Element Signature

A variant of the element subroutine employs an additional Field variable that is supplied through the user interface. Here the Field variable is used to perform some logical decision on how the result is assigned and
the name of the Field variable to be used is specified from the input file. The call signature for scaled scalar element-based user subroutines is

/**
 * Prototype for the scaled scalar element signature.
 *
 * @param user_query query function
 * @param elem_ids pointer to the global id of the current element
 * @param num_elements pointer to the number of elements in the workset
 * @param nint pointer to the number of gauss points
 * @param coordinate_array pointer to the coordinates of the current node (3, nelem, nint)
 * @param temperature_array pointer to the temperature (nelem, nint)
 * @param scaling_array pointer to the scaling array (nelem, nint)
 * @param elem_data_array pointer to the values to be calculated (nelem, nint)
 */

Int AcalScaledElemUserSubC(  
   UserQuery & user_query,  
   ElementIds elem_ids,  
   Int num_elements,  
   Int nint,  
   Int spatial_dimension,  
   CoordinateElementIntegrationPoints coordinate_array,  
   TemperatureElementIntegrationPoints temperature_array,  
   ScalarElementIntegrationPoints scaling_array,  
   ScalarElementIntegrationPoints element_data_array)

The Scaled Element Signature is supported only for all element property evaluations. A corresponding entry in the convective flux boundary condition command block might be:

begin convective flux boundary condition convect  
   add surface surface_2  
   reference temperature = 180.0  
   scaled convective coefficient subroutine is aucal_htcoef fieldname = c0  
end convective flux boundary condition convect

An example of the scaled scalar element signature subroutine is as follows:

#include <utility>
#include "math.h"
#include "Aria_Calore_User_Sub_Support.h"

int aucal_htcoef(  
   UserQuery & user_query,  
   ElementIds faceid,  
   int nelemf,  
   int nint,  
   int dimension,  
   CoordinateElementIntegrationPoints coords,  
   TemperatureElementIntegrationPoints t,

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 ScalarElementIntegrationPoints scaling,  
 ScalarElementIntegrationPoints htc)  
{
 for (int j = 0; j < nint; ++j)
{
    for (int i = 0; i < nelemf; ++i)
    {
        Real x = coords(0,i,j);  
        Real y = coords(1,i,j);  
        Real r = std::sqrt(x*x+y*y);  
        Real val = 0.0;  
        Real c0 = scaling(i,j);  
        if( c0 > 0.1 ) {
            val = 45.0;  
        } else {
            val = 20.0 * r;  
        }
        htc(i,j) = val;
    }
}
return 0;
}

RegisterFunction(ScaledElemUserSubC, aucal_htcoef, "aucal_htcoef");

One should note that the number of arguments for the Scaled Element Signature and Reference Temperature Dependent Element Signature are identical. Attempts to use a user subroutine in the wrong context will be detected while parsing the input file.

33.21.4 Reference Temperature Dependent Element Signature

In some problems the surface heat transfer coefficient is determined to be a function of reference temperature rather than just the surface temperature. One example of this would be when modeling problems of natural convection where one computes a Rayleigh number based upon the difference between surface temperature and reference temperature to determine the local Nusselt number and a local heat transfer coefficient. In these cases it is sometimes more convenient to utilize an existing model for reference temperature and use those values to evaluate the heat transfer coefficient based upon additional criteria within a user subroutine. Here the interface for the Element Signature is modified to also supply interpolated values of the heat transfer coefficient at the quadrature points with the following call signature

/**
 * Prototype for the scalar reference temperature dependent
 * heat transfer coefficient signature.
 *
 * user_query: query function
 * elem_ids: pointer to the global id of the current element
 * num_elements: pointer to the number of elements in the workset
 * nint: pointer to the number of gauss points
 * coordinate_array: pointer to the coordinates of the current node (3, nelem, nint)
 * ref_temperature_array: pointer to the reference temperature (nelem, nint)
 * temperature_array: pointer to the temperature (nelem, nint)
The Reference Temperature Dependent Element Signature is currently supported only for the heat transfer coefficient. A corresponding entry in the convective flux boundary condition command block might be:

```plaintext
begin convective flux boundary condition convect
  add surface surface_2
  reference temperature = 180.0
  ref temp convective coefficient subroutine is aucal_htcoef
end convective flux boundary condition convect
```

The corresponding implementation of the reference temperature heat transfer coefficient subroutine is shown below.

```c
#include <utility>
#include "math.h"
#include "Aria_Calore_User_Sub_Support.h"

int aucal_htcoef(
  UserQuery & user_query,
  ElementIds faceid,
  int nelemf,
  int nint,
  int dimension,
  CoordinateElementIntegrationPoints coords,
  ScalarElementIntegrationPoints ref_temp,
  TemperatureElementIntegrationPoints temp,
  ScalarElementIntegrationPoints htc)
{
  Real c1 = 1.0+std::pow(std::pow((0.492/.68),(9./16.)),(8./27));
  Real c2 = 30.0e-3/2.2;
  for (int j = 0; j < nint; ++j)
  {
    for (int i = 0; i < nelemf; ++i)
    {
      Real Ra = 9.0e6*(temp(i,j)-ref_temp(i,j));
      Real Nu = std::pow((0.825+ 0.387*std::pow(Ra,(1./6))/c1),2.0);
      htc(i,j) = c2 * Nu;
    }
  }
}
```
33.21.5 Scaled Reference Temperature Dependent Element Signature

In cases where the surface heat transfer coefficient is a function of reference temperature the coefficient and one wishes to either scale the heat transfer coefficient or assign it based upon values of another variable, the heat transfer coefficient can be defined via user subroutine using yet another alternative call signature is

```c
/**
 * Prototype for the scaled scalar reference temperature dependent
 * heat transfer coefficient signature.
 *
 * user_query: query function
 * elem_ids: pointer to the global id of the current element
 * num_elements: pointer to the number of elements in the workset
 * nint: pointer to the number of gauss points
 * coordinate_array: pointer to the coordinates of the current node (3, nelem, nint)
 * ref_temperature_array: pointer to the reference temperature (nelem, nint)
 * temperature_array: pointer to the temperature (nelem, nint)
 * scale_elem_data_array: pointer to the scaling values (nelem, nint)
 * elem_data_array: pointer to the values to be calculated (nelem, nint)
 */
Int AcalScaledElemUserSubHtcC(
    UserQuery & user_query,
    ElementIds element_ids,
    Int num_elements,
    Int num_integration_points,
    Int spatial_dimension,
    CoordinateElementIntegrationPoints coordinate_array,
    ScalarElementIntegrationPoints ref_temperature_array,
    TemperatureElementIntegrationPoints temperature_array,
    ScalarElementIntegrationPoints scaling_array,
    ScalarElementIntegrationPoints flux_array)
```

The Scaled Reference Temperature Dependent Element Signature is currently supported only for the heat transfer coefficient. A corresponding entry in the convective flux boundary condition command block might be:

```plaintext
begin convective flux boundary condition convect1
  add surface surface_1
  reference temperature = 215.0
  scaled ref_temp convective coefficient subroutine = aucal_htcoef fieldname = c0
end convective flux boundary condition convect1
```

A simple example using the scaled reference temperature heat transfer coefficient signature subroutine is as follows:
```cpp
#include <utility>
#include "math.h"
#include "Aria_Calore_User_Sub_Support.h"

int aucal_htcoef(
UserQuery & user_query,
ElementIds faceid,
int nelem,
int nint,
int dimension,
CoordinateElementIntegrationPoints coords,
ScalarElementIntegrationPoints ref_temp,
TemperatureElementIntegrationPoints temp,
ScalarElementIntegrationPoints scaling,
ScalarElementIntegrationPoints htc)
{
for (int j = 0; j < nint; ++j)
{
for (int i = 0; i < nelem; ++i)
{
Real val = 0.0;
Real c0 = scaling(i,j);
Real tref = ref_temp(i,j);
if (c0 > 0.1)
{
val = 45.0*tref*c0;
} else {
val = 10.0 * tref;
}
htc(i,j) = val;
}
}
return 0;
}

RegisterFunction(ScaledElemUserSubHtcC, aucal_htcoef, "aucal_htcoef");

33.21.6 Tensor Thermal Conductivity Signature

When the thermal conductivity is anisotropic and can be easily defined in the Cartesian coordinate frame one can specify the thermal conductivity tensor with a user subroutine. The call signature for anisotropic tensor thermal conductivity user subroutines is

/**
 * Prototype for the tensor element conductivity signature.
 *
 * user_query: query function
 * elem_ids: pointer to the global id of the current element
 * num_elements: pointer to the number of elements in the workset
 * nint: pointer to the number of gauss points
 */
```
* coordinate_array: pointer to the coordinates of the current node (3, nelem, nint)
* temperature_array: pointer to the temperature (nelem, nint)
* conductivity_array: pointer to the output principal conductivities (3, 3, nelem, nint)
*/

Int AcalElemTensorUserSubC(
    UserQuery & user_query,
    ElementIds elem_ids,
    Int num_elements,
    Int nint,
    Int spatial_dimension,
    CoordinateElementIntegrationPoints coordinate_array,
    TemperatureElementIntegrationPoints temperature_array,
    TensorElementIntegrationPoints conductivity_array);

In order to use this signature to assign the tensor thermal conductivity for a two dimensional problem the subroutine is first specified in the Aria material command block. Note that a tensor conductivity requires a generalized model of heat conduction.

Begin Aria Material mat1
.
    tensor thermal conductivity = calore_user_sub name = aucal_tensor_cond type=element_tensor
    .
    heat conduction = generalized
End Aria Material mat1

A sample implementation of the corresponding user subroutine could be as shown below.

#include <utility>
#include "math.h"
#include "Aria_Calore_User_Sub_Support.h"

int aucal_tensor_cond(
    UserQuery & user_query,
    ElementIds elemid,
    Int nelem,
    Int nint,
    Int spatial_dimension,
    CoordinateElementIntegrationPoints coords,
    TemperatureElementIntegrationPoints temp,
    TensorElementIntegrationPoints cond)
{
    cond.fill(0.0);

    for (int j = 0; j < nint; ++j) {
        for (int i = 0; i < nelem; ++i) {
            cond(0,0,i,j) = 400.0;
            cond(1,1,i,j) = 300.0;
        }
    }
}
Note that one could have just as easily defined this conductivity from the Aria material command block as:

```
BEGIN ARIA MATERIAL my_mat
  tensor thermal conductivity = constant t11=400.0 t22=300.0
  heat conduction = generalized
END ARIA MATERIAL my_mat
```

### 33.21.7 Transformed Tensor Thermal Conductivity Signature

When the thermal conductivity is anisotropic only in the reference frame of local coordinates the conductivity tensor must be transformed into the Cartesian reference frame for computation. Here the user subroutine will define the thermal conductivity tensor components and the principal directions associated with those components. The call signature for anisotropic tensor thermal conductivity user subroutines is

```c
/**
 * Prototype for the tensor element principal conductivity
 * and principle direction orientation signature.
 *
 * user_query: query function
 * elem_ids: pointer to the global id of the current element
 * num_elements: pointer to the number of elements in the workset
 * nint: pointer to the number of gauss points
 * coordinate_array: pointer to the coordinates of the current node (3, nelem, nint)
 * temperature_array: pointer to the temperature (nelem, nint)
 * conductivity_array: pointer to the output principal conductivities (3, 3, nelem, nint)
 * principle_direction_array: pointer to the output principal directions (2, 3, nelem, nint)
 */

Int AcalCondUserSubC(
  UserQuery & user_query,
  ElementIds elem_ids,
  Int num_elements,
  Int nint,
  Int spatial_dimension,
  CoordinateElementIntegrationPoints coordinate_array,
  TemperatureElementIntegrationPoints temperature_array,
  ConductivityElementIntegrationPoints conductivity_array,
  DirectionElementIntegrationPoints principle_direction_array);
```

Usage of the transformed tensor conductivity subroutine is considerably more detailed than other subroutines. Here we consider an example where the conductivity is known within an element and the orientation of principle directions relative to the Cartesian frame is also known.

Anisotropic Conductivity and Orientation from Input Data:
Note that here the data block is associated with thermal conductivity via the `data = cond_values` argument
load user plugin file ./aniso.so

Begin data block cond_values
  real cond_data = ( 400.0, 200.0, 400.0, 0.0, 0.0, 0.0 )
  real cond_vec1 = ( 0.707107, 0.707107, 0.0 )
  real cond_vec2 = ( -0.707107, 0.707107, 0.0 )
End data block

Begin Aria Material my_material
  density = constant rho = 0.1
  tensor thermal conductivity = calore_user_sub name = aucal_aniso_cond \\
    type = element_direction_tensor data = cond_values
  specific heat = constant cp = 385.0
  heat conduction = generalized
End Aria Material my_material

Included below is an example user subroutine implementation of anisotropic thermal conductivity. In this implementation the vectors are indicated as being supplied from the code interface, either from input file data or from the mesh database. Alternatively, one could conditionally define the orientation vector and thermal conductivity within the user subroutine itself. This approach potentially requires selection/identification of each element by spatial location.

Note that each of the modules contains the essential features, orientation vectors, normalization and conductivity values. Once these values are supplied then the transformation operations can be performed internal to the code. This examples also demonstrates the subroutine registration process.

Int
aucal_aniso_cond(
UserQuery &
ElementIds
Int
Int
Int
CoordinateElementIntegrationPoints
TemperatureElementIntegrationPoints
ConductivityElementIntegrationPoints
DirectionElementIntegrationPoints
{
  RealArray1d cond_data(6);
  RealArray1d cond_vec2(3);
  RealArray1d cond_vec1(3);

  user_query.getUserRealInstanceData(cond_data, "cond_data");
  user_query.getUserRealInstanceData(cond_vec1, "cond_vec1");
  user_query.getUserRealInstanceData(cond_vec2, "cond_vec2");

  // normalize the vectors
  Real rlen = std::sqrt(cond_vec1(0)*cond_vec1(0)
    + cond_vec1(1)*cond_vec1(1)
    + cond_vec1(2)*cond_vec1(2));
  cond_vec1(0) = cond_vec1(0) / rlen;
  cond_vec1(1) = cond_vec1(1) / rlen;
  cond_vec1(2) = cond_vec1(2) / rlen;

  rlen = std::sqrt(cond_vec2(0)*cond_vec2(0)
    + cond_vec2(1)*cond_vec2(1)
    + cond_vec2(2)*cond_vec2(2));
  cond_vec2(0) = cond_vec2(0) / rlen;
  cond_vec2(1) = cond_vec2(1) / rlen;
  cond_vec2(2) = cond_vec2(2) / rlen;

  // check orthonormality
  Real rdot = cond_vec1(0)*cond_vec2(0) + cond_vec1(1)*cond_vec2(1)
    + cond_vec1(2)*cond_vec2(2);
  if( std::abs(rdot) > 1.0e-5 ) {
    arialog << "[ERROR] Principal directions specified "
    << " are not orthonormal" << dendl;
    return;
  }

  for( Int e = 0; e<nelem; ++e )
  {
    for( Int n=0; n<nint; ++n )
    {
      elemCond(0, 0, e, n) = cond_data(0);
      elemCond(1, 1, e, n) = cond_data(1);
      elemCond(2, 2, e, n) = cond_data(2);
      // these assignments are not necessary
      // since the elemCond data is zeroed when subroutine is called
    }
  }
}
33.21.8 Chemistry Reaction Rate Signature

The call signature for the chemistry reaction rate subroutine is

```c
/**
 * Prototype for the chemical reaction rate signature.
 * user_query: query function
 * elem_id: element id
 * reaction_rates_array: reaction rates (output)
 * ak_array: kinetics coefficients (output)
 * temperature_array: temperature at the gauss points
 * species_array: species array
 * nint: number of gauss points on this element
 * num_species: number of species on this element
 * num_reactions: number of reactions on this element
 * steric_array: steric factors for each reaction
 * prex_array: log pre-exponential factors
 * aenergy_array: activation energies
 * amusp_array: concentration exponents
 */
Int AcalChemRRUserSubC(
    UserQuery & user_query,
    Int elem_id,
    ReactionIntegrationPoints reaction_rates_array,
    ReactionIntegrationPoints ak_array,
    ConstIntegrationPoints temperature_array,
    ConstSpeciesIntegrationPoints species_array,
    Int nint,
)
33.21.9 Chemistry Auxiliary Variables Signature

The other chemistry subroutine type is the one that is used to manipulate auxiliary variables. The call signature for this subroutine is as follows

```c
/**
 * This is the prototype for the chemical aux variable signature.
 *
 * user_query: query function
 * elem_id: element id
 * temperature_array: temperature at the gauss points
 * species_array: species array
 * aux_variables_array: auxiliary variable array (output)
 * nint: number of gauss points on this element
 * num_species: number of species on this element
 * num_reactions: number of reactions on this element
 * num_aux_variables: number of auxiliary variables on this element
 */
Int AcalChemAVUserSubC(
    UserQuery & user_query,
    Int elem_id,
    ConstIntegrationPoints temperature_array,
    ConstSpeciesIntegrationPoints species_array,
    AuxVariablesIntegrationPoints aux_variables_array,
    Int nint,
    Int num_species,
    Int num_reactions,
    Int num_aux_variables);
```

We note that each of the chemistry subroutines operate on a single element at a time.

33.22 FORTRAN-like C Interface Functions

A number of utility functions are supplied for use in the FORTRAN-like C functions. These utilities provide access to internal variables that one might wish to use in the C functions and enables manipulation of these Real and Integer variables to a certain extent. Examples of interface functions used to modify values would be for assignment of boundary condition, material data or global variables. Additionally, the interface functions provide access to values associated with data blocks. Details for usage of data block variables with user functions is briefly detailed in the paragraphs which follow.

Within the Aria code data blocks are generically known as Resources and the usage of specific Resources within the code are referred to as Region or Instance. Thus the appearance of Resources, Region or Instance in either the name or call signature of the interface function implies usage of a data block.
If an interface function is referenced within the input file with a `USE DATA BLOCK data_block_name` then that data block is in some sense bound to that scope and the data block variables become accessible without explicit reference to the data block name. When the `USE DATA BLOCK data_block_name` specification appears in the Region scope a user subroutine function can access the `data_block_name` variables using `Region` type call signatures. Likewise if the if the `USE DATA BLOCK data data_block_name` specification appears within a material property, boundary condition or volume heat command block then user subroutine functions defined within that command block can access the `data_block_name` variables using `Instance` type call signatures.

A more general usage of data block variables within user subroutine functions requires explicit reference to the data block, a `Resource` argument, within the user interface function call signature. This type of interface call proves useful when one wishes to access data from more than one data block or when one wants to ensure which data block will be used.

When more than one data block will be used within a command block an alternative would be to provide the data block as a `data = data_block_name` argument in the feature specification. For example: `Heat Transfer Coefficient = Calore_User_Sub name=hcoeff type=element data= data_block_name`. Of course, one could always employ this approach exclusively when dealing with data block variables in lieu of the `use data block data data_block_name` model usage.

Some of the remaining user interface function calls will reference specific program variables such as time or spatial dimension which cannot be altered (i.e. they will return non-mutable, `const`, program data). In most cases the interface functions will either retrieve values for use in the subroutine or populate values that are communicated back to the code. Here any variables not supplied through the subroutine interface must be explicitly declared within the user subroutine.

User interface function calls are part of the application code that are accessible to the user subroutine only through the `user_query` facility. As an example, in order to reference the the `currentTime()` function the subroutine body would contain a call as indicated here.

```c
int your_subroutine_interface( UserQuery & user_query,
                             more subroutine_arguments, . , . , . , .)
{
  int ierror = 0;
  .
  Real time = user_query.currentTime();
  .
  statements referencing time
  .
  return ierror;
}
```

Call signatures to the various user interface functions are provided below.

```c
Real currentTime() const;

Real currentTimeStepSize() const;

void getFunctionValues( Int, Real *, Real *, const String & name);

void getUserRegionData(RealArray1d &real_array, const String & real_label,
                        IntArray1d &int_array, const String & int_label);
```

937
void putUserRegionData(RealArray1d &real_array, const String & real_label,
        IntArray1d &int_array, const String & int_label);

void getUserRealRegionData(RealArray1d &real_array, const String & real_label);

void putUserRealRegionData(RealArray1d &real_array, const String & real_label);

void getUserIntRegionData(IntArray1d &int_array, const String & int_label);

void putUserIntRegionData(IntArray1d &int_array, const String & int_label);

void getUserInstanceData(RealArray1d &real_array, const String & real_label,
        IntArray1d &int_array, const String & int_label);

void putUserInstanceData(RealArray1d &real_array, const String & real_label,
        IntArray1d &int_array, const String & int_label);

void getUserRealInstanceData(RealArray1d &real_array, const String & real_label);

void putUserRealInstanceData(RealArray1d &real_array, const String & real_label);

void getUserIntInstanceData(IntArray1d &int_array, const String & int_label);

void putUserIntInstanceData(IntArray1d &int_array, const String & int_label);

void getUserMaterialData(RealArray1d &real_array, const String & real_label,
        IntArray1d &int_array, const String & int_label);

void putUserMaterialData(RealArray1d &real_array, const String & real_label,
        IntArray1d &int_array, const String & int_label);

void getUserRealMaterialData(RealArray1d &real_array, const String & real_label);

void putUserRealMaterialData(RealArray1d &real_array, const String & real_label);

void getUserIntMaterialData(IntArray1d &int_array, const String & int_label);

void putUserIntMaterialData(IntArray1d &int_array, const String & int_label);

void getUserRealMaterialData(Real &real_value, const String & label);

void putUserRealMaterialData(Real &real_value, const String & label);

void getIntFaceVar(Int id, const String & name, Int * s);

void getRealFaceVar(Int id, const String & name, Real * s);

void getRealElemVar(Int id, const String & name, Real * s);

void getIntNodeVar(Int id, const String & name, Real * s);

void getRealNodeVar(Int id, const String & name, Real * s);

void getIntFaceVar(Int id, const String & name, Int * s);
void getIntElemVar(Int id, const String & name, Int * s);
void getIntNodeVar(Int id, const String & name, Int * s);
void putRealFaceVar(Int id, const String & name, Real * s);
void putRealElemVar(Int id, const String & name, Real * s);
void putRealNodeVar(Int id, const String & name, Real * s);
void putIntFaceVar(Int id, const String & name, Int * s);
void putIntElemVar(Int id, const String & name, Int * s);
void putIntNodeVar(Int id, const String & name, Int * s);

Int currentTimeStepIndex();
Int getSpatialDimension();
void materialName(String & meshpart_name);
String getPartName(){ return m_block_name; }
void getContactMaterialName(Int, String &);
void getContactElement(const Int *, Int *, Int *);
void nodePredT(Int id, Real *);
void nodePrevT(Int id, Real *);
void TDotElementGauss(Int id, Real *tDot);
void getAuxVar(Int id, Real * a);
void setAuxVar(AuxVariablesIntegrationPoints aux);
Int getNumAuxVars();
void getSpecies(Int id, Real * s);
bool getElementVolume(Int id, Real * value);
bool getElementMass (Int id, Real * value);
bool getFieldIndex(const Real * fieldPtr, Int * index, const String & name);
Real or Int getEncoreGlobalVariable(const String & name)
void getEncoreGlobalVariable(Real or Int value, const String & name)
Real or Int getGlobalVariable(const String & name)
void getGlobalVariable(Real or Int value, const String &name)

void locallyUpdateUserGlobalVariable(Real or Int value, const String &name)

void getUserRealData(RealArray1d &real_array, const String &real_label, const String &resource, const String &user_sub_name);

void putUserRealData(RealArray1d &real_array, const String &real_label, const String &resource, const String &origin, const String &user_sub_name);

void getUserRealData(Real &real_value, const String &real_label, const String &resource, const String &user_sub_name);

void putUserRealData(Real &real_value, const String &real_label, const String &resource, const String &origin, const String &user_sub_name);

void getUserIntData(Int &int_value, const String &int_label, const String &resource, const String &user_sub_name);

void putUserIntData(Int &int_value, const String &int_label, const String &resource, const String &origin, const String &user_sub_name);

void getUserIntData(IntArray1d &int_array, const String &int_label, const String &resource, const String &user_sub_name);

void putUserIntData(IntArray1d &int_array, const String &int_label, const String &resource, const String &origin, const String &user_sub_name);

void push_query(UserQuery *query)

void pop_query(UserQuery *query)

33.22.1 CHEMEQ Species Variable Access

Occasionally one may wish to access the values of species from the CHEMEQ system within the FORTRAN-like C subroutine using the getSpecies(Int elem_id, Real species *) interface. In this case some knowledge of the returned data layout is required in order to access the correct values. Internal to the code species values for an element are defined at each of the element quadrature points and stored collectively in a single flat array. Thus in order to access species values for a specific component one must stride into the flat array of values.

As an example, consider the case of three species, num_species = 3, chemistry on a 2D quadrilateral which has four quadrature points. Within the subroutine storage must be provided for an array of length 12 denoted here as s,

RealArray1D s(12);
This implies the species data layout shown below

\[ s(0) \ s(1) \ s(2) \ s(3) \ s(4) \ s(5) \ s(6) \ s(7) \ s(8) \ s(9) \ s(10) \ s(11) \]

We note that if one is interested in the values for the second species these values, \( s(1), s(4), s(7), s(10) \), are offset by 1 into each subset of quadrature point values. For a given element id, \( \text{eid} \), the species values for that element are obtained with the user_query \( \text{getSpecies( \text{eid}, \&s )} \). Then to access each of the \( i \)th quadrature point values \((i=0,1,2,3)\) for the desired species one must stride into an appropriate offset position, \( \text{index} = 1 \), so that the species quadrature point value is

\[ \text{value} = s(\text{num\_species} \times i + \text{index}) \]

### 3.33.23 Debug Output From User Subroutines

Oftentimes one finds it instructive to output diagnostic information to file from the user subroutine. Aria provides three different ways of generating this output, output directly to a user specified file, output to the log file for all time and selective output to the log file. Each of these options has both advantages and disadvantages. In each of the cases information could be written each time the subroutine is called so the user must provide some code logic to limit the amount of output.

Outputting directly to a user specified file is most straightforward approach but then program output will be split over multiple files. Output to the log file is convenient but in this case the subroutine output will be interlaced with log file output that one would normally expect. Outputting to the log file for all time is appropriate only when diagnostics are always needed. Selective output to the log file is preferred in most cases since the amount of output is restricted to special instances. As an example of each approach the subroutine appearing in a previous section is augmented with the appropriate code. Here direct output to file is demonstrated with both C file specification and C++ file specification.

```c
Int
aucal_aniso_cond(
    UserQuery & user_query,
    ElementIds objid,
    Int nelem,
    Int nint,
    Int spatial_dimension,
    CoordinateElementIntegrationPoints coords,
    TemperatureElementIntegrationPoints temperature,
    ConductivityElementIntegrationPoints elemCond,
    DirectionElementIntegrationPoints elemVec)
{
    // main ingredients for separate file output

    static FILE *file_pointer = NULL;
    if(NULL == file_pointer ) // only open the file once
    {
        file_pointer = fopen("c.out", "w");
        ThrowRequire(file_pointer);
    }
```
// C++ file specification
static bool have_ofs = false;
static std::ofstream ofs;

if( !have_ofs )
{
    ofs.open("cpp.out", std::ios::out);
    have_ofs = true;
}

RealArray1d cond_data(6);
RealArray1d cond_vec2(3);
RealArray1d cond_vec1(3);

user_query.getUserRealInstanceData(cond_data, "cond_data");
user_query.getUserRealInstanceData(cond_vec1, "cond_vec1");
user_query.getUserRealInstanceData(cond_vec2, "cond_vec2");

// normalize the vectors
Real rlen = std::sqrt(cond_vec1(0)*cond_vec1(0)
    + cond_vec1(1)*cond_vec1(1)
    + cond_vec1(2)*cond_vec1(2));

cond_vec1(0) = cond_vec1(0) / rlen;
cond_vec1(1) = cond_vec1(1) / rlen;
cond_vec1(2) = cond_vec1(2) / rlen;

rlen = std::sqrt(cond_vec2(0)*cond_vec2(0)
    + cond_vec2(1)*cond_vec2(1)
    + cond_vec2(2)*cond_vec2(2));

cond_vec2(0) = cond_vec2(0) / rlen;
cond_vec2(1) = cond_vec2(1) / rlen;
cond_vec2(2) = cond_vec2(2) / rlen;

// check orthonormality
Real rdot = cond_vec1(0)*cond_vec2(0) + cond_vec1(1)*cond_vec2(1)
      + cond_vec1(2)*cond_vec2(2);

// orthonormality diagnostics
if( std::abs(rdot) > 1.0e-5 )
{
    // Information is always output to the c.out file C style
    fprintf(file_pointer, "[ERROR] Principal directions specified are not orthonormal.\n");
    // Information is always output to the cpp.out file C++ style
    ofs << "[ERROR] Principal directions specified "
        << " are not orthonormal" << std::endl;
    // Information is always output to log file
    arialog << "[ERROR] Principal directions specified "
            << " are not orthonormal" << endl;
    // Information is conditionally output to log file by supplying the
    // -O "-arialog user-sub" arguments to sierra command
    debuglog << "[ERROR] Principal directions specified "
             << " are not orthonormal" << endl;
    return;
}
for( Int e = 0; e<nelem; ++e )
{
    for( Int n=0; n<nint; ++n )
    {
        elemCond(0, 0, e, n) = cond_data(0);
        elemCond(1, 1, e, n) = cond_data(1);
        elemCond(2, 2, e, n) = cond_data(2);
        // these assignments are not necessary
        // since the elemCond data is zeroed when subroutine is called
        elemCond(0, 1, e, n) = cond_data(3);
        elemCond(1, 0, e, n) = cond_data(3);
        elemCond(0, 2, e, n) = cond_data(4);
        elemCond(2, 0, e, n) = cond_data(4);
        elemCond(1, 2, e, n) = cond_data(5);
        elemCond(2, 1, e, n) = cond_data(5);
        elemVec(0, 0, e, n) = cond_vec1(0);
        elemVec(0, 1, e, n) = cond_vec1(1);
        elemVec(0, 2, e, n) = cond_vec1(2);
        elemVec(1, 0, e, n) = cond_vec2(0);
        elemVec(1, 1, e, n) = cond_vec2(1);
        elemVec(1, 2, e, n) = cond_vec2(2);
    }
}

return;

RegisterFunction(ElemTensorUserSubC, aucal_aniso_cond, "aucal_aniso_cond");

In some instances one might wish to terminate execution when certain conditions arise in the user subroutine. Here the user can provide a diagnostic for termination when causing the program to stop including a conditional such as the one shown below.

if( condvec(2) == 0 )
    throw sierra::RuntimeError() << "condvec(2) = 0 on " << objid;

33.24 FORTRAN User Subroutines

An Aria FORTRAN user subroutine must be a text file of suffix .F or .f. The FORTRAN file content is made available to the application by either compiling the module and statically linking it to the application archive or by dynamically linking the module to the executable at run time. The static link approach is often preferred on capability high-performance computing platforms, where dynamically linked subroutines are either not supported or suffer a penalty in performance. User selection of static or dynamic linking is determined by a combination of commands used for job execution and input command lines.

For the static link approach consider the use of a FORTRAN subroutine module file named Aucal_ConvFlux.F. In order to use this module the input file must contain an input command line such as
user subroutine file = Aucal_ConvFlux.F

A static link of the module is then initiated by use the job command

```bash
sierra -make aria -i input.i
```

This command will invoke creation of an auxiliary FORTRAN module used to perform a registration of the subroutine in the executable. Then the subroutine and auxiliary module are compiled followed by a static link with the existing archive to create an executable written to a `UserSubsProject/bin` subdirectory. This executable must then be specifically called out in the job execution command.

For the dynamic link approach the user must supply the auxiliary FORTRAN module used for subroutine registration. Most often the auxiliary module is included in the subroutine file but can also be provided as a standalone file. In order to use this module the input file must contain an input command line such as

```bash
load user plugin file Aucal_ConvFlux.so using function conv_flux_register
```

where `conv_flux_register` is the name of the auxiliary module used for subroutine registration. An example of the auxiliary module is given in the section which follows 33.25. Note that this registration is equivalent to that of Aria User Plugin function registration 18. Invocation of the job command

```bash
sierra -make aria -i input.i
```

initiates a compilation that leads to creation of a shared object, `Aucal_ConvFlux.so`. During execution of the job the shared object will be automatically loaded and used in the execution.

Note that, since `Aucal_ConvFlux.F` is a filename, its input file name is case-sensitive, and considerations must be made concerning its location path. The user is free to name the file and subroutines in any way chosen, provided it is consistent with compiler and operating system limitations. For example, special characters such as '*' and ',' should be avoided. Finally actual use of the function by the application is triggered by association of the the user subroutine name in a particular command line, presently only Dirichlet boundary conditions, flux boundary conditions and material property evaluations.

### 33.25 FORTRAN Interfaces

While FORTRAN user subroutines could be defined to supply many different parameters, only two different signatures are currently provided in Aria. These signatures characterize the argument lists by which the subroutine interacts with the code. The various subroutine signatures are denoted as:

- node signature 33.25.1
- element signature 33.25.2
- convective flux signature 33.25.3
- radiative flux signature 33.25.4

Whereas the first C-Style subroutines focus on providing one data set or material property, some of the FORTRAN subroutines must be implemented as a pair or as a triplet of properties. The different subroutine call signatures are outlined below. With the exception of the `node`, these signatures will be used for subroutines that operate on quadrature point values. The FORTRAN user subroutines must be registered with a function signature similar to the example below given below.

c

944
c fortran subroutine registration

subroutine conv_flux_register

implicit none
external aucal_fort_elem_sub
external ht_coeff_test
external t_ref_test

c
    call fmwkusersub(aucal_fort_elem_sub, ht_coeff_test, 
                        'ht_coeff_test')
call fmwkusersub(aucal_fort_elem_sub, t_ref_test, 
                        't_ref_test')
c
end

Here the first argument to fmwkusersub is the internally defined interface signature name. The second argument is the name of the FORTRAN function (subroutine) and the third is the name to be associated with the subroutine from within the input file.

Perhaps the biggest difference between C-Style subroutines and FORTRAN subroutines will be the difference in how any problem specific input file data is provided to the user subroutine. While C-style subroutines provide flexibility to provide parameters through arbitrary DATA_BLOCKs, the FORTRAN subroutine parameters can only be provided through REAL DATA or INT DATA command lines within the command block making reference to the FORTRAN subroutine.

### 33.25.1 FORTRAN Node Subroutine

The FORTRAN Node subroutine is used primarily for applying Dirichlet boundary conditions and is designed to operate on sidesets. For example the subroutine call could be invoked using the following input command block:

```
Begin Temperature Boundary Condition bcblock1
    Temperature fortran subroutine = dirich_test
    Add surface surface_2
End Temperature Boundary Condition bcblock1
```

Here we demonstrate the node user subroutine in which the user supplied subroutine nodal values are assigned as constants. Note that time and coordinates are being supplied to the subroutine so the assigned values could also be functions of the supplied variables.

```c
Subroutine to handle coordinate and time-dependent
   Temperature Dirichlet BC

subroutine dirich_test(objid,nnodes,coords,temperature,
                        time,rdat,idat,ierror)
implicit none
integer nnodes
integer objid
double precision coords(2,nnodes)
```
double precision temperature(nnodes)
double precision time
integer ierror

c Note: rdat and idat must be sized at least 1,
even when not used
otherwise they must be sized consistent with the
data being provided from the input file

double precision rdat(1)
integer idat(1)

c INPUT ARGUMENTS:
objc(nelem): array of mesh object ID’s. Useful for passing through
to query methods
nelem: number of elements in the workset
nint: Number of integration points per element in the workset
coords(2,n,nint): array of the coordinates (x,y,z) of the Gauss points
rdat() real parameter data
idat() integer data

c OUTPUT ARGUMENTS:
temperature: temperature at the node
eerror: user defined error code for calore to test.
 zero means success.

c integer i, j

eerror = 0

do i=1,nnodes

    if( coords(2,i) .lt. 0.5 ) then
        temperature(i) = 1.5
    else
        temperature(i) = 0.1
    end if

end do

return
end

33.25.2 FORTRAN Element Subroutine

The FORTRAN element subroutine can be used to compute coefficients or material properties associated with either surface or volume elements. While the example given here it is associated with surface flux conditions, it is important to note that the element subroutine can be more broadly applied.

The FORTRAN element subroutine is used to applying a flux boundary condition and is here defined to operate on sidesets. For example the subroutine call could be invoked using the following input command block:
Here we demonstrate the flux user subroutine in which the user supplied subroutine heat flux values are assigned as constants. Note that temperature, time and coordinates are being supplied to the subroutine so the assigned values could also be functions of the supplied variables.

```c
Subroutine to handle coordinate, time and temperature
dependent flux

subroutine flux_test(objid, nelem, nint, coords, temperature, &
                      flux, time, rdat, idat, ierror)
  implicit none
  integer nelem, nint
  integer objid(nelem)
  double precision coords(3, nelem, nint)
  double precision temperature(nelem, nint)
  double precision flux(nelem, nint)
  double precision time
  integer ierror

double precision rdat(1)
integer idat(1)

INPUT ARGUMENTS:
  objid(nelem): array of mesh object ID’s. Useful for passing through
to query methods
  nelem: number of elements in the workset
  nint: Number of integration points per element in the workset
  coords(3,n,nint): array of the coordinates (x,y,z) of the Gauss points
  temperature(n,nint): temperature at the Gauss points
  rdat() real parameter data
  idat() integer data

OUTPUT ARGUMENTS:
  flux: array of length ( n, nint) containing
         the heat flux coefficient
         at integration points
  ierror: user defined error code for calore to test.
  zero means success.

integer i, j

ierror = 0
```
do i=1,nelem
    do j=1,nint
        if( coords(3,i,j) .gt. 4.0 ) then
            flux(i,j) = 4000.0
        else
            flux(i,j) = 0.0
        end if
    end do
end do
return
end

33.25.3 FORTRAN Convective Flux Subroutine

The FORTRAN Convective Flux subroutine is used to applying a flux boundary condition and is designed to operate on sidesets. For example the subroutine call could be invoked using the following input command block:

Begin Convective Flux Boundary Condition C1
    Add Surface surface_1
    convective coefficient fortran subroutine is ht_coeff_test
    reference temperature fortran subroutine is t_ref_test
    integer data 8, 12, 14
    real data 1.0 2.0
end

Here we demonstrate the convective flux user subroutine in which the user supplied subroutine values for heat transfer coefficient and reference temperature are assigned as constants. Note that temperature, time and coordinates are being supplied to the subroutine so the assigned values could also be functions of the supplied variables. Here the real and integer data prescribed in the input command block are supplied to the user subroutine but are not used.

c
    Subroutine to assign heat transfer coefficient

c
    subroutine ht_coeff_test(objid,nelem,nint,coords,temperature,
                            &
                            htccoeff,time,rdat,idat,ierror)
    implicit none
    integer nelem, nint
    integer objid(nelem)
    double precision coords(3,nelem,nint)
    double precision temperature(nelem,nint)
    double precision htccoeff(nelem,nint)
    double precision time
    integer ierror

    double precision rdat(2)
integer idat(3)

INPUT ARGUMENTS:
objid(nelem): array of mesh object ID’s. Useful for passing through
to query methods
nelem: number of elements in the workset
nint: Number of integration points per element in the workset
coords(3,n,nint): array of the coordinates (x,y,z) of the Gauss points
temperature(n,nint): temperature at the Gauss points
rdat() real parameter data
idat() integer data

OUTPUT ARGUMENTS:
htcoef: array of length ( n, nint) containing
the heat transfer coefficient
at integration points
ierror: user defined error code for calore to test.
zero means success.

integer i, j

do i=1,nelem

do j=1,nint

    htc[100,200] = 50.0

end do
end do

return
end

Subroutine to assign reference temperature

subroutine t_ref_test(objid,nelem,nint,coords,temperature,
& tref,time,rdat,idat,ierror)
imPLICIT none
integer nelem, nint
integer objid(nelem)
double precision coords(3,nelem,nint)
double precision temperature(nelem,nint)
double precision tref(nelem,nint)
double precision time
integer ierror

double precision rdat(2)
integer idat(3)

INPUT ARGUMENTS:
objid(nelem): array of mesh object ID’s. Useful for passing through
to query methods
nelem: number of elements in the workset
33.25.4 FORTRAN Radiative Flux Subroutine

The FORTRAN Radiative Flux subroutine is used to applying a flux boundary condition and is designed to operate on sidesets. For example the subroutine call could be invoked using the following input command block:

Begin Radiative Flux Boundary Condition fortran
  Add Surface surface_1
  emissivity fortran subroutine is emissivity_test
  form factor fortran subroutine is form_factor_test
  reference temperature fortran subroutine is t_ref_test
  integer data 8, 12
  real data 1.0 2.0
end

Here we demonstrate the radiative flux user subroutine in which the user supplied subroutine values for form factor, emissivity and reference temperature are assigned as constants. Note that temperature, time and coordinates are being supplied to the subroutine so the assigned values could also be functions of the supplied variables. Here the real and integer data prescribed in the input command block are supplied to the user subroutine but are not used.

Subroutine to assign form factor

subroutine form_factor_test(objid,nelem,nint,coords,temperature, &
  ffactor,time,rdat,idat,ierror)
implicit none

integer nelem, nint
integer objid(nelem)
double precision coords(3,nelem,nint)
double precision temperature(nelem,nint)
double precision ffactor(nelem,nint)
double precision time
integer ierror

double precision rdat(2)
integer idat(3)

INPUT ARGUMENTS:
objcid(nelem): array of mesh object ID’s. Useful for passing through
    to query methods
nelem: number of elements in the workset
nint: Number of integration points per element in the workset
coords(3,n,nint): array of the coordinates (x,y,z) of the Gauss points
temperature(n,nint): temperature at the Gauss points
rdat(): real parameter data
idat(): integer data

OUTPUT ARGUMENTS:
ffactor: array of length ( n, nint) containing
    the form factor at integration points
ierror: user defined error code for aria to test.
    zero means success.

integer i, j

do i=1,nelem
  do j=1,nint
    form_factor(i,j) = 0.92
  end do
end do

end do

return
end

Subroutine to assign emissivity

subroutine emissivity_test(objid,nelem,nint,coords,temperature,
 &     emis,time,rdat,idat,ierror)
  implicit none
  integer nelem, nint
  integer objid(nelem)
double precision coords(3,nelem,nint)
double precision temperature(nelem,nint)
double precision emis(nelem,nint)
Table 1: Mean and Standard Deviation of Temperature for Different Coarse Grids and Aliasing Conditions

<table>
<thead>
<tr>
<th>Coarse Grid</th>
<th>Aliasing Condition</th>
<th>Mean Temperature</th>
<th>Standard Deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grid 1</td>
<td>None</td>
<td>20.5°C</td>
<td>1.2°C</td>
</tr>
<tr>
<td></td>
<td>Aliasing</td>
<td>20.0°C</td>
<td>1.0°C</td>
</tr>
</tbody>
</table>

**Notes:**
- Mean temperature and standard deviation are calculated from 1000 samples.
- Aliasing reduces the mean temperature by 0.5°C and the standard deviation by 0.2°C.

**Further Analysis:**
- The results indicate that aliasing significantly affects the temperature measurements.
- Additional experiments are needed to validate these findings.

**References:**
INPUT ARGUMENTS:

- objid(nelem): array of mesh object ID’s. Useful for passing through to query methods
- nelem: number of elements in the workset
- nint: Number of integration points per element in the workset
- coords(3,n,nint): array of the coordinates (x,y,z) of the Gauss points
- temperature(nelem,nint): temperature at the Gauss points
- time
- rdat() real parameter data
- idat() integer data

OUTPUT ARGUMENTS:

- t_ref: array of length (nelem, nint) containing the heat transfer coefficient at integration points
- ierror: user defined error code for aria to test. zero means success.

```fortran
integer i, j

do i=1,nelem
    do j=1,nint
        tref(i,j) = 300.0
    end do
end do
return
end
```

### 33.26 FORTRAN Subroutine Interface Functions

A number of utility functions are supplied for use in the FORTRAN subroutines. These utilities provide access to internal variables that one might wish to use within the subroutine and enables manipulation of these Double Precision and Integer variables to a certain extent. Examples of interface functions used to modify values associated with boundary condition, material data or global variables.

User interface function calls are part of the application code that are accessible from the user subroutine. As an example, in order to retrieve a global variable named \( T_{AVERAGE} \) the subroutine body would contain a an external definition of the retrieval function and the function call as indicated here.

```fortran
subroutine your_subroutine_interface( objid,nelem,nint,coords,temperature, &
    source,time,rdat,idat,ierror)
    implicit none
    integer nelem, nint
    integer objid(nelem)
    double precision coords(3,nelem,nint)
    double precision temperature(nelem,nint)
    double precision source(nelem,nint)
    double precision time
    integer ierror
```
double precision t_avg
character*9 cvar

external aria_get_global_real_var

cvar = "T_AVERAGE"
ierror = 0
.
call aria_get_global_real_var(t_avg,1,cvar)
.
C statements referencing t_avg
 .
return
end
}

Call signatures to the various FORTRAN user interface functions are provided below.

aria_get_time(double precision time)

aria_get_timestep_index(integer k)

aria_get_material_name(integer length, character name)

aria_get_contact_material_name(integer face_id, integer length, character name)

aria_get_contact_element(integer face_id, integer elem_id, integerfound)

aria_get_pred_nodal_t(integer id, double precision t, integer found)

aria_get_prev_nodal_t(integer id, double precision t, integer found)

aria_get_species(integer id, double precision s)

aria_get_aux_var(integer id, double precision a)

aria_get_element_tdot(integer id, double precision tdot)

aria_get_global_real_var(double precision value, integer len, character varName)

aria_get_global_int_var(integer value, integer len, character varName)

aria_lupdate_global_real_var( double precision deltaValue, integer len, character varName)

aria_lupdate_global_int_var( integer deltaValue, integer len, character varName)

aria_get_element_volume(integer id, double precision value)

aria_get_element_mass(integer id, double precision value)

aria_get_real_nodal_var(double precision value, integer id, integer nameLen, character varName)

aria_get_int_nodal_var(integer value, integer id, integer nameLen, character varName)
aria_put_real_nodal_var(double precision value, integer id, integer nameLen, character varName)
aria_put_int_nodal_var(integer value, integer id, integer nameLen, character varName)
aria_get_real_elem_var(double precision value, integer id, integer nameLen, character varName)
aria_get_int_elem_var(integer value, integer id, integer nameLen, character varName)
aria_put_real_elem_var(double precision value, integer id, integer nameLen, character varName)
aria_put_int_elem_var(integer value, integer id, integer nameLen, character varName)
aria_get_real_face_var(double precision value, integer id, integer nameLen, character varName)
aria_get_int_face_var(integer value, integer id, integer nameLen, character varName)
aria_put_real_face_var(double precision value, integer id, integer nameLen, character varName)
aria_put_int_face_var(integer value, integer id, integer nameLen, character varName)
aria_get_material_real_data(double precision addr, integer length, character name, integer name_length)
aria_put_material_real_data(double precision addr, integer length, character name, integer name_length)
aria_get_material_int_data(integer addr, integer length, character name, integer name_length)
aria_put_material_int_data(integer addr, integer length, character name, integer name_length)
aria_get_region_real_data(double precision addr, integer length, character name, integer name_length)
aria_put_region_real_data(double precision addr, integer length, character name, integer name_length)
aria_get_region_int_data(integer addr, integer length, character name, integer name_length)
aria_put_region_int_data(integer addr, integer length, character name, integer name_length)
aria_get_instance_real_data(double precision addr, integer length, character name, integer name_length)
aria_put_instance_real_data(double precision addr, integer length, character name, integer name_length)
aria_get_instance_int_data(integer addr, integer length, character name, integer name_length)
aria_put_instance_int_data(integer addr, integer length, character name, integer name_length)

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33.27 Solution Mapping Methodology

Numerical simulations of full system models are oftentimes intensely consuming calculations. During the design process the need often arises to study the same problem with minor changes to a subcomponent of the original model. To fulfill this need a solution mapping methodology that simulates the thermal response of a subassembly, without having to repeat a simulation of the full assembly model has been developed. This capability is particularly useful in cases where the analyst must evaluate alternative subassembly designs and provide timely input to engineering design studies. This section describes how to apply this solution mapping methodology.

The basic idea of the Aria solution mapping methodology is to use existing simulation results (e.g. temperature or heat flux) from a full assembly model to set boundary conditions (BCs) for a separate simulation of a smaller, but possibly more geometrically resolved, subassembly or component. In effect, the Aria thermal simulation of the full assembly serves as the “Donor” of the solution results, while the subassembly model becomes the “Recipient”. This methodology uses existing Aria code features to map solution results to the appropriate boundary surfaces of the subassembly model. It is important to mention that the methodology permits the analyst to use different meshes and mesh resolutions at the interface between the full and subassembly models.

33.27.1 Setting up Simulation for Donor Model

To set up a Donor model, the user modifies the Aria input file for the full assembly model so that it outputs the solution information that will be transferred or mapped to the outer surfaces of the Recipient subassembly model or mesh. The user can choose to set either Dirichlet boundary conditions (i.e. map temperature) or Neumann type boundary conditions (i.e. map heat fluxes) on the outer surface of the recipient model. Numerical studies conducted by the Aria development team indicate that temperature mapping generally provides much more reliable estimation of the subassembly temperatures than using heat flux mapping. In fact, the user is cautioned to avoid the flux mapping transfer capability, especially for long-duration transients when the error accumulates in time and can become quite substantial.

In either type of solution mapping, the user must compile and specify the material blocks surrounding and contacting the subassembly, which are to be used in the solution mapping. It is recommended that, where possible, the user should map the solution from the Donor model to an enclosing region that extends slightly beyond the subassembly, rather than mapping solutions directly on the subassembly exterior. This approach provides some measure of buffering of the effect of sudden changes in time varying BCs.

First, add a new “Begin Results Output” (described in 25.1) block to the input file used specifically for mapping, such as the example shown below. This output block is in addition to the typical output block that writes the nodal field for the entire model domain.
Begin Results Output Solution_Mapping
   Database Name = Donor_solution.e
   Include = BlockList

   # ... Use for Temperature Mapping Only ...
   Nodal Variables = solution->Temperature as T

   # ... Use for Heat Flux Mapping Only ...
   Nodal variables = pp->Heat_Conduction as Heat_Flux
   At Time 0.0, Increment = 1.0
   Timestep Adjustment Interval is 4
End

In the above syntax example, the BlockList is the list of block IDs that will be used in the solution mapping. If heat flux mapping is used, then add the following command to the “Region” level of the input file:

Postprocess Heat_Conduction on all_blocks

With above command lines added, execute the thermal simulation for the full assembly model and create the donor-solution exodus output file.

33.27.2 Setting up Simulation for Recipient Model

In most cases, the user can prepare the Aria input file for the Recipient model by copying and simplifying the input file from the original full assembly model. The input file for Recipient model will reference the subassembly mesh and the subassembly material definitions. With the input file stylized for the subassembly, the user should add:

1. A “Begin Finite Element Model” block
2. A “Begin Input_Output Region” block
3. Transfer Commands
4. Transfer definitions via the “Begin Transfer” block
5. BC XFER command (for temperature mapping only).

For the case where heat flux is used in the solution mapping, the last step is replaced with:

1. Addition of the user field command (see Chapter 9 and
2. Addition of a “Flux Vector Node Variable” (see Chapter 33.1)

1. Finite Element Model

The purpose of the new Finite Element Block is to identify the exodus file generated by the Donor model. The syntax of this command block is described in Chapter 2. An example of this block is shown below:
Begin Finite Element Model Input_Transfer
  Database Name = Donor_solution.e
  Database Type = exodusII
End

This block is placed in the Domain level of the input file. Explanation of Domain, Procedure and Region levels of the input file can be found in Chapter 2 of the manual.

2. Begin Input_Output Block

A description of the “Begin Input_Output Region” block can be found in Section 15.1. This command block is placed in the Procedure level of the input file. An example is given below:

Begin Input_Output Region Transfer_Region
  Use Finite Element Model Input_Transfer
End

3. Transfer Commands

A detailed explanation of Transfer command syntax and usage is presented in Section 15.1 of this user manual. These commands are added to the “Begin System Main” block. An example of these transfer commands is shown below:

Begin Solution Control Description
  Use System Main
  Begin System Main
   Simulation Start Time = 0.0
   Simulation Termination Time = 3600.0
   Begin Transient solution_block_1
    Advance transfer_region
    Transfer trans_to_aria
    Advance myRegion
   End
   .
   .
  End System Main
  .
  .
End Solution Control Description

The above command block is located in the Procedure level of the input file.

4. Transfer Definitions

Syntax and usage of transfer definitions is presented in Section 15.1 of the manual. Here is an example of the transfer definitions for

Begin Transfer trans_to_aria
Interpolate Volume Nodes from transfer_region to myRegion
Nodes Outside Region = project #or extrapolate, #or truncate
# ... Use for Temperature transfer only ...
Send Block block_1 block_2 ... to surface_1 surface_2 ...
Send Field T State None to solution->Temperature State New
# ... Use for heat flux transfer only ...
# Send Block block_1 block_2 ... to block_1 block_2 ...
# Send Field heat_flux State None to surf_flux State None
# ... Tolerance settings ...
Search Geometric Tolerance is 0.001
Search Surface Gap Tolerance is 0.01
End

The above example has transfer definitions for both temperature and heat flux, however, only one set of commands can be used.

5a. BC XFER Command

As explained previously, the BC XFER command is used to set the temperature BC on the Recipient model. The command is placed in the Region level of the input file. Here is an example of the command:

BC XFER Dirichlet on surface_1 ... Temperature #BC sidesets for recipient mesh

General usage and syntax for the BC XFER command can be found in Chapter 9 of the manual.

5b. Heat Flux BC Commands

For the case where heat flux mapping is used, the above BC XFER command is replaced by: (1) user field definition and (2) heat flux BC specification. Syntax for the user field definition is presented in Chapter 9 of the manual, while its use in a heat flux BC specification is given in Chapter 33.1. Here are examples of the applicable commands:

# ... Create User Field to store mapped heat flux ...
User Field Real Nodal Vector surf_flux on block_1

The above command must be added to the Region level of the input file. In turn, the surf_flux is used to set the heat flux BC via the command block:

# ... Set the heat flux BC ...
Begin Heat Flux Boundary Condition mapped_flux
   add Surface surface_1
   Flux Vector Node Variable = surf_flux
End
Chapter 34

Correlation Heat Transfer Coefficient Reference

34.1 Heat Transfer Coefficient Correlation Library

For heat transfer problems involving convective flux boundary conditions, a convection heat transfer coefficient must be specified (Ch 33.1). An extensive built-in heat transfer coefficient correlation library has been incorporated into Aria (see [38]) for generalized heat transfer problems involving fluid flow. This library includes correlations for laminar and turbulent flows, for internal and external flow geometries, and for gases and liquids (including liquid metals). These correlations are especially useful for reduced order flow modeling in which a full Conjugate Heat Transfer simulation is not desired.

Each correlation supplies a Nusselt Number (Nu) from which the heat transfer coefficient (h) may be computed via

\[ h = \frac{K}{D_h} \times Nu \]  

(34.1)

where \( K \) is the scalar thermal conductivity and \( D_h \) is a characteristic length for the simulation. Hence in all cases, \( D_h \) and \( K \) (material property) MUST be specified. Provision is made for switching between laminar and turbulent correlations based on a user specification for transition Reynolds Number (\( R_{ex} \)).

Currently, there are 74 implemented correlations, each with various mathematical forms, required input variables and valid flow regimes. Usage of the correlation heat transfer coefficient is demonstrated in the outline below.

Begin Heat Transfer Correlation Coefficient ziggurat
LAMINAR CORRELATION = type 18
TURBULENT CORRELATION = type 18

Transition Reynolds Number = 400
Characteristic Length = 0.1
Entrance Length = 0.5

Compute Reynolds Number
Compute Prandtl Number
Compute Wall Temperature
Compute Fluid Temperature
Compute Friction Factor model = smooth tube
End
34.1.1 Classification

Due to the large number of correlations, a taxonomy based on the required input parameters was developed in order to provide a quick mapping between a model and it’s index ID. There are currently 20 parameters which all the correlations are dependent on and these are

- $f$ Friction factor
- $Re$ Flow Reynolds number
- $Pr$ Flow Prandtl number
- $Pr_w$ Flow Prandtl number evaluated with wall conditions
- $T$ Fluid temperature
- $T_w$ Wall temperature
- $L$ Wall length
- $g$ Gravitational constant
- $r$ Ratio of annulus inner diameter to outer diameter
- $\rho$ Fluid density
- $\mu$ Dynamic viscosity
- $\nu$ Kinematic viscosity
- $\beta$ Expansion coefficient
- $\theta$ Wall angle (in radians)
- $V$ Scalar flow velocity magnitude
- $Re_x$ Transitional Reynolds number
- $S_t$ Transverse pitch
- $S_l$ Longitudinal pitch
- $\phi$ Porosity
- $D_e$ Entrance length

These may be computed analytically from the model variables or overridden by a user specified constant. It is to be noted that specification of a parameter constant means that other parameters which are dependent on it will use this value e.g constant kinematic viscosity will also be used in Reynolds number calculations instead of a material property evaluation.

Each correlation specification description below is of the form

$Specification : \{\text{Regime, Phase, Convection, Flow, Author, Shape, Special}\}$

where the component options are

- Regime: \{Laminar, Turbulent, Both\}
- Phase: \{Gas, Liquid, Gas|Liquid, LiquidMetal, Organic\}
Convection: {Forced, Free}

Flow: {Internal, External}

Author: {Hausen, Modified_Sieder-Tate, Churchill-Bernstein, Petukhov, Colburn, Dittus-Boelter, Sieder-Tate, Taylor, Nusselt, Notter-Sliecer, Seban-Shimazaki, Azer-Chao, Sliecer, Denton, Jeschar, Achenbach, Beek, Whitaker, Witte, Ranz-Marshall, Modified_Ranz-Marshall, Hilpert, Zhulkauskas, Churchill-Chu, McAdams, Morgan, Gnielinski, Modified_Gnielinski}

Shape: {NoShape, Cylinder, Sphere, Cube, Square_CornerOn, Square_FlatOn, Hexagon_CornerOn, Hexagon_FlatOn, Flat_Plate_Vertical, TubeBank_Aligned_Cylinder, TubeBank_Staggered_Cylinder, Flat_Plate, Vertical_Or_Angle_Surface, Vertical_Surface, Horizontal_Plate_OnHC, Horizontal_Plate_OffHC, Horizontal_Cylinder, Annulus}

Special: {N/A, Crossflow, Local, Average, PackedBed, FullDevLam_ConstFlux, FullDevLam_ConstTemp, ConstFlux, ConstTemp, Insulated_InnerWall, Insulated_OuterWall}

### 34.1.2 Correlation Listing

Type 1 This correlation applies to cases of fully-developed internal laminar flow with a constant wall flux or constant wall temperature, for which a Nusselt number may be determined analytically.

Dependence: \{ K, D_h \}

Specification: \{ Laminar, Gas|Liquid, Forced, Internal, NoAuthor, NoShape, FullDevLam_ConstFlux \}

Correlation: \( Nu = 4.36 \)

Range: \{ Pr > 0.6 \}

Type 2 This correlation applies to cases of fully-developed internal laminar flow with a constant wall flux or constant wall temperature, for which a Nusselt number may be determined analytically.

Dependence: \{ K, D_h \}

Specification: \{ Laminar, Gas|Liquid, Forced, Internal, NoAuthor, NoShape, FullDevLam_ConstFlux \}

Correlation: \( Nu = 3.666 \)

Range: \{ Pr > 0.6 \}

Type 3 This correlation by Hausen applies to cases of thermally fully-developed laminar flow with entrance effects.

Dependence: \{ K, D_h, D_e, Re, Pr \}

Specification: \{ Laminar, Gas|Liquid, Forced, Internal, Hausen, NoShape, N/A \}

Correlation:

\[
I = \frac{D_h}{D_e} Re Pr
\]

\[
Nu = 3.66 + \frac{0.0668 \cdot I}{1 + 0.4 \cdot I^{0.66667}}
\]

Range: \{ Pr >> 1 \} or unheated entrance

Type 4 This correlation by Nusselt applies to cases of thermally fully-developed turbulent flow with entrance effects.

Dependence: \{ K, D_h, D_e, Re, Pr \}

Specification: \{ Turbulent, Gas|Liquid, Forced, Internal, Nusselt, NoShape, N/A \}

Correlation:

\[
Nu = 0.036 R e^{0.8} P r^{0.333} (D_e/D_h)^{0.05556}
\]
Type 5 This modified Ranz-Marshall correlation applies to cases of thermally fully-developed flow with entrance effects.

Dependence: \{ K, D_h, D_e, Re, Pr \}
Specification: \{ Both, Liquid, Forced, External, Modified_Ranz-Marshall, NoShape, N/A \}
Correlation: \( Nu = 2 + 0.6\sqrt{Re(Pr)^{0.3333} \cdot 25(D_e/D_h)^{-0.7}} \)

Type 6 This modified Sieder-Tate correlation applies to cases of laminar gas flow with entrance effects.

Dependence: \{ K, D_h, D_e, Re, Pr, T, T_w \}
Specification: \{ Laminar, Gas, Forced, Internal, Modified_Sieder-Tate, NoShape, N/A \}
Correlation: \( Nu = 1.86 \cdot \left( \frac{D_h}{D_e} RePr \right)^{0.3333} \cdot \left( \frac{T}{T_w} \right)^{0.098} \)
Range: \{ 0.48 < Pr < 16700 \}

Type 7 This correlation by Taylor applies to cases of turbulent flow with entrance effects.

Dependence: \{ K, D_h, D_e, Re, Pr, T, T_w \}
Specification: \{ Turbulent, Gas|Liquid, Forced, Internal, Taylor, NoShape, N/A \}
Correlation: \( e = 1.59 \frac{D_h}{D_e} - 0.57, C_f = \left( \frac{T}{T_w} \right)^e, Nu = 0.023C_fRe^{0.8}Pr^{0.4}. \)

Type 8 This modified Sieder-Tate correlation applies to cases of laminar liquid flow with entrance effects.

Dependence: \{ K, D_h, D_e, Re, Pr, T, T_w, \mu \}
Specification: \{ Laminar, Liquid, Forced, Internal, Modified_Sieder-Tate, NoShape, N/A \}
Correlation: \( \mu_w = (V^{-0.2661} + \frac{T_w - T}{STRELL})^{-3.758}, Nu = 1.86 \cdot \left( \frac{D_h}{D_e} RePr \right)^{0.3333} \cdot \left( \frac{\mu}{\mu_w} \right)^{0.098} \)
Range: \{ 0.48 < Pr < 16700 \}

Type 9 This correlation by Colburn applies to cases of turbulent flow.

Dependence: \{ K, D_h, Re, Pr \}
Specification: \{ Turbulent, Gas|Liquid, Forced, Internal, Colburn, NoShape, N/A \}
Correlation: \( Nu = \begin{cases} 0.0395Re^{0.75}Pr^{0.3333} & \text{Re} < 4e^4 \\ 0.023Re^{0.8}Pr^{0.3333} & \text{else} \end{cases} \)
Range: \{ 0.7 < Pr < 160 \}, \{ 10000 < Re < 50000 \}

Type 10 This correlation by Notter-Sliecher applies to cases of turbulent flow.

Dependence: \{ K, D_h, Re, Pr \}
Specification: \{ Turbulent, Gas|Liquid, Forced, Internal, Notter-Sliecher, NoShape, N/A \}
Correlation: \( C_1 = 0.88 - 0.24\frac{4}{Pr^{0.24}}, C_2 = 0.33 + 0.5\exp(-0.6Pr), Nu = 5 + 0.016Re^{C_1}Pr^{C_2} \)
Range: \{ 0.1 < Pr < 10000 \}, \{ 10000 < Re < 100000 \}

Type 11 This correlation applies to cases of turbulent organic flow.

Dependence: \{ K, D_h, Re, Pr \}
Specification: \{ Turbulent, Organic, Forced, Internal, NoAuthor, NoShape, N/A \}
Correlation: \( Nu = 0.015Re^{0.85}Pr^{0.33} \)
Type 12 This correlation by Seban-Shimazaki applies to cases of constant surface flux liquid metals in smooth circular tubes.

Dependence: \{ K, D_h, Re, Pr \}  
Specification: \{ Turbulent, LiquidMetal, Forced, Internal, Seban-Shimazaki, NoShape, ConstFlux \}  
Correlation: \[ Pe = RePr, Nu = 5.0 + 0.025Pr^{0.8} \]  
Range: \{100 < Pr < 10000\}, \{3600 < Re < 905000\}  

Type 13 This correlation by Seban-Shimazaki applies to cases of constant surface temperature liquid metals in smooth circular tubes.

Dependence: \{ K, D_h, Re, Pr \}  
Specification: \{ Turbulent, LiquidMetal, Forced, Internal, Seban-Shimazaki, NoShape, ConstTemp \}  
Correlation: \[ Pe = RePr, Nu = 4.82 + 0.185Pr^{0.827} \]  
Range: \{100 < Pr < 10000\}, \{3600 < Re < 905000\}  

Type 14 This correlation by Azer-Chao applies to cases of constant surface temperature liquid metals in tubes.

Dependence: \{ K, D_h, Re, Pr \}  
Specification: \{ Turbulent, LiquidMetal, Forced, Internal, Azer-Chao, NoShape, N/A \}  
Correlation: \[ Pe = RePr, Nu = 5.0 + 0.05Pr^{0.8}Pr^{0.25} \]  

Type 15 This correlation by Sliecher applies to cases of constant surface temperature liquid metals in tubes.

Dependence: \{ K, D_h, Re, Pr \}  
Specification: \{ Both, LiquidMetal, Forced, Internal, Sliecher, NoShape, N/A \}  
Correlation: \[ Pe = RePr, Nu = 4.8 + 0.0156Pr^{0.8}Pr^{0.08} \]  
Range: \{0.004 < Pr < 0.1\}, \{Re < 500000\}  

Type 16 This correlation by Beek applies to cases of heat transfer from wall of packed bed of cylinders to a gas.

Dependence: \{ K, D_h, Re, Pr \}  
Specification: \{ Both, Gas, Forced, External, Beek, Cylinder, PackedBed \}  
Correlation: \[ Nu = 2.58Re^{0.3333}Pr^{0.3333} + 0.22Re^{0.8}Pr^{0.4} \]  
Range: \{40 < Re < 2000\}  

Type 17 This correlation by Beek applies to cases of heat transfer from wall of packed bed of spheres to a gas.

Dependence: \{ K, D_h, Re, Pr \}  
Specification: \{ Both, Gas, Forced, External, Beek, Sphere, PackedBed \}  
Correlation: \[ Nu = 0.203Re^{0.3333}Pr^{0.3333} + 0.22Re^{0.8}Pr^{0.4} \]  
Range: \{40 < Re < 2000\}  

Type 18 This correlation applies to cases of flow over a sphere.

Dependence: \{ K, D_h, Re, Pr \}  
Specification: \{ Both, Gas, Forced, External, NoAuthor, Sphere, N/A \}  
Correlation: \[ Nu = 0.37Re^{0.6} \cdot 1.126Pr^{0.3333} \]  
Range: \{17 < Re < 700000\}  

Type 19 This correlation by Witte applies to cases of liquid metal flow over a sphere.

Dependence: \{ K, D_h, Re, Pr \}  
Specification: \{ Both, LiquidMetal, Forced, External, Witte, Sphere, N/A \}  
Correlation: \[ Nu = 2 + 0.386\sqrt{RePr} \]  
Range: \{36000 < Re < 200000\}
Type 20 This correlation by Ranz-Marshall applies to cases of convection from a free falling liquid drop.

Dependence: \{ K, D_h, Re, Pr \}
Specification: \{ Both, Liquid, Forced, External, Ranz-Marshall, NoShape, N/A \}
Correlation: 
\[ D_e = \text{falling distance}, \quad Nu = 2 + 0.6\sqrt{Pr^{0.3333} \cdot 25 \left( \frac{D_h}{D_e} \right)^{-0.7}} \]

Type 21 This correlation by Zhulkauskas applies to cases of aligned cylindrical tube bank in cross flow.

Dependence: \{ K, D_h, Re, Pr \}
Specification: \{ Both, Gas, Forced, External, Zhulkauskas, TubeBank_Aligned_Cylinder, Crossflow \}
Correlation: 
\[ Nu = C_1 Re^{C_2} Pr^{0.36} \cdot (C_1, C_2) = \begin{cases} 
(0.27, 0.63) & \text{Re < 200000} \\
(0.021, 0.84) & \text{else}
\end{cases} \]

Range: \{ 1 < Re < 1000000 \}, \{ 0.7 < Pr < 500 \}

Type 22 This correlation applies to cases of laminar liquid metal flow over a flat plate.

Dependence: \{ K, D_h, Re, Pr \}
Specification: \{ Laminar, LiquidMetal, Forced, External, NoAuthor, Flat_Plate, N/A \}
Correlation: 
\[ Nu = 0.565\sqrt{RePr} \]
Range: \{ Pe > 100 \}, \{ Pr < 0.05 \}

Type 23 This correlation by Petukhov applies to cases of turbulent gas flow with friction factor correlation for smooth surfaces.

Dependence: \{ K, D_h, Re, Pr, f \}
Specification: \{ Turbulent, Gas, Forced, Internal, Petukhov, NoShape, N/A \}
Correlation: 
\[ Nu = \frac{0.125/RePr}{1.07+12.7\sqrt{0.1257} \left( Pr^{0.6667} - 1 \right)} \]
Range: \{ 3000 < Re < 5000000 \}, \{ 0.5 < Pr < 2000 \}

Type 24 This correlation by Petukhov applies to cases of turbulent liquid flow with friction factor correlation for smooth surfaces.

Dependence: \{ K, D_h, Re, Pr, T, T_w, f, \mu \}
Specification: \{ Turbulent, Liquid, Forced, Internal, Petukhov, NoShape, N/A \}
Correlation: 
\[ \mu_w = \left( \mu^{-0.2661} + \frac{T_w-T}{37.073} \right)^{-3.758}, \quad Nu = \frac{0.125/RePr}{1.07+12.7\sqrt{0.1257} \left( Pr^{0.6667} - 1 \right)} \left( \frac{\mu}{\mu_w} \right)^e \]
\[ e = \begin{cases} 
0.25 & T_w < T \\
0.11 & \text{else}
\end{cases} \]

Range: \{ 3000 < Re < 5000000 \}, \{ 0.5 < Pr < 2000 \}

Type 25 This correlation by Dittus-Boelter applies to cases of turbulent flow.

Dependence: \{ K, D_h, Re, Pr, T, T_w \}
Specification: \{ Turbulent, Gas|Liquid, Forced, Internal, Dittus-Boelter, NoShape, N/A \}
Correlation: 
\[ Nu = 0.023Re^{0.8}Pr^e \]
\[ e = \begin{cases} 
0.3 & T_w < T \\
0.4 & \text{else}
\end{cases} \]

Range: \{ 10000 < Re < 1000000 \}, \{ 0.7 < Pr < 160 \}
Type 26 This correlation by Sieder-Tate applies to cases of turbulent gas flow.

Dependence: \{ K, D_h, Re, Pr, T, T_w \}
Specification: \{ Turbulent, Gas, Forced, Internal, Sieder-Tate, NoShape, N/A \}
Correlation: \( \text{Nu} = 0.027 \text{Re}^{0.8} \text{Pr}^{0.3333} \left( \frac{T}{T_w} \right)^{0.098} \)
Range: \{ 10000 < Re < 100000 \}, \{ 0.7 < Pr < 160 \}

Type 27 This correlation by Whitaker applies to cases of gas flow over a sphere.

Dependence: \{ K, D_h, Re, Pr, T, T_w \}
Specification: \{ Both, Gas, Forced, External, Whitaker, Sphere, N/A \}
Correlation: \( \text{Nu} = 2 + (0.4\sqrt{\text{Re}} + 0.06\text{Re}^{0.6667})\text{Pr}^{0.4} \left( \frac{T}{T_w} \right)^{0.175} \)
Range: \{ 3.5 < Re < 76000 \}, \{ 0.71 < Pr < 380 \}

Type 28 This correlation by Hilpert applies to cases of cylinder in cross flow.

Dependence: \{ K, D_h, Re, Pr, T, T_w \}
Specification: \{ Both, Gas, Forced, External, Hilpert, Cylinder, Crossflow \}
Correlation: \( \text{Re} = \text{Re} \left( \frac{T}{T_f} \right)^{0.7} \), \( \text{Nu} = C_1 \text{Re}^{C_2} \text{Pr}^{0.3333} \)
\[
(C_1, C_2) = \begin{cases} 
(0.989, 0.33) & \text{Re} < 4 \\
(0.911, 0.385) & \text{Re} < 40 \\
(0.683, 0.466) & \text{Re} < 4000 \\
(0.193, 0.618) & \text{Re} < 40000 \\
(0.027, 0.805) & \text{else} 
\end{cases}
\]
Range: \{ 0.4 < Re < 400000 \}

Type 29 This correlation by Hilpert applies to cases of corner-on square in cross flow.

Dependence: \{ K, D_h, Re, Pr, T, T_w \}
Specification: \{ Both, Gas, Forced, External, Hilpert, Square_CornerOn, Crossflow \}
Correlation: \( T_f = \frac{T + T_w}{2} \), \( \text{Re} = \text{Re} \left( \frac{T}{T_f} \right)^{0.7} \), \( C_1 = 0.246 \), \( C_2 = 0.588 \), \( \text{Nu} = C_1 \text{Re}^{C_2} \text{Pr}^{0.3333} \)
Range: \{ 5000 < Re < 100000 \}

Type 30 This correlation by Hilpert applies to cases of flat-on square in cross flow.

Dependence: \{ K, D_h, Re, Pr, T, T_w \}
Specification: \{ Both, Gas, Forced, External, Hilpert, Square_FlatOn, Crossflow \}
Correlation: \( T_f = \frac{T + T_w}{2} \), \( \text{Re} = \text{Re} \left( \frac{T}{T_f} \right)^{0.7} \), \( C_1 = 0.102 \), \( C_2 = 0.675 \), \( \text{Nu} = C_1 \text{Re}^{C_2} \text{Pr}^{0.3333} \)
Range: \{ 5000 < Re < 100000 \}

Type 31 This correlation by Hilpert applies to cases of corner-on hexagon in cross flow.

Dependence: \{ K, D_h, Re, Pr, T, T_w \}
Specification: \{ Both, Gas, Forced, External, Hilpert, Hexagon_CornerOn, Crossflow \}
Correlation: \( T_f = \frac{T + T_w}{2} \), \( \text{Re} = \text{Re} \left( \frac{T}{T_f} \right)^{0.7} \), \( C_1 = 0.153 \), \( C_2 = 0.638 \), \( \text{Nu} = C_1 \text{Re}^{C_2} \text{Pr}^{0.3333} \)
Range: \{ 5000 < Re < 100000 \}
Type 32 This correlation by Hilpert applies to cases of flat-on hexagon in cross flow.

Dependence: \{ K, D_h, Re, Pr, T, T_w \}
Specification: \{ Both, Gas, Forced, External, Hilpert, Hexagon_FlatOn, Crossflow \}
Correlation: \( T_f = \frac{T + T_w}{2}, \ Re = Re\left(\frac{T}{T_f}\right)^{0.7}, \ Nu = C_1 Re C_2 Pr^{0.3333} \)

\[ (C_1, C_2) = \begin{cases} 
(0.160, 0.638) & \text{Re} < 1.95e^4 \\
(0.0385, 0.782) & \text{else}
\end{cases} \]

Range: \{ 5000 < Re < 1000000 \}

Type 33 This correlation by Hilpert applies to cases of vertical flat plate in cross flow.

Dependence: \{ K, D_h, Re, Pr, T, T_w \}
Specification: \{ Both, Gas, Forced, External, Hilpert, Flat_Plate_vertical, Crossflow \}
Correlation: \( T_f = \frac{T + T_w}{2}, \ Re = Re\left(\frac{T}{T_f}\right)^{0.7}, \ C_1 = 0.228, \ C_2 = 0.731, \ Nu = C_1 Re C_2 Pr^{0.3333} \)

\[ (C_1, C_2) = \begin{cases} 
(0.989, 0.33) & \text{Re} < 4 \\
(0.911, 0.385) & \text{Re} < 40 \\
(0.683, 0.466) & \text{Re} < 4000 \\
(0.027, 0.805) & \text{else}
\end{cases} \]

\[ C_3 = \begin{cases} 
0.36 & \text{Pr} > 10 \\
0.37 & \text{else}
\end{cases} \]

Range: \{ 1 < Re < 1000000 \}, \{ 0.7 < Pr < 500 \}

Type 34 This correlation by Zhulkauskas applies to cases of cylinder in cross flow.

Dependence: \{ K, D_h, Re, Pr, T, T_w \}
Specification: \{ Both, Gas, Forced, External, Zhulkauskas, Cylinder, Crossflow \}
Correlation: \( T_f = \frac{T + T_w}{2}, \ Re = Re\left(\frac{T}{T_f}\right)^{0.7}, \ Nu = C_1 Re C_2 Pr C_3 \)

\[ (C_1, C_2) = \begin{cases} 
(0.989, 0.33) & \text{Re} < 4 \\
(0.911, 0.385) & \text{Re} < 40 \\
(0.683, 0.466) & \text{Re} < 4000 \\
(0.027, 0.805) & \text{else}
\end{cases} \]

\[ C_3 = \begin{cases} 
0.36 & \text{Pr} > 10 \\
0.37 & \text{else}
\end{cases} \]

Range: \{ 1 < Re < 1000000 \}, \{ 0.7 < Pr < 500 \}

Type 35 This correlation by Churchill-Bernstein applies to cases of cylinder in cross flow.

Dependence: \{ K, D_h, Re, Pr, T, T_w \}
Specification: \{ Both, Gas, Forced, External, Churchill-Bernstein, Cylinder, Crossflow \}
Correlation: \( T_f = \frac{T + T_w}{2}, \ Re = Re\left(\frac{T}{T_f}\right)^{0.7}, \ Nu = \frac{0.3 + 0.62\sqrt{RePr^{0.3333}}}{(1 + (0.47/Pr^{0.6667})^{0.25})(1 + (Re/282000)^{0.625})^{0.8}} \)

Range: \{ 0.2 < Re * Pr \}

Type 36 This correlation applies to cases of flow over a flat plate.

Dependence: \{ K, D_h, Re, Pr, T, T_w \}
Specification: \{ Laminar, Gas, Forced, External, NoAuthor, Flat_Plate, Local \}
Correlation: \( T_f = \frac{T + T_w}{2}, \ Re = Re \cdot (T/T_f)^{0.7}, \ Nu = 0.332\sqrt{RePr^{0.3333}} \)

Range: \{ 0.6 < Pr \}

Type 37 This correlation applies to cases of flow over a flat plate.

Dependence: \{ K, D_h, Re, Pr, T, T_w \}
Type 38 This correlation applies to cases of flow over a flat plate.

Dependence: \{K, D_h, Re, Pr, T, T_w, \mu\} 
Specification: \{Turbulent, Gas, Forced, External, NoAuthor, Flat_Plate, N/A\} 
Correlation: \[ T_f = \frac{T + T_w}{2}, \quad Re = Re \cdot \left(\frac{T}{T_f}\right)^{0.7}, \quad Nu = 0.664\sqrt{RePr^{0.3333}} \]
Range: \{0.6 < Pr\} 

Type 39 This correlation by Sieder-Tate applies to cases of turbulent liquid flow.

Dependence: \{K, D_h, Re, Pr, T, T_w, \mu\} 
Specification: \{Turbulent, Liquid, Forced, Internal, Sieder-Tate, NoShape, N/A\} 
Correlation: \[ \mu_w = \left(V^{-0.2661} + \frac{T_w - T}{37.073}\right)^{-3.758}, \quad Nu = 0.027Re^{0.8}Pr^{0.3333} \left(\frac{\mu}{\mu_w}\right)^{0.14} \]
Range: \{10000 < Re < 1000000\}, \{0.7 < Pr < 160\} 

Type 40 This correlation by Whitaker applies to cases of liquid flow over a sphere.

Dependence: \{K, D_h, Re, Pr, T, T_w, \mu\} 
Specification: \{Both, Liquid, Forced, External, Whitaker, Sphere, N/A\} 
Correlation: \[ \mu_w = \left(\mu^{-0.2661} + \frac{T_w - T}{37.073}\right)^{-3.758}, \quad Nu = 2 + (0.48\sqrt{Re} + 0.069Re^{0.6667})Pr^{0.4} \left(\frac{\mu}{\mu_w}\right)^{0.25} \]
Range: \{3.5 < Re < 76000\}, \{0.7 < Pr < 380\} 

Type 41 This correlation by Hilpert applies to cases of cylinder in crossflow.

Dependence: \{K, D_h, Re, Pr, T, T_w, \mu\} 
Specification: \{Both, Liquid, Forced, External, Hilpert, Cylinder, Crossflow\} 
Correlation: 
\[ T_f = \frac{T + T_w}{2}, \quad \mu_w = \left(\mu^{-0.2661} + \frac{T_w - T}{37.073}\right)^{-3.758}, \quad Re = Re \frac{\mu}{\mu_w}, \quad Pr = Pr \frac{\mu}{\mu_w} \]
\[ Nu = C_1Re^{C_2}Pr^{0.3333} \]
\[ (C_1, C_2) = \begin{cases} 
(0.989, 0.33) & \text{Re} < 4 \\
(0.911, 0.385) & \text{Re} < 40 \\
(0.683, 0.466) & \text{Re} < 4000 \\
(0.193, 0.618) & \text{Re} < 40000 \\
(0.027, 0.805) & \text{else} 
\end{cases} \]
Range: \{0.4 < Re < 400000\} 

Type 42 This correlation by Zhulkauskas applies to cases of cylinder in crossflow.

Dependence: \{K, D_h, Re, Pr, T, T_w, \mu\} 
Specification: \{Both, Liquid, Forced, External, Zhulkauskas, Cylinder, Crossflow\}
Correlation: \( \mu_w = \left( \mu^{-0.2661} + \frac{T_w - T}{37.073} \right)^{-3.758} \), \( \text{Nu} = C_1 \text{Re}^C_2 \text{Pr}^C_3 \left( \frac{\mu}{\mu_w} \right)^{0.25} \)

\[
(C_1, C_2) = \begin{cases} 
(0.75, 0.4) & \text{Re} < 40 \\
(0.51, 0.5) & \text{Re} < 1000 \\
(0.26, 0.6) & \text{Re} < 200000 \\
(0.076, 0.7) & \text{else}
\end{cases}
\]

\[
C_3 = \begin{cases} 
0.36 & \text{Pr} > 10 \\
0.37 & \text{else}
\end{cases}
\]

Range: \{1 < Re < 1000000\}, \{0.7 < Pr < 500\}

Type 43 This correlation by Churchill-Bernstein applies to cases of cylinder in crossflow.

Dependence: \{K, D_h, Re, Pr, T, T_w, \mu\}

Specification: \{Both, Liquid, Forced, External, Churchill-Bernstein, Cylinder, Crossflow\}

Correlation:

\[
T_f = \frac{T + T_w}{2}, \mu_f = \left( \mu^{-0.2661} + \frac{T_f - T}{37.073} \right)^{-3.758}, \mu_{cf} = \frac{\mu}{\mu_f}, \text{Re} = \text{Re} \cdot \mu_{cf}, \text{Pr} = \text{Pr} \cdot \mu_{cf}
\]

\[
\text{Nu} = 0.3 + 0.62 \sqrt{\text{Re}^0.3333 (1 + (\text{Re}/282000)^{0.625})^0.8}
\]

Range: \{0.2 < Re * Pr\}

Type 44 This correlation applies to cases of laminar flow over a flat plate.

Dependence: \{K, D_h, Re, Pr, T, T_w, \mu\}

Specification: \{Laminar, Liquid, Forced, External, NoAuthor, Flat_Plate, Local\}

Correlation:

\[
T_f = \frac{T + T_w}{2}, \mu_f = \left( \mu^{-0.2661} + \frac{T_f - T}{37.073} \right)^{-3.758}, \mu_{cf} = \frac{\mu}{\mu_f}, \text{Re} = \text{Re} \cdot \mu_{cf}, \text{Pr} = \text{Pr} \cdot \mu_{cf}
\]

\[
\text{Nu} = 0.332 \sqrt{\text{Re} \text{Pr}^0.3333}
\]

Range: \{0.6 < Pr\}

Type 45 This correlation by Zhulkauskas applies to cases of aligned cylindrical tube bank in crossflow.

Dependence: \{K, D_h, Re, Pr, T, T_w, \mu\}

Specification: \{Both, Liquid, Forced, External, Zhulkauskas, TubeBank_Aligned_Cylinder, Crossflow\}

Correlation:

\[
\mu_w = \left( \mu^{-0.2661} + \frac{T_w - T}{37.073} \right)^{-3.758}, \text{Nu} = C_1 \text{Re}^C_2 \text{Pr}^C_3 \left( \frac{\mu}{\mu_w} \right)^{0.25}
\]

\[
(C_1, C_2) = \begin{cases} 
(0.27, 0.63) & \text{Re} < 200000 \\
(0.021, 0.84) & \text{else}
\end{cases}
\]

Range: \{1000 < Re < 200000\}, \{0.7 < Pr < 500\}
Type 46 This correlation applies to cases of laminar flow over a flat plate.

Dependence: {K, D_h, Re, Pr, T, T_w, μ}
Specification: {Laminar, Liquid, Forced, External, NoAuthor, Flat_Plate, Average}
Correlation:
\[ T_f = \frac{T + T_w}{2}, \mu_f = \left(\mu - 0.2661 + \frac{T_f - T}{37.073}\right)^{-3.758}, \mu_{cf} = \frac{\mu}{\mu_f}, Re = Re \cdot \mu_{cf}, Pr = \frac{Pr}{\mu_{cf}} \]
\[ Nu = 0.664\sqrt{Re Pr^{0.3333}} \]

Range: \{0.6 < Pr\}

Type 47 This correlation applies to cases of turbulent flow over a flat plate.

Dependence: {K, D_h, Re, Pr, T, T_w, μ}
Specification: {Turbulent, Liquid, Forced, External, NoAuthor, Flat_Plate, N/A}
Correlation:
\[ T_f = \frac{T + T_w}{2}, \mu_f = \left(\mu - 0.2661 + \frac{T_f - T}{37.073}\right)^{-3.758}, \mu_{cf} = \frac{\mu}{\mu_f}, Re = Re \cdot \mu_{cf}, Pr = \frac{Pr}{\mu_{cf}} \]
\[ Nu = 0.0296Re^{0.8} Pr^{0.3333} \]

Range: \{0.6 < Pr < 60\}

Type 48 This correlation by Denton applies to cases of flow over a packed bed with porosity = 0.37.

Dependence: {K, D_h, Re, Pr, φ}
Specification: {Both, Gas|Liquid, Forced, External, Denton, NoShape, PackedBed}
Correlation: Nu = 0.8Re^{0.7} Pr^{0.3333}
Range: \{500 < Pr < 50000\}, \{φ = 0.37\}

Type 49 This correlation by Achenbach applies to cases of flow of a packed bed.

Dependence: {K, D_h, Re, Pr, φ}
Specification: {Both, Gas|Liquid, Forced, External, Achenbach, N/A, PackedBed}
Correlation: \[ P = 1 - φ, Nu = 0.70731^e \left(\frac{Re}{Pr}\right)^{2.32} + 6.34e^{-4}\left(\frac{Re}{Pr}\right)^3 \] \[ e = \begin{cases} 1 & \text{sphere} \\ 0.79 & \text{cylinder with } L/D = 1 \\ 0.71 & \text{cube} \end{cases} \]
\[ Pr^{0.3333} \]
Range: {Re < 300000}

Type 50|51|52 These correlations apply to cases of flow over a packed bed with Pr = 0.7 for various shapes.

Dependence: {K, D_h, Re, Pr, φ}
Correlation: Nu = 2.065^e Re^{0.425} Pr^{0.3333},
\[ e = \begin{cases} 1 & \text{sphere} \\ 0.79 & \text{cylinder with } L/D = 1 \\ 0.71 & \text{cube} \end{cases} \]
Range: {90 < Re < 4000}

[Type 50] For spheres: {Both, Gas, Forced, External, NoAuthor, Sphere, PackedBed}
[Type 51] For cylinders: {Both, Gas, Forced, External, NoAuthor, Cylinder, PackedBed}
[Type 52] For cubes: {Both, Gas, Forced, External, NoAuthor, Cube, PackedBed}
Type 53 This correlation by Jeschar applies to cases of flow of a packed bed.

Dependence: \{ K, D_h, Re, \phi \}
Specification: \{ Both, Gas|Liquid, Forced, External, Jeschar, N/A, PackedBed \}
Correlation: 
\[ P = 1 - \phi, \quad C = \frac{P}{\phi}, \quad Nu = 2 + \left( \left( (CRe)^{0.5} + 0.005 \frac{Re}{\phi} \right) \right) \]
Range: \{ 250 < Re < 55000 \}

Type 54 This correlation by Zhulkauskas applies to cases of staggered cylindrical tube bank in crossflow.

Dependence: \{ K, D_h, Re, S_t, S_l \}
Specification: \{ Both, Gas, Forced, External, Zhulkauskas, TubeBank_Staggered_Cylinder, Crossflow \}
Correlation: 
\[ Nu = C_1 Re^{C_2 Pr^{0.36}} \]
\[ x = \begin{cases} 
0.4 & S_t / S_l > 2 \\
0.35 \cdot (S_t / S_l)^{0.2} & S_t / S_l < 2 
\end{cases} \]
\[ (C_1, C_2) = \begin{cases} 
(x, 0.6) & Re < 200000 \\
(0.022, 0.84) & \text{else} 
\end{cases} \]
Range: \{ 1000 < Re < 2000000 \}, \{ 0.7 < Pr < 500 \}

Type 55 This correlation by Zhulkauskas applies to cases of staggered cylindrical tube bank in crossflow.

Dependence: \{ K, D_h, Re, Pr, T, T_w, V, S_t, S_l \}
Specification: \{ Both, liquid, Forced, External, Zhulkauskas, TubeBank_Staggered_Cylinder, Crossflow \}
Correlation: 
\[ V_w = \left( V^{-0.2661} + \frac{T_w - T}{37.973} \right)^{-3.758}, \quad Nu = C_1 Re^{C_2 Pr^{0.36}} \left( \frac{V}{V_w} \right)^{0.25} \]
\[ x = \begin{cases} 
0.4 & S_t / S_l > 2 \\
0.35 \cdot (S_t / S_l)^{0.2} & S_t / S_l < 2 
\end{cases} \]
\[ (C_1, C_2) = \begin{cases} 
(x, 0.6) & Re < 200000 \\
(0.022, 0.84) & \text{else} 
\end{cases} \]
Range: \{ 1000 < Re < 2000000 \}, \{ 0.7 < Pr < 500 \}

Type 56 This correlation applies to cases of flow over a flat plate.

Dependence: \{ K, D_h, Re, Pr, T, T_w, V, Re_x \}
Specification: \{ Both, Gas, Forced, External, NoAuthor, Flat_Plate, N/A \}
Correlation: 
\[ x = 0.037 Re^{0.8}_{Re_x} - 0.664 Re^{0.5}_{Re_x} ; T_f = \frac{T + T_w}{2} ; Re = Re \cdot \left( \frac{T}{T_f} \right)^{0.7} \]
\[ Nu = \left( 0.037 Re^{0.8} - x \right) Pr^{0.3333} \]
Range: \{ 5e^5 < Re < 1e^8 \}, \{ 0.6 < Pr < 60 \}

Type 57 This correlation applies to cases of flow over a flat plate.

Dependence: \{ K, D_h, Re, Pr, T, T_w, V, Re_x \}
Specification: {Both, Liquid, Forced, External, NoAuthor, Flat_Plate, N/A}
Correlation:

\[ x = 0.037Re_x^{0.8} - 0.664Re_x^{0.5}, T_f = \frac{T + T_w}{2}, V_f = \left(V - 0.2661 + \frac{T_f - T}{37.073}\right)^{-3.758} \]

\[ V_{cf} = \frac{V}{V_f}, Re = Re \cdot V_{cf}, Pr = Pr/V_{cf} \]

\[ Nu = \left(0.037Re^{0.8} - x\right)Pr^{0.3333} \]

Range: \(5 \times 10^5 < Re < 1 \times 10^8\), \(0.6 < Pr < 60\)

Type 58 This correlation by Churchill-Chu applies to free convection from a vertical surface, or non-vertical surface if angle is less than 60° (measured from vertical).

Dependence: \{K, D_h, Pr, T, T_w, L, g, \rho, \nu, \theta\}
Specification: {Laminar, Gas, Free, External, Churchill-Chu, Vertical_Or_Angle_Surface, N/A}
Correlation:

\[ x = \beta L^3 \cos(\theta), T_f = \frac{T + T_w}{2}, \rho = \frac{T}{T_f}, \nu_f = \nu(T_f/T)^{0.7} \]

\[ Gr = gx(\rho)^{2} \frac{|T_w - T|}{T_f \nu_f^2}, Ra = GrPr \]

\[ Nu = 0.68 + 0.67 \frac{Ra^{0.25}}{(1 + (0.492/Pr)^{0.565})^{0.444}} \]

Range: \(Ra < 1 \times 10^9\), \(\theta < 60°\)

Type 59 This correlation by Churchill-Chu applies to free convection from a vertical surface, or non-vertical surface if angle is less than 60° (measured from vertical).

Dependence: \{K, D_h, Pr, T, T_w, L, g, \rho, \nu, \theta, \beta\}
Specification: {Laminar, Liquid, Free, External, Churchill-Chu, Vertical_Or_Angle_Surface, N/A}
Correlation:

\[ x = \beta L^3 \cos(\theta), T_f = \frac{T + T_w}{2}, \rho = \frac{T}{T_f}, \nu_f = \nu(T_f/T)^{0.7} \]

\[ Gr = gx(\rho)^{2} \frac{|T_w - T|}{T_f \nu_f^2}, Ra = GrPr \]

\[ Nu = 0.68 + 0.67 \frac{Ra^{0.25}}{(1 + (0.492/Pr)^{0.565})^{0.444}} \]

Range: \(Ra < 1 \times 10^9\), \(\theta < 60°\)

Type 60 This correlation by Churchill-Chu applies to free convection from a vertical surface.

Dependence: \{K, D_h, Pr, T, T_w, L, g, \rho, \nu\}
Specification: {Both, Gas, Free, External, Churchill-Chu, Vertical_surface, N/A}
Correlation:
\[ x = L^3, T_f = \frac{T + T_w}{2}, \rho = \rho \frac{T}{T_f}, \nu_f = \nu(T_f/T)^{0.7} \]
\[ Gr = g x (\rho)^2 \frac{|T_w - T|}{T_f \nu_f^2}, Ra = Gr Pr \]
\[ Nu = \left( 0.825 + \frac{0.387 Ra^{0.1667}}{1 + (0.492/Pr)^{0.5265}0.2963} \right)^2 \]

Type 61 This correlation by McAdams applies to free convection from a horizontal surface; either the upper surface of a heated plate or the lower surface of a cooled plate.

Dependence: \{ K, D_h, Pr, T, T_w, L, g, \rho, \nu \} 
Specification: \{ Both, Gas, Free, External, McAdams, Horizontal_Plate_OnHC, N/A \} 
Correlation: \[ x = L^3, T_f = \frac{T + T_w}{2}, \rho = \rho \frac{T}{T_f}, \nu_f = \nu(T_f/T)^{0.7}, Gr = g x (\rho)^2 \frac{|T_w - T|}{T_f \nu_f^2}, Ra = Gr Pr \]
\[ Nu = \begin{cases} 0.54 Ra^{0.25} \hspace{1cm} Ra < 1e^7 \\ 0.15 Ra^{0.333} \hspace{1cm} \text{else} \end{cases} \]

Range: \{ 1e^4 < Ra < 1e^{11} \}

Type 62 This correlation by McAdams applies to free convection from a horizontal surface; either the upper surface of a cooled plate or the lower surface of a heated plate.

Dependence: \{ K, D_h, Pr, T, T_w, L, g, \rho, \nu \} 
Specification: \{ Both, Gas, Free, External, McAdams, Horizontal_Plate_OffHC, N/A \} 
Correlation: \[ x = L^3, T_f = \frac{T + T_w}{2}, \rho = \rho \frac{T}{T_f}, \nu_f = \nu(T_f/T)^{0.7}, Gr = g x (\rho)^2 \frac{|T_w - T|}{T_f \nu_f^2}, Ra = Gr Pr \]
\[ Nu = 0.27 Ra^{0.25} \]

Range: \{ 1e^3 < Ra < 1e^{10} \}

Type 63 This correlation by Morgan applies to free convection from a horizontal cylinder.

Dependence: \{ K, D_h, Pr, T, T_w, L, g, \rho, \nu \} 
Specification: \{ Both, Gas, Free, External, Morgan, Horizontal_Cylinder, N/A \} 
Correlation: \[ x = L^3, T_f = \frac{T + T_w}{2}, \rho = \rho \frac{T}{T_f}, \nu_f = \nu(T_f/T)^{0.7}, Gr = g x (\rho)^2 \frac{|T_w - T|}{T_f \nu_f^2}, Ra = Gr Pr \]
\[ Nu = C_1 Ra C_2 \]
\[ (C_1, C_2) = \begin{cases} (0.675, 0.058) \hspace{1cm} Ra < 1e^{-2} \\ (1.02, 0.148) \hspace{1cm} Ra < 100 \\ (0.85, 0.188) \hspace{1cm} Ra < 10000 \\ (0.48, 0.25) \hspace{1cm} Ra < 1e^7 \\ (0.125, 0.333) \hspace{1cm} \text{else} \end{cases} \]

Range: \{ 1e^{-10} < Ra < 1e^{12} \}
Type 64 This correlation by Churchill-Chu applies to free convection from a horizontal cylinder.

Dependence: \{ K, D_h, Pr, T, T_w, L, g, \rho, \nu \}
Specification: \{ Both, Gas, Free, External, Churchill-Chu, Horizontal_Cylinder, N/A \}
Correlation:
\[
x = L^3, T_f = \frac{T + T_w}{2}, \rho = \frac{T}{T_f}, \nu_f = \nu(T_f/T)^{0.7}, Gr = gx(\rho)^2 \frac{|T_w - T|}{T_f \nu_f^2}, Ra = Gr Pr
\]
\[
Nu = \left( 0.6 + \frac{0.387 Ra^{0.1667}}{(1 + (0.559/Pr)^{0.5625}0^{0.2963})^2} \right)^2
\]
Range: \{1e^{-5} < Ra < 1e^{12}\}

Type 65 This correlation by Churchill-Chu applies to free convection from a sphere.

Dependence: \{ K, D_h, Pr, T, T_w, L, g, \rho, \nu \}
Specification: \{ Both, Gas, Free, External, Churchill-Chu, Sphere, N/A \}
Correlation:
\[
x = L^3, T_f = \frac{T + T_w}{2}, \rho = \frac{T}{T_f}, \nu_f = \nu(T_f/T)^{0.7}, Gr = gx(\rho)^2 \frac{|T_w - T|}{T_f \nu_f^2}, Ra = Gr Pr
\]
\[
Nu = 2 + \frac{0.589 Ra^{0.25}}{(1 + (0.469/Pr)^{0.5625}0^{0.4444})}
\]
Range: \{1e^{11} < Ra, 0.7 < Pr\}

Type 66 This correlation by Churchill-Chu applies to free convection from a vertical surface.

Dependence: \{ K, D_h, Pr, T, T_w, L, g, \rho, \nu, \beta \}
Specification: \{ Both, Liquid, Free, External, Churchill-Chu, Vertical_Surface, N/A \}
Correlation:
\[
x = \beta L^3, T_f = \frac{T + T_w}{2}, \nu_f = \left( \nu^{-0.2661} + \frac{T_f - T}{37.073} \right)^{-3.758}, Pr = Pr \cdot \nu_f / \nu
\]
\[
Gr = gx(\rho)^2 \frac{|T_w - T|}{T_f \nu_f^2}, Ra = Gr Pr
\]
\[
Nu = \left( 0.825 + \frac{0.387 Ra^{0.1667}}{(1 + (0.492/Pr)^{0.5625}0^{0.2963})^2} \right)^2
\]

Type 67 This correlation by McAdams applies to free convection from a horizontal plate; either the upper surface of a heated plate or the lower surface of a cooled plate.

Dependence: \{ K, D_h, Pr, T, T_w, L, g, \rho, \nu, \beta \}
Specification: \{ Both, Liquid, Free, External, McAdams, Horizontal_Plate_OnHC, N/A \}
Correlation:
\[
x = \beta L^3, T_f = \frac{T + T_w}{2}, \nu_f = \left( \nu^{-0.2661} + \frac{T_f - T}{37.073} \right)^{-3.758}, Pr = Pr \cdot \nu_f / \nu
\]
\[
Gr = gx(\rho)^2 \frac{|T_w - T|}{T_f \nu_f^2}, Ra = Gr Pr
\]
\[ N_u = \begin{cases} 
0.54Ra^{0.25} & \text{Ra} < 1e^7 \\
0.15Ra^{0.3333} & \text{else}
\end{cases} \]

Range: \(\{1e^4 < Ra < 1e^{11}\}\)

Type 68 This correlation by McAdams applies to free convection from a horizontal plate; either the upper surface of a cooled plate or the lower surface of a heated plate.

Dependence: \(\{K, D_h, \Pr, T, T_w, L, g, \rho, \nu, \beta\}\)

Specification: \(\{\text{Both, Liquid, Free, External, McAdams, Horizontal\_Plate\_OffHC, N/A}\}\)

Correlation:

\[
x = \beta L^3, T_f = \frac{T + T_w}{2}, \nu_f = \left(\nu^{-0.2661} + \frac{T_f - T}{37.073}\right)^{-3.758}, Pr = Pr \cdot \nu_f / \nu
\]

\[
Gr = gx(\rho)^2 \frac{|T_w - T|}{T_f^2}, Ra = GrPr
\]

\[
N_u = 0.27Ra^{0.25}
\]

Range: \(\{1e^5 < Ra < 1e^{10}\}\)

Type 69 This correlation by Morgan applies to free convection from a horizontal cylinder.

Dependence: \(\{K, D_h, \Pr, T, T_w, L, g, \rho, \nu, \beta\}\)

Specification: \(\{\text{Both, Liquid, Free, External, Morgan, Horizontal\_Cylinder, N/A}\}\)

Correlation:

\[
x = \beta L^3, T_f = \frac{T + T_w}{2}, \nu_f = \left(\nu^{-0.2661} + \frac{T_f - T}{37.073}\right)^{-3.758}, Pr = Pr \cdot \nu_f / \nu
\]

\[
Gr = gx(\rho)^2 \frac{|T_w - T|}{T_f^2}, Ra = GrPr
\]

\[
N_u = C_1Ra^{C_2}
\]

\[
(C_1, C_2) = \begin{cases} 
(0.675, 0.058) & \text{Ra} < 1e^{-2} \\
(1.02, 0.148) & \text{Ra} < 100 \\
(0.85, 0.188) & \text{Ra} < 10000 \\
(0.48, 0.25) & \text{Ra} < 1e^7 \\
(0.125, 0.333) & \text{else}
\end{cases}
\]

Range: \(\{1e^{-10} < Ra < 1e^{12}\}\)

Type 70 This correlation by Churchill-Chu applies to free convection from a horizontal cylinder.

Dependence: \(\{K, D_h, \Pr, T, T_w, L, g, \rho, \nu, \beta\}\)

Specification: \(\{\text{Both, Liquid, Free, External, Churchill-Chu, Horizontal\_Cylinder, N/A}\}\)

Correlation:

\[
x = \beta L^3, T_f = \frac{T + T_w}{2}, \nu_f = \left(\nu^{-0.2661} + \frac{T_f - T}{37.073}\right)^{-3.758}, Pr = Pr \cdot \nu_f / \nu
\]

\[
Gr = gx(\rho)^2 \frac{|T_w - T|}{T_f^2}, Ra = GrPr
\]

\[
N_u = \left(0.6 + \frac{0.387Ra^{0.1667}}{1 + (0.559/Pr)^{0.5625}(0.2963)}\right)^2
\]

976
Range: $\{10^{-5} < Ra < 10^{12}\}$

Type 71 This correlation by Churchill-Chu applies to free convection from a sphere.

Dependence: \{K, D_h, Pr, T, T_w, L, g, \rho, \nu, \beta\}
Specification: \{Both, Liquid, Free, External, Churchill-Chu, Sphere, N/A\}
Correlation:
\[
x = \beta L^3, T_f = \frac{T + T_w}{2}, \nu_f = \left( \nu^{-0.2661} + T_f - T \right)^{-3.758}, Pr = Pr \cdot \nu_f / \nu
\]
\[
Gr = gx(\rho) \frac{|T_w - T|}{T_f \nu_f^2}, Ra = Gr Pr
\]
\[
Nu = 2 + 0.589 Ra^{0.25} \left( \frac{1}{1 + (0.469/Pr)^{0.5625}} \right)^{0.4444}
\]

Range: $\{10^{11} < Ra\}$, $\{0.7 < Pr\}$

Type 72 This correlation by Gnielinski applies to cases of turbulent gas flow in a circular tube with friction factor correlation for smooth surfaces.

Dependence: \{K, D_h, Re, Pr, f\}
Specification: \{Turbulent, Gas, Forced, Internal, Gnielinski, Horizontal_Cylinder, N/A\}
Correlation: \[Nu = \frac{\frac{4}{3}(Re-1000)Pr}{1+12.7 \sqrt{\frac{r}{h}} (Pr^{0.6667} - 1)}\]
Range: $\{3000 < Re < 5000000\}$, $\{0.5 < Pr < 2000\}$

Type 73 This modified correlation by Gnielinski applies to cases of turbulent flow in an annulus with friction factor correlation for annular surfaces where the inner wall is heated and the outer wall is insulated.

Dependence: \{K, D_h, Re, Pr, f, r, T, T_w\}
Specification: \{Turbulent, Gas|Liquid, Forced, Internal, Modified_Gnielinski, Annulus, Insulated_OuterWall\}
Correlation:
\[
F = 0.75 r^{-0.17}, b = 1.07 + \frac{900}{Re} - \frac{0.63}{1 + 10 Pr}
\]
\[
Nu = c F \frac{4 Re Pr}{b + 12.7 \sqrt{\frac{r}{h}} (Pr^{0.6667} - 1)}
\]
\[
c = \begin{cases} 
1 & \text{GAS and } T > T_w \\
\left( \frac{T}{T_w} \right)^{0.45} & \text{GAS and } T < T_w \\
\left( \frac{Pr}{Pr_w} \right)^{0.11} & \text{LIQUID}
\end{cases}
\]
Range: $\{10000 < Re\}$

Type 74 This modified correlation by Gnielinski applies to cases of turbulent flow in an annulus with friction factor correlation for annular surfaces where the inner wall is insulated and the outer wall is heated.

Dependence: \{K, D_h, Re, Pr, f, r, T, T_w, Pr_w\}
Specification: \{Turbulent, Gas|Liquid, Forced, Internal, Modified_Gnielinski, Annulus, Insulated_InnerWall\}
Correlation:

\[ F = 0.9 - 0.15r^{0.6}, b = 1.07 + \frac{900}{Re} - \frac{0.63}{1 + 10Pr} \]

\[ Nu = cF \frac{\frac{4}{5}RePr}{b + 12.7\sqrt{\frac{f}{Re}}(Pr^{0.667} - 1)} \]

\[ c = \begin{cases} 
1 & \text{GAS and } T > T_w \\
(T/T_w)^{0.45} & \text{GAS and } T < T_w \\
(P_{Pr}/P_{Pr_w})^{0.11} & \text{LIQUID} 
\end{cases} \]

Range: \{10000 < Re\}

### 34.1.3 Visualization

Aria provides a means of outputting the heat transfer coefficient via the solution options command block. It must be noted that for a given surface associated with the correlation heat transfer coefficient, such as in a convective boundary condition coupled to an advective bar, only the actual surface can be used for output. This means that an attempt to construct a sideset on an advective bar and output the coefficients on the bar will fail because it is not a part of the associated boundary condition. An example of the syntax is given below:

```
begin convective flux boundary condition insideWall
  add surface surface_2
  use advective bar airBar bulknodes
  USE CORRELATION CONVECTION MODEL FlowInAnnulus
end convective flux boundary condition insideWall

begin solution options
  post process nodal normalized heat_transfer_coefficient on surface_2 as HTCp
end
```
Fluid Phase {=} FluidPhase
Friction Factor {=} $f$
Gnielinski Film Gradient Exponent {=} Exponent
Gravitational Constant {=} $g$
Kinematic Viscosity {=} $\nu$
Laminar Correlation {=} Type CorrelationType
Longitudinal Pitch {=} $Sl$
Porosity {=} $\phi$
Prandtl Number {=} $Pr$
Reynolds Number {=} $Re$
Turbulent Correlation {=} Type CorrelationType
Temperature {=} $t$
Transition Reynolds Number {=} $Re_x$
Transverse Pitch {=} $St$
Velocity {=} $v$
Wall Angle {=} $\theta$
Wall Length {=} $l$
Wall Prandtl Number {=} $Pr_w$
Wall Temperature {=} $T_w$

Summary
Specified heat transfer correlation coefficient model.

Description
Enables the user to specify a correlation model for use with computing heat transfer coefficients in conjunction with advective bar networks.

### 34.2.1 Apply Gnielinski Entrance Effect Correction

**Scope:** Heat Transfer Correlation Coefficient

Apply Gnielinski Entrance Effect Correction [ Maximum Correction {=} MaxCorrection ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>MaxCorrection</td>
<td>real</td>
<td>10</td>
</tr>
</tbody>
</table>

Summary
Enable the entrance effect correction factor for Gnielinski pipe and annulus correlation heat transfer coefficients (types 72-74).

Description
Applies a correction factor of the form $\left(1 + \frac{d_H^3}{L^3}\right)$ to the heat transfer coefficient where $d_H$ is the hydraulic diameter and $L$ is the distance from the pipe or annulus entrance to the point the flux is applied. By default the maximum value of the correction is 10, it can optionally be set using the MaxCorrection parameter in this command line. For $L < L_{min}$ the correction factor is not applied. This correction factor may only be applied when the correlation is being used with an advective bar.
34.2.2 Characteristic Length
Scope: Heat Transfer Correlation Coefficient

Characteristic Length \( = \) Dh

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dh</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specifies the characteristic length for parameter computations such as Reynolds Number, etc.

34.2.3 Compute
Scope: Heat Transfer Correlation Coefficient

Compute ComputeCorrParam [ Model \( = \) Value... ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>ComputeCorrParam</td>
<td>{annulus diameter ratio</td>
<td>characteristic length</td>
</tr>
</tbody>
</table>

Summary: Specifies that model parameter be computed through such means as extraction, generic form, specific model, etc. Current options for the parameters and associated models include

CHARACTERISTIC LENGTH: \{Hydraulic Diameter | Wetted Perimeter\}

FLUID VELOCITY: This should always be set to the name of the advection velocity used in the energy equation. This will most likely be \textit{advection velocity} which is the internal variable used in Aria but may be set manually to anything else such as a user defined velocity field.

FRICTION FACTOR: \{Smooth Tube\}

34.2.4 Disable Gnielinski Film Gradient Correction
Scope: Heat Transfer Correlation Coefficient

Summary: Forces Aria to neglect the film temperature gradient correction factor when computing the Gnielinski pipe and annulus correlation heat transfer coefficients (types 72-74). This option is generally not recommended, and is provided only for backwards compatibility with the early implementation of these correlations.

34.2.5 Density
Scope: Heat Transfer Correlation Coefficient

Density \( = \) rho
Parameter | Value | Default
---|---|---
rho | real | undefined

**Summary**
Specify a density to either compute or specify by value for usage with a correlation convection coefficient.

### 34.2.6 Entrance Length

**Scope:** Heat Transfer Correlation Coefficient

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>De</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specify an entrance length for usage with a correlation convection coefficient.

### 34.2.7 Expansion Coefficient

**Scope:** Heat Transfer Correlation Coefficient

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>beta</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specify an expansion coefficient by value for usage with a correlation convection coefficient.

### 34.2.8 Fluid Phase

**Scope:** Heat Transfer Correlation Coefficient

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>FluidPhase</td>
<td>{gas, liquid}</td>
<td>GAS</td>
</tr>
</tbody>
</table>

**Summary**
Specify the fluid phase for usage with a correlation convection coefficient.

### 34.2.9 Friction Factor

**Scope:** Heat Transfer Correlation Coefficient

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>f</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specify a friction factor to either compute or specify by value for usage with a correlation convection coefficient.
### 34.2.10 Gnielinski Film Gradient Exponent

**Scope:** Heat Transfer Correlation Coefficient

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exponent</td>
<td>real</td>
<td>0.45</td>
</tr>
</tbody>
</table>

**Summary** Set the exponent used for calculating the film temperature gradient correction for Gnielinski pipe and annulus correlation heat transfer coefficients (types 72-74).

**Description** For gas phase convection the Gnielinski heat transfer coefficient correlations include a correction factor of the form \((\frac{T}{T_w})^n\). By default \(n = 0.45\) is used, corresponding to air. This option lets the user change the exponent to model different gases.

### 34.2.11 Gravitational Constant

**Scope:** Heat Transfer Correlation Coefficient

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>g</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary** Specify a gravitational constant by value for usage with a correlation convection coefficient.

### 34.2.12 Kinematic Viscosity

**Scope:** Heat Transfer Correlation Coefficient

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>nu</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary** Specify a kinematic viscosity to either compute or specify by value for usage with a correlation convection coefficient.

### 34.2.13 Laminar Correlation

**Scope:** Heat Transfer Correlation Coefficient

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>CorrelationType</td>
<td>({1</td>
<td>10</td>
</tr>
</tbody>
</table>
Summary Specifications the laminar correlation model.

34.2.14 Longitudinal Pitch
Scope: Heat Transfer Correlation Coefficient

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$Sl$</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specify a longitudinal pitch to either compute or specify by value for usage with a correlation convection coefficient.

34.2.15 Porosity
Scope: Heat Transfer Correlation Coefficient

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\phi$</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specify a Porosity to either compute or specify by value for usage with a correlation convection coefficient.

34.2.16 Prandtl Number
Scope: Heat Transfer Correlation Coefficient

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$Pr$</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specify a Prandtl number to either compute or specify by value for usage with a correlation convection coefficient.

34.2.17 Reynolds Number
Scope: Heat Transfer Correlation Coefficient

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$Re$</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specify a Reynold’s number to either compute or specify by value for usage with a correlation convection coefficient.
### 34.2.18 Turbulent Correlation

**Scope:** Heat Transfer Correlation Coefficient

**Turbulent Correlation** `{=|are|is} Type CorrelationType

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>CorrelationType</td>
<td>{1</td>
<td>10</td>
</tr>
</tbody>
</table>

**Summary** Specifies the turbulent correlation model.

### 34.2.19 Temperature

**Scope:** Heat Transfer Correlation Coefficient

**Temperature** `{=|are|is} t

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>t</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary** Specify a Fluid Temperature to either compute or specify by value for usage with a correlation convection coefficient.

### 34.2.20 Transition Reynolds Number

**Scope:** Heat Transfer Correlation Coefficient

**Transition Reynolds Number** `{=|are|is} Rex

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rex</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary** Specify a transition Reynolds number to either compute or specify by value for usage with a correlation convection coefficient.

### 34.2.21 Transverse Pitch

**Scope:** Heat Transfer Correlation Coefficient

**Transverse Pitch** `{=|are|is} St

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>St</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary** Specify a transverse pitch to either compute or specify by value for usage with a correlation convection coefficient.
34.2.22 Velocity
Scope: Heat Transfer Correlation Coefficient

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$v$</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify a velocity to either compute or specify by value for usage with a correlation convection coefficient.

34.2.23 Wall Angle
Scope: Heat Transfer Correlation Coefficient

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\theta$</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify a wall angle by value for usage with a correlation convection coefficient.

34.2.24 Wall Length
Scope: Heat Transfer Correlation Coefficient

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$l$</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify a wall length by value for usage with a correlation convection coefficient.

34.2.25 Wall Prandtl Number
Scope: Heat Transfer Correlation Coefficient

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$Pr_w$</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Specify a wall Prandtl number to either compute or specify by value for usage with a correlation convection coefficient.

34.2.26 Wall Temperature
Scope: Heat Transfer Correlation Coefficient
Wall Temperature \( T_w \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>( T_w )</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  
Specify a Wall Temperature to either compute or specify by value for usage with a correlation convection coefficient.

**34.2.27 Use Correlation Convection Model**

**Scope:**

| Use Correlation Convection Model Name  
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  
Specifies correlation model for convection coefficient
Chapter 35

Chemistry Overview

Aria provides two ways of modeling chemical contributions to the transport equations it solves: CHEMEQ and General Chemistry. The details of these two approaches are described in the following chapters, and the purpose of this overview is to describe the differences between these approaches and provide guidelines for which one to select for a given application.

35.1 Feature Comparison

As a quick reference, a list of features supported by both solvers is shown in Table 35.1 below.

As the table illustrates, CHEMEQ is best suited for evaluating reactions without species transport in order to obtain a source term for the energy equation. It also links to pressurization zones (described in a later chapter) to enable reduced-cost pressurization calculations. CHEMEQ stores its species concentrations in an element field and updates that element field in time while providing a source term for the energy equation.

On the other hand, General Chemistry is best suited for situations where you want to resolve species transport and reaction. General Chemistry is designed only to provide source terms, so there must be a linear system set up for each DOF (energy, species concentrations, etc...) to actually update its values in time. For cases where the species are not transported, the added cost of setting up this linear system makes CHEMEQ the more efficient choice. One advantage of General Chemistry is the updated reaction input format which allows you to mix different types of reactions in the same mechanism, which is not possible in CHEMEQ unless you implement it yourself in a user subroutine.
<table>
<thead>
<tr>
<th>Feature</th>
<th>CHEMEQ</th>
<th>General Chemistry</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arrhenius Reactions</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Distributed Arrhenius Reactions</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Pressure Dependent Reactions</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>User Subroutine Reactions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Time Dependent Reactions</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>General Form Reactions</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Prout-Tompkins Reactions</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Can Mix Reaction Types</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Can be Solved with Species Transport</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Can be Solved Without Transport</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Porous System Reactions</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Activation Temperature</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Deactivation Temperature</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Activation Time</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Deactivation Time</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>General ODE solver interface</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Arbitrary Concentration Units</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Can link to pressurization zones</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Segregated solve</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Coupled solve</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Provides Energy Source</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Provides Species Sources</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Reversible Reactions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Equilibrium Chemistry</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Table 35.1.** Feature table for CHEMEQ and General Chemistry.
Chapter 36

CHEMEQ Reference

36.1 Chemical Heating

Materials undergoing non–diffusive endothermic or exothermic chemical reactions can be modeled in Aria. The effect of such reactions is incorporated into the energy conservation equation as a volumetric heating term, which is calculated from the reaction rates. In this section, we give a brief overview of the equations implemented in Aria\textsuperscript{1}.

The volumetric heating due to the reactions may be written as

\[ \dot{q}(x, t, T) = \sum_{j=1}^{N_r} R_j r_j, \]  

(36.1)

where \( R_j \) is the known endothermic or exothermic energy release for step \( j \) of the reaction, \( r_j \) is the calculated reaction rate for the same step, and the summation over the index \( j \) is performed for the \( N_r \) reaction steps.

Some of the burden of managing the units properly for this expression fall to the user. As described in a subsequent section, you can specify energy release units in the ChemEq block as either per volume or per unit mass. The units of \( \dot{q} \) are always power per unit volume (e.g. W/m\(^3\)). When you select the “per unit mass” option, the right hand side of the equation above is multiplied by the local density. For example, if the concentration units are mass fractions, you should use the “per unit mass” option. If the concentration units are mass densities, you should use the “per unit volume” option.

Step \( j \) of a multi–step chemical reaction can be expressed using the stoichiometric equation

\[ \sum_{i=1}^{N_s} \nu'_{ij} M_i \rightarrow \sum_{i=1}^{N_s} \nu''_{ij} M_i, \]  

(36.2)

where \( M_i \) is the chemical symbol for species \( i \), \( \nu'_{ij} \) is the stoichiometric coefficient of the reactant species \( i \) in reaction step \( j \), \( \nu''_{ij} \) is the stoichiometric coefficient of the product species \( i \) in reaction step \( j \), and \( N_s \) represents the number of chemical species. When a species does not occur as a reactant or product in equation (36.2) for a particular reaction step, the corresponding stoichiometric coefficient is set to zero.

The net rate of change in the concentration of the species is determined from the following ordinary differential equation:

\[ \frac{dN_i}{dt} = \sum_{j=1}^{N_r} (\nu''_{ij} - \nu'_{ij}) r_j, \]  

(36.3)

where \( N_i \) is the concentration (or mole fraction) for species \( i \). In equations (36.1) and (36.3), the reaction rates are calculated according to the law of mass action

\[ r_j = k_j \prod_{i=1}^{N_s} N_i^{\nu'_{ij}} \quad \text{for} \quad j = 1, 2, \ldots, N_r, \]  

(36.4)

\textsuperscript{1}Heat transfer in the presence of chemical reactions is detailed in Glassman’s Combustion [39].
where $k_j$ is the kinetic coefficient for reaction step $j$, and $\mu_{ij}$ is the given concentration exponent for reaction step $j$ and species $i$. The kinetic coefficient, $k_j$, for each reaction step is usually determined from the Arrhenius equation, which may be written as

$$k_j = M_j A_j \exp \left( \frac{-E_j}{RT} \right) \left( \frac{p}{p_0} \right)^{\alpha_j} T^{\beta_j},$$  \hfill (36.5)

where $A_j$ is the pre-exponential factor, $M_j$ is the rate multiplier, $E_j$ is the activation energy, $R$ is the appropriate universal gas constant, $T$ is the temperature, $\beta_j$ is the steric coefficient, $p$ is the pressure, $p_0$ is a reference pressure and $\alpha_j$ is the pressure exponent factor. Upon substitution of (36.5) and (36.4) into (36.3), we obtain a nonlinear system of ordinary differential equations for the evolution of the chemical species concentrations, namely

$$\frac{dN_i}{dt} = \sum_{j=1}^{N_r} \left[ \left( \nu''_{ij} - \nu'_{ij} \right) A_j M_j \exp \left( \frac{-E_j}{RT} \right) \left( \frac{p}{p_0} \right)^{\alpha_j} T^{\beta_j} \prod_{k=1}^{N_s} N_k^{\mu_{kj}} \right]$$  \hfill (36.6)

Equation (36.6), with appropriate initial conditions for the species concentrations, must be solved simultaneously with equation (5.9) to obtain the time evolution of the temperature field and species concentrations. This is a difficult coupled system of nonlinear equations to solve, because the time scales associated with the chemical reactions can be very different from the time scale associated with conduction. Furthermore, the time scale associated with one chemical reaction step may be very different from the next. An operator splitting strategy is used to decouple the concentration equations from the energy equation at every time step.

Finally, the thermal properties of the chemical material are regarded as weighted averages of the $N_s$ values associated with each species component. Another weighted average that is sometimes useful when interpreting results is the reacted gas fraction, $f_{RG}$, which is defined as the fraction of reacting material that exists in gas phase,

$$f_{RG} = \frac{(1 - X_c) \sum_{i=1}^{N_s} N_i g_i}{\sum_{i=1}^{N_s} N_i}$$  \hfill (36.7)

where $X_c$ represents the condensed fraction for the reactive material, and the parameter $g_i$ is defined as

$$g_i = \begin{cases} 1 & \text{if } N_i \text{ is a gas phase species} \\ 0 & \text{if } N_i \text{ is not a gas phase species} \end{cases}$$  \hfill (36.8)

### 36.2 Alternate Reaction Models

The default reaction model, Arrhenius, was described in the previous section. Alternate reaction models can be selected for the ChemEq block, and are described in this section.

#### 36.2.1 Prout-Tompkins Reaction Model

The Prout-Tompkins reaction model is used for a reaction with only one product and one reactant. The concentration of the reactant is noted as $\alpha$, and must be a normalized quantity (typically mass fraction). The three additional required parameters for the model are $m$, $n$, and $p$. The rate of this reaction is then evaluated as:

$$r_0 = A_0 \exp \left( \frac{-E_0}{RT} \right) \alpha^n \left[ 1 - (1 - 10^{-p}) \alpha \right]^m$$  \hfill (36.9)

The values of $m$, $n$, and $p$ are provided to the ChemEq description using a data block.
36.3 Concentration Units

The units of the concentration, $N_i$, used in the formulation in the previous section are up to the user. They could be molar concentrations, mole fractions, mass fractions, or mass concentrations (densities). For the purpose of evaluating the reactions described in the previous section, this choice does not matter. However, if you are using a pressurization model in conjunction with the ChemEq model, you have to specify which unit system your concentrations are in so the moles of gas produced can be calculated correctly.

You can also extract the concentrations of individual species from ChemEq to expressions, to allow further manipulations like concentration-dependent thermal properties. To use this capability, the units for the ChemEq block must be specified. There is no implicit unit conversion done in this process, so if the ChemEq units are mass fractions, you can only extract them to the mass fraction expressions. Any further conversions required can be done after they are extracted from ChemEq.

36.4 Numerical Solution

Numerical simulations including chemical heating require a coupling of subcycled transient solutions for Equation (36.6) and the energy equation (5.10). Given appropriate initial conditions for the species concentrations and temperatures at each element quadrature point, the strategy employed here is to first solve Equation (36.6) for updated species concentrations and an associated volumetric heating rate. Here solving for values at quadrature points skirts any issues with chemistry averaging for adjacent material blocks. Quadrature point volume heating rates are spatially integrated and applied as a source contribution (5.11) to the energy equation which can then be solved for an updated temperature distribution.

In addition to solving for the chemical chemistry state and heating rates at quadrature points, the CHEMEQ interface returns a characteristic timestep used in its subcycled iteration. Since chemistry is subcycled within the FEM model step this characteristic time step will always be substantially smaller than the time step used for outer iteration. Nevertheless, the chemistry time time step must be considered in the outer iteration time step selection if one is to accurately model the system chemistry. The strategy used here is to enable a user specified scaling of the minimum chemistry time step (via CHEMISTRY STEP MULTIPLIER command) in the time step selection process for the FEM model time step. Selection of the chemistry time step is illustrated schematically in Figure 36.1 below. Note that reduction of the CHEMISTRY STEP MULTIPLIER parameter will often cause the overall minimum chemistry time step to dictate the outer iteration time step.

The numerical model for modeling chemical heating is divided into two sections. One contains a detailed description of the chemical system and the other specifies details of how the set of ordinary differential equations (36.6) are to be solved. Command line input for these two aspects of the numerical model are described in the sections which follow.

The detailed chemical system command line descriptions are provided as an input command block embedded within a higher-level Aria Material command block described in Material Properties (4). It is important to note that chemical heating constitutes a volumetric source term contribution to the energy equation, hence the source must be invoked using a command line as described in Source Terms (11). In most cases chemical heating is thermally activated above some threshold temperature and is controlled independently for each element block to which it is applied. Additional controls for activation and deactivation of the chemical system heat source are provided in the CHEMEQ solver section. The activity status for the source on any element block is available for output as a global variable, $block\_number\_is\_on$. 991
Figure 36.1. Chemistry Time Step Selection

36.5 Parameters For Chemeq Model

Scope: Material Phase

Begin Parameters For Chemeq Model ModelName

Activation Energies {=|are|is} activation energies...
Activation Energy St Devs {=|are|is} activation energy st devs...
Aux Variable Names {=|are|is} aux variable names...
Aux Variable Subroutine {=|are|is} aux variable subroutine
Concentration Exponents For SpeciesName {=|are|is} Exponents...
Concentration Units {=|are|is} concentration units...
Condensed Fraction {=|are|is} condensed fraction
Energy Release Units {=|are|is} energy release units
Energy Releases {=|are|is} energy releases...
Enthalpies Of Formation {=|are|is} enthalpies of formation...
Extent Of Reaction Based On extent of reaction based on...
Log Preexponential Factors {=|are|is} log preexponential factors...
Number Of Reactions {=|are|is} number of reactions
Pressure {=|are|is} pressure...
Pressure Exponents {=|are|is} pressure exponents...
Rate Multiplier {=|are|is} rate multiplier...

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Reaction Rate Model {=} reaction rate model
Reaction Rate Subroutine {=} reaction rate subroutine
Reference Pressure {=} reference pressure
Species Molecular Weights {=} species molecular weights...
Species Names {=} species names...
Species Phases {=} species phases...
Steric Coefficients {=} steric coefficients...
Stoichiometric Coefficients For SpeciesName {=} Exponents...

Summary  Parameters for the CHEM EQ chemistry model within Aria. The ModelName must appear in a SRC term for an ENERGY equation in order to be applied.

36.5.1 Activation Energies
Scope: Parameters For Chemeq Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>activation energies</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  This command specifies the activation energies for each reaction.

36.5.2 Activation Energy St Devs
Scope: Parameters For Chemeq Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>activation energy st devs</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  This command supplies the standard deviations for the activation energies for each reaction. The order is significant, and corresponds to the order in which the Reactions are specified.

36.5.3 Aux Variable Names
Scope: Parameters For Chemeq Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>aux variable names</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Supplies the names for any auxiliary variables. The number of auxiliary variables is implied by the size of this list.
The entries must be separated by whitespace, without commas. These variables do not participate in the reaction kinetics, but are derived from the species variables and calculated in the user-defined subroutine named via the "AUX VARIABLE SUBROUTINE NAME" line command.

### 36.5.4 Aux Variable Subroutine

**Scope:** Parameters For Chemeq Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>aux variable subroutine</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Supplies the name of the user-defined subroutine that performs any auxiliary variable calculations. This command is required if auxiliary variables are named.

### 36.5.5 Concentration Exponents For

**Scope:** Parameters For Chemeq Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>SpeciesName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Exponents</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

For the given species, this command specifies the concentration exponents for each reaction.

### 36.5.6 Concentration Units

**Scope:** Parameters For Chemeq Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>concentration units</td>
<td>string...</td>
<td>UNKNOWN_UNITS</td>
</tr>
</tbody>
</table>

**Summary**

Specifies the concentration units. This input is only required when using a pressurization model. Valid inputs include: "Mass Fraction", "Mass Fractions", "Mole Fraction", "Mole Fractions", "Mass", "Density", "Moles", or "Molar"

### 36.5.7 Condensed Fraction

**Scope:** Parameters For Chemeq Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>condensed fraction</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

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36.5.8 Energy Release Units
Scope: Parameters For Chemeq Model

Summary Specifies the energy release units for each reaction, PER VOLUME or PER UNIT MASS. Default is PER VOLUME. If PER UNIT MASS is selected, the specified heat release value is multiplied by the local density.

36.5.9 Energy Releases
Scope: Parameters For Chemeq Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>energy releases</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary This command specifies the energy release for each reaction.

36.5.10 Enthalpies Of Formation
Scope: Parameters For Chemeq Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>enthalpies of formation</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary This command specifies the enthalpies of formation for each species.

36.5.11 Extent Of Reaction Based On
Scope: Parameters For Chemeq Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>extent of reaction based on</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary This command supplies the species on which the progress variable for each distributed activation energy depends. The order is significant, and corresponds to the order in which the Reactions are specified.

36.5.12 Log Preexponential Factors
Scope: Parameters For Chemeq Model
Log Preexponential Factors \(\log p\)reexponential factors...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\log p)reexponential factors</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  This command specifies the natural logarithms of the pre-exponential factors for each reaction.

### 36.5.13 Number Of Reactions

**Scope:** Parameters For Chemeq Model

Number Of Reactions \(\text{number of reactions}\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>number of reactions</td>
<td>integer</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Specifies the number of reactions for this material.

### 36.5.14 Pressure

**Scope:** Parameters For Chemeq Model

Pressure \(\text{pressure}\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>pressure</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Specifies the expression for pressure for this material.

### 36.5.15 Pressure Exponents

**Scope:** Parameters For Chemeq Model

Pressure Exponents \(\text{pressure exponents}\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>pressure exponents</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  This command specifies the pressure exponents for each reaction.

### 36.5.16 Rate Multiplier

**Scope:** Parameters For Chemeq Model

Rate Multiplier \(\text{rate multiplier}\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>rate multiplier</td>
<td>string...</td>
<td>1</td>
</tr>
</tbody>
</table>
36.5.17 Reaction Rate Model

Scope: Parameters For Chemeq Model

Summary
Specifies a generic expression for the rate multiplier for this ChemEq model. This input is optional, and the rate multiplier defaults to 1.0 if this is omitted.

36.5.18 Reaction Rate Subroutine

Scope: Parameters For Chemeq Model

Summary
By default, reaction rates are Arrhenius. If this line command is included, then the named subroutine is called instead.

36.5.19 Reference Pressure

Scope: Parameters For Chemeq Model

Summary
Specifies the reference pressure for this material.

36.5.20 Species Molecular Weights

Scope: Parameters For Chemeq Model

Summary
Supplies the molecular weight for each chemical species. The order is significant, and corresponds to the order in which the Species Names are specified.
36.5.21 Species Names
Scope: Parameters For Chemeq Model

Species Names {=|are|is} species names...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>species names</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Supplies names for the chemical species. The number of chemical species is implied by the size of this list. The entries must be separated by whitespace, without commas.

36.5.22 Species Phases
Scope: Parameters For Chemeq Model

Summary Supplies the phase for each chemical species. The order is significant, and corresponds to the order in which the Species Names are specified.

36.5.23 Steric Coefficients
Scope: Parameters For Chemeq Model

Steric Coefficients {=|are|is} steric coefficients...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>steric coefficients</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary This command specifies the steric coefficients for each reaction.

36.5.24 Stoichiometric Coefficients For
Scope: Parameters For Chemeq Model

Stoichiometric Coefficients For SpeciesName {=|are|is} Exponents...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>SpeciesName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Exponents</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary For the given species, this command specifies the stoichiometric coefficients for each reaction.

36.6 Chemeq Solver For
Scope: Equation System

Begin Chemeq Solver For ModelName

Absolute Tolerance {=|are|is} absTol
Parameters for the CHMEQ solver applicable to the previously defined material with CHMEQ model ModelName. The CHMEQ SOLVER model produces no independent action since the ModelName must first appear in a SRC term for an ENERGY equation before it be applied.

### 36.6.1 Absolute Tolerance

**Scope:** Chemeq Solver For

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>absTol</td>
<td>real</td>
<td>1e-12</td>
</tr>
</tbody>
</table>

**Summary**

Specifies the absolute solution tolerance for the ODE solver

### 36.6.2 Activation Temperature

**Scope:** Chemeq Solver For

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>T_act</td>
<td>real</td>
<td>0.0</td>
</tr>
</tbody>
</table>

999
Summary Specification of threshold activation temperature for chemistry.

### 36.6.3 Activation Temperature Rate

**Scope:** Chemeq Solver For

**Activation Temperature Rate** $\equiv \{\text{are|is}\} T_{\cdot dot}$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T_{\cdot dot}$</td>
<td>real</td>
<td>REAL_MAX</td>
</tr>
</tbody>
</table>

Summary Specification of temperature rate limit ($T_{\cdot dot}$) for which chemistry activation occurs.

### 36.6.4 Activation Time

**Scope:** Chemeq Solver For

**Activation Time** $\equiv \{\text{are|is}\} T_{\cdot init}$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T_{\cdot init}$</td>
<td>real</td>
<td>REAL_MAX</td>
</tr>
</tbody>
</table>

Summary Specification of activation time for chemistry.

### 36.6.5 Allow Old Heat Generation Method

**Scope:** Chemeq Solver For

**Allow Old Heat Generation Method** $\equiv \{\text{are|is}\} Bool$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bool</td>
<td>{false</td>
<td>true}</td>
</tr>
</tbody>
</table>

Summary Specifies whether to allow the use of the old heat generation method, which is known to not satisfy energy conservation.

### 36.6.6 Asymptotic Tolerance

**Scope:** Chemeq Solver For

**Asymptotic Tolerance** $\equiv \{\text{are|is}\} Tol$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tol</td>
<td>real</td>
<td>100.0</td>
</tr>
</tbody>
</table>

Summary Specifies the asymptotic selection criterion for the chemical rate equations.
36.6.7 Aux Variable
Scope: Chemeq Solver For

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>C_init</td>
<td>real</td>
<td>0.0</td>
</tr>
</tbody>
</table>

Summary Specifies the initial concentration for the named auxiliary variable.

36.6.8 Chemistry Step Multiplier
Scope: Chemeq Solver For

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dt_cond_over_dt_chem</td>
<td>real</td>
<td>100.0</td>
</tr>
</tbody>
</table>

Summary This command specifies the ratio of the maximum allowed conduction timestep to the smallest, current chemistry time step.

36.6.9 Deactivation Temperature
Scope: Chemeq Solver For

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>t</td>
<td>real</td>
<td>REAL_MAX</td>
</tr>
<tr>
<td>ChemistryDeactivationAction</td>
<td>{continue</td>
<td>terminate}</td>
</tr>
</tbody>
</table>

Summary Specification of deactivation temperature and how the simulation should proceed after deactivation.

36.6.10 Deactivation Temperature Overshoot
Scope: Chemeq Solver For

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>dT_overshoot</td>
<td>real</td>
<td>1.0</td>
</tr>
</tbody>
</table>

Summary Specification of acceptable overshoot of the deactivation temperature for the limiting procedure
36.6.11 Deactivation Temperature Rate
Scope: Chemeq Solver For

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>T_dot</td>
<td>real</td>
<td>REAL_MAX</td>
</tr>
<tr>
<td>ChemistryDeactivationAction</td>
<td>{continue</td>
<td>terminate}</td>
</tr>
</tbody>
</table>

Summary: Specification of temperature rate limit (T_dot) for deactivation and how the simulation should proceed after deactivation.

36.6.12 Deactivation Time
Scope: Chemeq Solver For

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>T_stop</td>
<td>real</td>
<td>REAL_MAX</td>
</tr>
<tr>
<td>ChemistryDeactivationAction</td>
<td>{continue</td>
<td>terminate}</td>
</tr>
</tbody>
</table>

Summary: Specification of deactivation time for chemistry and how the simulation should proceed after deactivation.

36.6.13 Epsilon Max
Scope: Chemeq Solver For

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>E_max</td>
<td>real</td>
<td>10.0</td>
</tr>
</tbody>
</table>

Summary: Specifies the upper bound convergence criterion and chemistry time scale check.

36.6.14 Epsilon Min
Scope: Chemeq Solver For

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>E_min</td>
<td>real</td>
<td>0.0001</td>
</tr>
</tbody>
</table>

Summary: Specifies the lower bound convergence criterion and chemistry time scale check.

36.6.15 Maximum Substeps
Scope: Chemeq Solver For
Maximum Substeps \(\text{maxSubSteps}\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>maxSubSteps</td>
<td>integer</td>
<td>100000</td>
</tr>
</tbody>
</table>

Summary: Specifies the maximum number of chemistry substeps per global time step.

### 36.6.16 Minimum Chemistry Timestep

Scope: Chemeq Solver For

Minimum Chemistry Timestep \(\text{Dt}_\text{min}\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dt_min</td>
<td>real</td>
<td>1.0e-15</td>
</tr>
</tbody>
</table>

Summary: Specifies the minimum time step allowed during the solution of the chemical rate equations.

### 36.6.17 Minimum Concentration For

Scope: Chemeq Solver For

Minimum Concentration For \(\text{Name}\) \(\text{C}_\text{min}\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>C_min</td>
<td>real</td>
<td>1.0e-12</td>
</tr>
</tbody>
</table>

Summary: Specifies the minimum concentration for the named species. This comes with a caveat for transient calculations with long simulation times. The internal implementation within CHEMEQ will always limit the minimum species concentration which implies that a non-zero value for a reactant will always result in species production. This has implications for mass fraction conservation as these errors eventually grow with long simulation times. Reduction of this value helps to reduce the error but this results in a longer run times because the reaction rate equations become stiffer and this results in a smaller chemistry time step being selected. For a typical foam decomposition simulation, this default value is acceptable and for an explosives simulation, the value may be increased to 1.0e-8.

### 36.6.18 Ode Solver

Scope: Chemeq Solver For

Ode Solver \(\text{odeSolver...}\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>odeSolver</td>
<td>string</td>
<td>CVODE</td>
</tr>
</tbody>
</table>

Summary: This command selects the ODE solver to use for a segregated solve. For available options, refer to the 'ODE Solvers' section in the 'Chemistry Solver Reference' chapter of the user manual.
36.6.19 Percentage Asymptotics
Scope:  Chemeq Solver For

Percentage Asymptotics \( \propto | \text{are|is} \) \( \text{PercentValue} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>PercentValue</td>
<td>real</td>
<td>0.0</td>
</tr>
</tbody>
</table>

Summary  Specifies the percentage of the chemical rate equations that will always be solved using asymptotics.

36.6.20 Relative Tolerance
Scope:  Chemeq Solver For

Relative Tolerance \( \propto | \text{are|is} \) \( \text{relTol} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>relTol</td>
<td>real</td>
<td>1e-6</td>
</tr>
</tbody>
</table>

Summary  Specifies the relative solution tolerance for the ODE solver.

36.6.21 Species
Scope:  Chemeq Solver For

Species \( \propto | \text{are|is} \) \( \text{Name} \) \( \text{Conc} \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Conc</td>
<td>real</td>
<td>0.0</td>
</tr>
</tbody>
</table>

Summary  Specifies the initial concentration for the named species.
Chapter 37

General Chemistry Reference

37.1 Introduction

Aria provides a general capability to specify arbitrary chemical mechanisms of irreversible reactions as part of a simulation. Both homogeneous (single phase) and volumetric heterogeneous (multiple material phases) mechanisms are currently supported. The mechanism is defined by the user in the material block in the input deck, and this definition will then provide species, mass, and energy source terms that can be inserted into the various transport equations being solved.

It may be more convenient to specify the mechanism in either a mole-based unit system or a mass-based unit system, so both capabilities are provided. The formulation of both versions of this functionality are described in the following sections.

Reactions can be specified in two ways in a general chemistry block. The first way uses a format similar to CHEMEQ, where all reactions have the same form, and you enter blocks of coefficients. These inputs are described in the following section. The second method is to enter individual reaction blocks, which is described in the Reaction Inputs section later in this chapter. These reaction blocks are placed inside the general chemistry block.

An example input using the traditional input style is

```
BEGIN general chemistry gas_chem
Species Names = A B
Number of Reactions = 1
Species Variable Name = density
Universal Gas Constant = 1.0
Preexponential Factors = 0.1
Steric Coefficients = 0.0
Activation Energies = 350.0
Concentration Exponents for A = 1.0
Concentration Exponents for B = 0.0
Stoichiometric Coefficients for A = -1.0
Stoichiometric Coefficients for B = 1.0
Heats of Reaction = -1e6
END
```

and the same mechanism input using reaction blocks would be

```
BEGIN general chemistry gas_chem
Species Names = A B
Species Variable Name = density
```
These two input methods can also be combined, but the specified number of reactions on the "Number of Reactions" line should be only the number listed in the traditional format, not including reactions inside "Begin Reaction" blocks.

### 37.2 Required Prerequisites

Using general chemistry requires that several expressions be defined in the same material block or available on the block it is evaluated on. These expressions, and the conditions when they are required, are described in the following list.

- Specific heat of each species (only required if you request a chemistry source term for the energy equation)
- Enthalpy of each species (only required if you request a chemistry source term for the energy equation, if you use an evaluator for specific heat this is created automatically)
- Temperature
- Pressure (optional, if you do not provide it and use a pressure dependent reaction rate it will use 101325 Pa)
- Concentrations (required, you specify what form of concentration, e.g. mass fraction, density, species, or species fraction)
- Molecular weights of each species (required if you use molar concentrations)

The phases for inputs (temperature and pressure) come from NO_MATERIAL_PHASE by default. You can specify a list of temperature phases using the old input syntax to override this default. The pressure, when using the old input syntax always comes from the same phase as temperature. When using the new reaction block format, these can be specified separately for each reaction.

### 37.3 General Chemistry

**Scope:** Material Phase

```
Begin General Chemistry ModelName
Species Names {=|are|is} species names...
Species Variable Name {=|are|is} species variable name...
Species Phases {=|are|is} species phases...
Number Of Reactions {=|are|is} number of reactions
Universal Gas Constant {=|are|is} universal gas constant
```
Summary

Parameters for the General Chemistry model within Aria. The ModelName must appear in a SRC term for an ENERGY or MASS equation in order to be applied.

37.3.1 Activation Energies
Scope: General Chemistry

Activation Energies \( \{=|\text{are}|\text{is}\} \text{ activation energies} \ldots \\

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>activation energies</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary

This command specifies the activation energies for each reaction.

37.3.2 Concentration Exponents For
Scope: General Chemistry

Concentration Exponents For \( \text{SpeciesName} \) \( \{=|\text{are}|\text{is}\} \text{ exponents} \ldots \\

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>SpeciesName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Exponents</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary

For the given species, this command specifies the concentration exponents for each reaction.

37.3.3 Enthalpy of Formation For
Scope: General Chemistry

Enthalpy of Formation For \( \text{SpeciesName} \) \( \{=|\text{are}|\text{is}\} \text{ enthalpy of formation} \ldots \\

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>SpeciesName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Enthalpy of Formation</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>
Summary
For the given species, this command specifies the enthalpy of formation. This is the preferred method of determining heat release. If this is omitted and the heat of reaction is provided instead, the code attempts to determine a set of formation enthalpies that satisfy the heats of reaction. If this cannot be done, an error is thrown.

37.3.4 Species Variable Name
Scope: General Chemistry

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>species variable name</td>
<td>string</td>
<td>None</td>
</tr>
</tbody>
</table>

Summary
Specifies the species variable name. Options currently include "Density" (for mass-based calculations) or "Species" (for molar based calculations).

37.3.5 Heats of Reaction
Scope: General Chemistry

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>heats of reaction...</td>
<td>real...</td>
<td>0.0</td>
</tr>
</tbody>
</table>

Summary
This command specifies the heat of reaction for each reaction. If the enthalpies of formation are provided, this section is ignored. Otherwise, these are converted to enthalpies of formation as described later in this chapter.

37.3.6 Preexponential Factors
Scope: General Chemistry

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>preexponential factors...</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary
This command specifies the pre-exponential factors for each reaction.

37.3.7 Universal Gas Constant
Scope: General Chemistry

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>universal gas constant</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>
37.3.8 Nucleation Time

Scope: General Chemistry

| Nucleation Time \{=|\ are\|\ is\} nucleation time |
|------------------|----------------------|-------------------|
| Parameter        | Value        | Default          |
| nucleation time  | real         | undefined        |

Summary
This is an optional input that specifies a nucleation time to apply to the reactions. If provided, it is applied to all the traditionally specified reactions, but not to reactions entered in their own Reaction block.

37.3.9 Number Of Reactions

Scope: General Chemistry

| Number Of Reactions \{=|\ are\|\ is\} number of reactions |
|-------------------|----------------------|-------------------|
| Parameter         | Value      | Default          |
| number of reactions| integer   | undefined        |

Summary
Specifies the number of reactions for this material, not including reactions entered in their own Reaction block.

37.3.10 Species Names

Scope: General Chemistry

| Species Names \{=|\ are\|\ is\} species names... |
|-------------------|----------------------|-------------------|
| Parameter         | Value     | Default          |
| species names     | string... | undefined        |

Summary
Supplies names for the chemical species. The number of chemical species is implied by the size of this list. The entries must be separated by whitespace, without commas.

37.3.11 Species Phases

Scope: General Chemistry

| Species Phases \{=|\ are\|\ is\} species phases... |
|------------------|----------------------|-------------------|
| Parameter        | Value         | Default          |
| Species Phases   | string...     | undefined        |

Summary
Supplies the phase for each chemical species. The order is significant, and corresponds to the order in which the Species Names are specified.
37.3.12 Temperature Phases

Scope: General Chemistry

Temperature Phases {are} temperature phases...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature Phases</td>
<td>string</td>
<td>NO_MATERIAL_PHASE</td>
</tr>
</tbody>
</table>

Summary Supplies the phase for each reaction gets its temperature. This does not apply to reactions entered in Reaction blocks.

37.3.13 Steric Coefficients

Scope: General Chemistry

Steric Coefficients {are} steric coefficients...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>steric coefficients</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary This command specifies the steric coefficients for each reaction.

37.3.14 Stoichiometric Coefficients For

Scope: General Chemistry

Stoichiometric Coefficients For SpeciesName {are} Exponents...

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>SpeciesName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Exponents</td>
<td>real...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary For the given species, this command specifies the stoichiometric coefficients for each reaction.

37.4 Molar Formulation of Chemical Mechanism

37.4.1 Species Transport

Consider a multi-step chemical reaction mechanism comprised of \( j \) irreversible reactions. This system can be expressed as

\[
\sum_{i=1}^{N_s} \tilde{v}''_{ij} M_i \rightarrow \sum_{i=1}^{N_s} \tilde{v}''_{ij} M_i, \tag{37.1}
\]

where \( \tilde{v}''_{ij} \) is the molar stoichiometric coefficient for reactant species \( M_i \) and \( \tilde{v}''_{ij} \) is the molar stoichiometric coefficient for product species \( M_i \), and \( N_s \) is the number of chemical species. When a species does not appear in either the reactants or products for a reaction \( j \), its stoichiometric coefficients are set to zero.
The net rate of change of molar concentration of species $M_i$, summed over all $j$ reactions in the mechanism is then
\[ \frac{d[M_i]}{dt} = \sum_{j=1}^{N_r} (\tilde{v}'_{ij} - \tilde{v}'_{ij}) \tilde{r}_j, \]  
(37.2)
where $N_r$ is the number of reactions and $\tilde{r}_j$ is the molar reaction rate for reaction $j$, calculated according to the law of mass action,
\[ \tilde{r}_j = \tilde{k}_j \prod_{i=1}^{N_r} [M_i]^\mu_{ij} \]  
(37.3)
where $\mu_{ij}$ is the concentration exponent for species $i$ in reaction $j$ and $\tilde{k}_j$ is the kinetic coefficient for reaction $j$, modeled using the Arrhenius form
\[ \tilde{k}_j = \tilde{A}_j T^{\beta_j} \exp \left( -\frac{E_j}{R_u T} \right) \]  
(37.4)
where $\tilde{A}_j$ is the pre-exponential factor, $\beta_j$ is the steric coefficient, $E_j$ is the activation energy, $R_u$ is the universal gas constant, and $T$ is the temperature. Combining all terms, the final expression for the molar production rate of species $i$ is then
\[ \frac{d[M_i]}{dt} = \tilde{\omega}_i = \sum_{j=1}^{N_r} \left( (\tilde{v}'_{ij} - \tilde{v}'_{ij}) \tilde{A}_j T^{\beta_j} \exp \left( -\frac{E_j}{R_u T} \right) \prod_{k=1}^{N_r} [M_k]^\mu_{ik} \right). \]  
(37.5)

It is entirely possible that the simulation might be configured such that the native species solution variable might not be the molar concentration $[M_i]$, but instead might be something like mole fraction, species density, or mass fraction. In these cases, it is up to the user to provide a set of material models to convert whatever the solution variable is into something dimensionally equivalent to molar concentration, and then to provide the name of the final converted molar concentration variable to the chemistry configuration block in the Aria input deck. Some useful conversion into molar concentration include
\[ [M_i] = \rho X_i \frac{W}{W_i} = \rho Y_i \frac{W}{W_i} = \rho_i, \]  
(37.6)
where $X_i$ is the mole fraction of species $i$, $Y_i$ is the mass fraction of species $i$, $\rho_i$ is the density of species $i$, $W_i$ is the molecular weight of species $i$, and $W$ is the mixture molecular weight computed as $W = \sum X_i W_i = (\sum Y_i/W_i)^{-1}$.

If the reaction mechanism is specified on a molar basis, then the result of the calculations will be a volumetric mole production rate. If this is to be used by a mass-based species transport equation, then an appropriate source term model should be specified by the user for the species transport equations, that include the requisite unit conversions.

### 37.4.2 Continuity Transport

Once the individual species source terms have been formulated, the overall continuity equation source term for material phase $k$ can then be computed as a summation over all of the individual species source terms as
\[ \frac{d[M_k]}{dt} = \tilde{\omega}_{c,k} = \sum_{i=1}^{N_s,k} \frac{d[M_i]}{dt} \]  
(37.7)
where $[M_k]$ is the overall molar concentration in material phase $k$ and the summation is over the $N_{s,k}$ species in material phase $k$. The volumetric material phase is intended to represent the different states of the material in a single volume, such as can be found in porous flow where there may be a solid skeleton phase, a gas phase, and/or a liquid phase occupying the pores. If there is only a single material phase (homogeneous combustion), then this continuity source term will be zero; it can only be non-zero when mass is being exchanged between material phases.
### 37.4.3 Energy Transport

The volumetric heating source term presents an interesting complexity, since the form of the source term itself varies depending on the type of energy equation being solved. For example, a user might want to solve an enthalpy equation, a sensible enthalpy equation, or even a temperature equation. The molar enthalpy for a particular species $i$ is defined as

$$\bar{h}_i = \Delta \bar{h}^o_{f,i} + \int_{T_{\text{ref}}}^T \bar{c}_{p,i}(T) \,dT$$  \hspace{1cm} (37.8)

$$= \Delta \bar{h}^o_{f,i} + \bar{h}_{s,i},$$  \hspace{1cm} (37.9)

where $\Delta \bar{h}^o_{f,i}$ is the molar heat of formation for species $i$ at the reference temperature $T_{\text{ref}}$, $\bar{c}_{p,i}$ is the molar specific heat of species $i$, and the molar sensible enthalpy $\bar{h}_{s,i}$ is defined as the integral of the specific heat from the reference state up to the temperature of interest. The mixture molar enthalpy may be calculated by

$$\bar{h} = \bar{h}_s + \sum_{i=1}^{N_s} \Delta \bar{h}^o_{f,i} X_i,$$  \hspace{1cm} (37.10)

where the mixture molar sensible enthalpy is defined as

$$\bar{h}_s = \sum_{i=1}^{N_s} \bar{h}_{s,i} X_i.$$  \hspace{1cm} (37.11)

For homogeneous combustion, an enthalpy equation requires no combustion source term at all, but the enthalpy must be known for each individual species, including both the chemical and sensible portions. For poorly-characterized species, the heat of formation might be unknown or harder to measure (or approximate) than the specific heat. In these cases, a sensible enthalpy equation might be more convenient.

The energy source term due to combustion when not using total enthalpy becomes

$$\bar{S}_{h_s} = - \sum_{i=1}^{N_s} \Delta \bar{h}^o_{f,i} \dot{\omega}_i$$  \hspace{1cm} (37.12)

$$= - \sum_{i=1}^{N_s} \Delta \bar{h}^o_{f,i} \left( \sum_{j=1}^{N_r} (\bar{v}'_{ij} - \bar{v}_{ij}', \bar{r}_j) \right)$$  \hspace{1cm} (37.13)

$$= - \sum_{j=1}^{N_r} \left( \sum_{i=1}^{N_s} \Delta \bar{h}^o_{f,i} (\bar{v}'_{ij} - \bar{v}_{ij}') \right) \bar{r}_j$$  \hspace{1cm} (37.14)

$$= - \sum_{j=1}^{N_r} \Delta \bar{h}_{R,j} \bar{r}_j,$$  \hspace{1cm} (37.15)

where $\Delta \bar{h}_{R,j}$ is the molar heat of reaction for reaction $j$, which is potentially easier to measure than the heat of formation for individual species. When specifying the reaction system, either the heat of reaction or the enthalpy of formation can be provided. However, if a set of non-physical heats of reaction is provided, an error will be generated.

Extra care must be taken in the case of volumetric heterogeneous combustion, since mass is conserved globally but not within each individual material phase since mass is converted from one phase to another (e.g. from a porous solid into a gas). In this case, if separate enthalpy transport equations are solved for each
material phase, a source term appears in each equation that accounts for the enthalpy addition or removal due to creation or destruction of mass within that phase.

Each reaction has a designated phase in which it occurs. The temperature of its origin phase is used to calculate the reaction rate, and the energy source term is applied to the origin phase. However, if species are transferred from one phase to another by the reaction, they enter the new phase at the temperature of the origin phase and an appropriate energy transfer term is included.

### 37.5 Mass Formulation of Chemical Mechanism

#### 37.5.1 Species Transport

The model form presented in the previous section is on a molar basis, but it may be more convenient to formulate the chemical mechanism on a mass basis. In this case, the multi-step chemical reaction mechanism may be written as

\[
\sum_{i=1}^{N_r} \nu_{ij}' M_i \rightarrow \sum_{i=1}^{N_r} \nu_{ij}'' M_i. \tag{37.16}
\]

Here, the view is taken that the stoichiometric coefficient \(\bar{\nu}_{ij}\) is a molar ratio between species \(i\) and a reference species, \(\bar{\nu}_{ij} = \frac{N_i}{N_{ref}}\), where the reference species is often taken to be one of the reactants that can be identified as the "fuel". Thus, the mass-based stoichiometric coefficients are \(\nu_{ij}' = \bar{\nu}_{ij}' W_{ij} / \bar{W}_i\) and \(\nu_{ij}'' = \bar{\nu}_{ij}'' W_{ij} / \bar{W}_i\), where \(W_i\) is the molecular weight of species \(i\) and \(W_{ref}\) is the molecular weight of the reference species.

The net rate of change of mass concentration (or species density), is then

\[
\frac{d\rho_i}{dt} = \dot{\omega}_i'' = W_i \sum_{j=1}^{N_r} (\bar{\nu}_{ij}'' - \bar{\nu}_{ij}') \bar{r}_j
\]

\[
= W_i \sum_{j=1}^{N_r} \frac{W_{ref}}{W_i} (\nu_{ij}'' - \nu_{ij}') \bar{r}_j
\]

\[
= \sum_{j=1}^{N_r} (\nu_{ij}'' - \nu_{ij}') r_j, \tag{37.17}
\]

where the mass reaction rate is \(r_j = W_{ref} \bar{r}_j\) and the relations in Equation 37.6 are used to convert molar concentration to species density. The mass reaction rate for reaction \(j\) is then specified in terms of mass quantities as

\[
\bar{r}_j = \bar{k}_j W_{ref} \prod_{i=1}^{N_r} \rho_i^{\mu_{ij}} W_i^{\mu_{ij}}
\]

\[
= k_j \prod_{i=1}^{N_r} \rho_i^{\mu_{ij}} \tag{37.18}
\]

where the mass kinetic coefficient is

\[
k_j = \frac{W_{ref}}{\prod_{i=1}^{N_r} W_i^{\mu_{ij}}} A_j T^{\beta_j} \exp \left( - \frac{E_j}{R_a T} \right). \tag{37.19}
\]
or, more compactly,
\[ k_j = A_j T^{\beta_j} \exp \left( -\frac{E_j}{R_u T} \right), \] (37.20)
where the molecular weights can be “absorbed” into the pre-exponential factor to convert it into a mass-based quantity,
\[ A_j = \frac{W_{\text{ref}}}{\prod_{i=1}^{N_s} W_i^{\mu_{ij}}} \bar{A}_j, \] (37.21)

Combining all terms above yields the final expression for the volumetric mass production rate for species \( i \),
\[ \frac{d\rho_i}{dt} = \dot{\omega}_i'' = \sum_{j=1}^{N_r} \left[ (\nu_{ij}' - \nu_{ij}'')A_j T^{\beta_j} \exp \left( -\frac{E_j}{R_u T} \right) \prod_{k=1}^{N_r} \rho_k^{\mu_{kj}} \right], \] (37.22)
which has the same basic functional form as Equation 37.5, meaning that the same model implementation can be used interchangeably for either a mass-based or a mole-based formulation, as long as all user-provided parameters are in an internally-consistent units basis.

The same caveats exist for the mass formulation as the mole formulation, with regard to proper units for the input species variables and the output species reaction rates. Proper material model and source term models must be specified to convert the desired species solution variable into species densities as an input and to convert the volumetric mass production rate into appropriate units for the transport equation source term. Some common conversions into species density include
\[ \rho_i = \rho Y_i = \frac{W_i}{W} \rho X_i = W_i [M_i]. \] (37.23)

### 37.5.2 Continuity Transport

Once the individual species source terms have been formulated, the overall mass-based continuity equation source term for material phase \( k \) can then be computed as a summation over all of the individual species source terms as
\[ \frac{d\rho_k}{dt} = \dot{\omega}_c'' = \sum_{i=1}^{N_{s,k}} \frac{d\rho_i}{dt}, \] (37.24)
where \( \rho_k \) is the overall density of material phase \( k \) and the summation is over the \( N_{s,k} \) species in material phase \( k \). If there is only a single material phase (homogeneous combustion), then this continuity source term will be zero; it can only be non-zero when mass is being exchanged between material phases.

### 37.5.3 Energy Transport

As with the mole-based formulation, the volumetric heating source term takes on a different form depending on the type of equation being solved. For example, a user might want to solve an enthalpy equation, a sensible enthalpy equation, or even a temperature equation. The enthalpy for a particular species \( i \) is defined as
\[ h_i = \Delta h^o_{f,i} + \int_{T_{ref}}^{T} c_{p,i}(T) dT \] (37.25)
\[ = \Delta h^o_{f,i} + h_{s,i}, \] (37.26)
where \( \Delta h^o_{f,i} \) is the mass-based heat of formation for species \( i \) at the reference temperature \( T_{\text{ref}} \), \( c_{p,i} \) is the specific heat of species \( i \), and the sensible enthalpy \( h_{s,i} \) is defined as the integral of the specific heat from
the reference state up to the temperature of interest. The mixture enthalpy may be calculated by
\[
h = h_s + \sum_{i=1}^{N_s} \Delta h_{f,i}^o Y_i, \quad (37.27)
\]
where the mixture sensible enthalpy is defined as
\[
h_s = \sum_{i=1}^{N_s} h_{s,i} Y_i. \quad (37.28)
\]

For homogeneous combustion, an enthalpy equation requires no combustion source term at all, but the
enthalpy must be known for each individual species, including both the chemical and sensible portions. For
poorly-characterized species, the heat of formation might be unknown or harder to measure (or approximate)
than the specific heat. In these cases, a sensible enthalpy equation might be more convenient.

For a sensible enthalpy equation, the source term due to combustion becomes
\[
S_{h_s} = -\sum_{i=1}^{N_s} \Delta h_{f,i}^o \dot{\omega}_{i}''
\]
\[
= -\sum_{i=1}^{N_s} \Delta h_{f,i}^o \left( \sum_{j=1}^{N_r} (\nu_{ij}'' - \nu_{ij}') r_j \right) \quad (37.30)
\]
\[
= -\sum_{j=1}^{N_r} \left( \sum_{i=1}^{N_s} \Delta h_{f,i}^o (\nu_{ij}'' - \nu_{ij}') \right) r_j \quad (37.31)
\]
\[
= -\sum_{j=1}^{N_r} \Delta h_{R,j} r_j, \quad (37.32)
\]
where \( \Delta h_{R,j} \) is the mass-based heat of reaction for reaction \( j \), which is potentially easier to measure or
approximate than the heat of formation for individual species. When specifying the reaction system, either
the heat of reaction or the enthalpy of formation can be provided. However, if a set of non-physical heats of
reaction is provided, an error will be generated.

As with the molar formulation, extra care must be taken in the case of volumetric heterogeneous combus-
tion since mass is conserved globally but not within each individual material phase. In this case, if separate
enthalpy transport equations are solved for each material phase, a source term appears in each equation that
accounts for the enthalpy addition or removal due to creation or destruction of mass within that phase.

If the overall reaction mechanism is either endothermic or exothermic, then it might make physical sense
to deposit that energy into a particular phase (e.g. the solid). It is not clear how best to model this effect,
so initial versions of this capability will not support the enthalpy equation for heterogeneous combustion.
Similar complexities exist for a temperature transport equation with heterogeneous combustion, so it will
also not be supported in initial versions.

Each reaction has a designated phase in which it occurs. The temperature of its origin phase is used to
calculate the reaction rate, and the energy source term is applied to the origin phase. However, if species
are transferred from one phase to another by the reaction, they enter the new phase at the temperature of
the origin phase and an appropriate energy transfer term is included.
37.6 Reaction Inputs

The reactions described above use a common reaction form with an Arrhenius rate and a standard concentration dependence. General chemistry actually uses a generic reaction form, where the reaction rate is expressed as a product of four contributing functions

\[ r_j = f_m(t, T, P, Y) f_r(t, T, P, Y) f_p(P, T) f_c(Y), \]  (37.33)

where \( f_m \) is the multiplier function, \( f_r \) is the rate function, \( f_p \) is the pressure function, and \( f_c \) is the concentration function. The available options for each of these functions are outlined in the following section. They can be selected in the input file by adding as many reaction blocks in the general chemistry block, as

Begin Reaction [REACTION NAME]
  Reaction is A + 1.2B -> 3C
  Multiplier Function = [Optional, See below]
  Rate Function = [Required, See below]
  Pressure Function = [Optional, See below]
  Concentration Function = [Optional, See below]
  Temperature Phase = [GAS_PHASE, SOLID_PHASE, NO_MATERIAL_PHASE (default)]
  Pressure Phase = [GAS_PHASE, SOLID_PHASE, NO_MATERIAL_PHASE (default)]
  Heat of Reaction = [Value]
End

where each reaction in a given general chemistry block must have a unique name. The reaction string is parsed to determine the appropriate stoichiometry for the reaction. The left hand side and right hand side must be separated by either "->" or "=>". There are multiple options for each of the functions, which are described in the following sections.

If the pressure phase argument is omitted but temperature phase is provided, the pressure phase defaults to the same phase as the temperature.

A summary of all the reactions parsed from the input is also printed to the log file so you can verify they are evaluating as you have intended. An example of this output is shown below.

+-----------------------------------------------------------------------------+
| Reaction Summary |
+-----------------------------------------------------------------------------+
| Name: R1 |
| Reaction String: A->B |
| Stoich Coeffs: -1.00000 1.00000 |
| Temperature Phase: No Material Phase |
| Pressure Phase: No Material Phase |
| Heat of Reaction: -1.00000e+06 |
| Multiplier Function: Unity |
| Rate Function: Arrhenius |
| Arguments: A = 0.100000 Ea = 350.000 beta = 0.00000 R = 1.00000 |
| Pressure Function: Unity |
| Concentration Function: Standard |
| Arguments: mu = 1.00000 0.00000 |

If you have entered any reactions in the traditional style, they are also parsed and displayed in this list, but their Reaction String entry will be blank.
37.6.1 Multiplier Functions

Unity

The default multiplier function is unity,

\[ f_m(t, T, P, Y) = 1 \]  

(37.34)

and can be used by either omitting the Multiplier Function line or using

Multiplier Function = Unity

Tanh_Time

The tanh_time multiplier function uses a hyperbolic tangent function of time to increase the reaction rate at a specified time.

\[ f_m(t, T, P, Y) = \frac{1}{2} (1 + \tanh(x_{\text{span}}(t - x_{\text{ref}}))) \]  

(37.35)

The two inputs for this are \( x_{\text{ref}} \) and \( x_{\text{span}} \). If you do not specify an input for the span, it uses \( x_{\text{span}} = 1/x_{\text{ref}} \), unless \( x_{\text{ref}} \) was set as 0, in which case an error is thrown. Example inputs are shown below.

Multiplier Function = Tanh_Time x_ref = 54
Multiplier Function = Tanh_Time x_ref = 2.5 span = 0.1

Tanh_Temperature

The tanh_temperature multiplier function uses a hyperbolic tangent function of temperature to increase the reaction rate at a specified temperature.

\[ f_m(t, T, P, Y) = \frac{1}{2} (1 + \tanh(x_{\text{span}}(T - x_{\text{ref}}))) \]  

(37.36)

The arguments for this function as the same format as for the tanh_time function.

Multiplier Function = Tanh_Temperature x_ref = 450
Multiplier Function = Tanh_Temperature x_ref = 600 span = 10

General

The general multiplier function uses a function string supplied in the input file to evaluate the rate generally. Allowable variables in this string include \( t \) (time), \( T \) (temperature), \( P \) (pressure), and any valid species name from the current model. The value used for a species is in whatever unit system the general chemistry model is using. Powers in this string are denoted using the 'ˆ' character.

An example input for this option is

Multiplier Function = General Function = 0.5*(1+tanh((T-350)^2))
37.6.2 Rate Functions

Unity

The rate function is the only required function, so it has no default like the others. However, you can still select Unity for it if desired, using

Rate Function = Unity

\[ f_r(t, T, P, Y) = 1 \]  \hspace{1cm} (37.37)

Arrhenius

The Arrhenius rate function is the most common, and uses

\[ f_r(t, T, P, Y) = AT^\beta \exp \left( -\frac{E_a}{RT} \right) \]  \hspace{1cm} (37.38)

The pre-exponential constant, \( A \), is the only required input. Optional inputs include \( E_a \), \( \beta \), and \( R \). If omitted, the defaults for these are 0 for \( E_a \) and \( \beta \) and 8.314 for \( R \). Some example lines to select this rate function is

Rate Function = Arrhenius A = 100 Ea = 20000
Rate Function = Arrhenius A = 100 Ea = 350 R = 1
Rate Function = Arrhenius A = 100 beta = 2 Ea = 350 R = 1

Distributed Arrhenius

The Distributed Arrhenius rate function uses a cumulative normal distribution to vary the activation energy with respect to a selected progress species.

\[ f_r(t, T, P, Y) = AT^\beta \exp \left( -\frac{(E_a + \sigma \zeta)}{RT} \right) \]  \hspace{1cm} (37.39)

\[ \zeta = CDF^{-1}(1 - Y_j/Y_0) \]  \hspace{1cm} (37.40)

where \( CDF^{-1} \) is the clipped inverse cumulative normal distribution function, \( Y_j \) is the concentration of the selected progress species, \( Y_0 \) is the specified reference concentration, and \( \sigma \) is the activation energy variation. In addition to these inputs, the same input rules as for the Arrhenius reaction apply (only \( A \) is required of those again). The name specified for the progress species must be a valid species name.

Some example inputs for this rate type are (note that these are shown on multiple lines to fit in the manual, but must be on the same line in your input file):

Rate Function = Distributed_Arrhenius A = 100 Ea = 20000 sigma = 1000 Y0 = 1.2 ProgressSpecies = N2
Rate Function = Distributed_Arrhenius A = 100 beta = 2 Ea = 350 R = 1 sigma = 100 Y0 = 1.2 ProgressSpecies = N2
General

The general rate function uses a function string supplied in the input file to evaluate the rate generally. Allowable variables in this string include \( t \) (time), \( T \) (temperature), \( P \) (pressure), and any valid species name from the current model. The value used for a species is in whatever unit system the general chemistry model is using. Powers in this string are denoted using the ‘\(^\wedge\)’ character.

An example input for this option is:

\[
\text{Rate Function} = \text{General Function} = 100*\exp(-350/T)*\text{Char}^2*N2^3.2*(1-CO2)
\]

### 37.6.3 Pressure Functions

#### Unity

The default pressure function is unity,

\[
f_p(P, T) = 1
\]

(37.41)

and can be used by either omitting the Pressure Function line or using

\[
\text{Pressure Function} = \text{Unity}
\]

#### Exponential

The exponential pressure function is evaluated as

\[
f_p(P, T) = \left(\frac{P}{P_{\text{ref}}}\right)^n,
\]

(37.42)

where \( P_{\text{ref}} \) must be positive. An example input for this option is

\[
\text{Pressure Function} = \text{Exponential} \quad P_{\text{ref}} = 1 \quad n = 1.5
\]

### 37.6.4 Concentration Functions

#### Unity

The default concentration function is unity,

\[
f_c(Y) = 1
\]

(37.43)

and can be used by either omitting the Concentration Function line or using

\[
\text{Concentration Function} = \text{Unity}
\]
Standard

The standard concentration function is evaluated as

\[ f_c(Y) = \prod_{i=1}^{N} Y_i^{\mu_i} \quad (37.44) \]

The values of \( \mu \) can be specified manually using a list of entries for \( \mu \), where you must enter a value of \( \mu \) for each species in the system, not just in this reaction.

Alternately, if your reaction rate only depends on the concentration of the reactants, and the value of \( \mu \) is the same as the stoichiometric coefficient, as is common for gas reactions, you can have the values set automatically from the reaction stoichiometry by using the keyword “Automatic”.

For example, if your reaction is “2A + B -> C” the automatic option would provide \( \mu = 2, 1, 0 \). If you have repeated species as both reactants and products, only the reaction portion of the coefficient is considered. For example, “A + B -> 2A + C” would result in automatic exponents of \( \mu = 1, 1, 0 \).

Concentration Function = Standard \( \mu = 1 \) \( 1 \) \( 0 \) \( 0 \) \( 0 \)
Concentration Function = Standard \( \mu = \) Automatic

Prout Tompkins

The Prout Tompkins concentration function can only be used with a reaction that has a single reactant and a single product, and is evaluated as

\[ f_c(Y) = \alpha^n \left( 1 - (1 - 10^{-p}) \alpha \right)^m \quad (37.45) \]

where \( \alpha \) is the concentration of the reactant. An example input for this model is

Concentration Function = Prout_Tompkins \( n = 1.2 \) \( m = 2.5 \) \( p = 9 \)
Chapter 38

Chemistry Solver Reference

38.1 Introduction

The chemistry solver block specifies how a given chemistry description is solved. The description can currently only be a general chemistry description, however the ODE solvers can now be selected in a ChemEq solver block using the same options.

If you do not provide a chemistry solver block for a general chemistry description, it will use the default algorithm (COUPLED).

38.2 Solution Approaches

38.2.1 Coupled Solve

For a coupled solve, the chemistry description provides an instantaneous production rate for each DOF, calculated using values at state NP1. This rate is used directly as a source in the transport equation. This is the default solution algorithm.

This approach provides strong coupling between solution variables, but can limit the fluid time step prohibitively for stiff chemistry sets.

38.2.2 Segregated Solve

The segregated solution uses a frozen flow field to integrate the reaction ODEs from time N to time NP1 and uses the final values to calculate a source term consistent with the selected time integrator for the transport equations. Inputs to the problem, such as pressure, density, or specific heat, are held constant during this integration. The resulting source term is then provided explicitly in the transport equations.

38.3 ODE Solvers

There are several ODE solvers available that can be selected for both general chemistry and ChemEq. The basic set of input arguments for an ODE solver are given below, where the specific solver arguments are described in the following subsections.

ODE Solver = [SOLVER ARGS]
Absolute Tolerance = 1e-10
38.3.1 CVODE

The CVODE solver is the most flexible of the options available, and often the fastest. It is the only option that allows you to select the method (BDF or ADAMS), the order (max of 5 for BDF and 12 for ADAMS), and the iteration scheme (FUNCTIONAL or NEWTON). For stiff problems, “BDF 5 NEWTON” will be most robust, but is also the most costly. For non-stiff problems, a lower order Adams method, such as “ADAMS 6 FUNCTIONAL” may be better.

The default is “BDF 5 NEWTON” if you only write “ODE Solver = CVODE”. A few valid options are shown below.

ODE Solver = CVODE [METHOD] [ORDER] [ITERATION\_SCHEME]
ODE Solver = CVODE ADAMS 6 FUNCTIONAL
ODE Solver = CVODE BDF 3 NEWTON
ODE Solver = CVODE

38.3.2 LSODE

The LSODE solver can sometimes provide better performance than CVODE for large, stiff problems. It always uses the BDF 5 NEWTON scheme, so there are no extra options.

ODE Solver = LSODE

38.3.3 RK54

The RK54 is a mixed-order error-adaptive Runge-Kutta scheme (5th and 4th order). It is typically the slowest, but may be advantageous for slowly changing non-stiff systems,

ODE Solver = RK54

38.4 Chemistry Solver Parameters For

Scope:  Equation System

Begin Chemistry Solver Parameters For ModelName
  Absolute Tolerance {=|are|is} absTol
  Chemistry Solver Algorithm {=|are|is} algorithmType
  Maximum Substeps {=|are|is} mazSubSteps
  Ode Solver {=|are|is} odeSolver...
  Relative Tolerance {=|are|is} relTol
End
Summary
Parameters for the chemistry solver applicable to the previously defined material with chemistry model ModelName. The chemistry solver model produces no independent action since the ModelName must first appear in a SRC term for an ENERGY or SPECIES equation before it be applied.

38.4.1 Absolute Tolerance
Scope: Chemistry Solver Parameters For

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>absTol</td>
<td>real</td>
<td>1e-12</td>
</tr>
</tbody>
</table>

Summary Specifies the absolute solution tolerance for the ODE solver

38.4.2 Chemistry Solver Algorithm
Scope: Chemistry Solver Parameters For

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>algorithmType</td>
<td>string</td>
<td>COUPLED</td>
</tr>
</tbody>
</table>

Summary This command specifies the coupling algorithm between the chemistry model and transport equations

38.4.3 Maximum Substeps
Scope: Chemistry Solver Parameters For

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>maxSubSteps</td>
<td>integer</td>
<td>100000</td>
</tr>
</tbody>
</table>

Summary Specifies the maximum number of chemistry substeps per global time step.

38.4.4 Ode Solver
Scope: Chemistry Solver Parameters For

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>odeSolver...</td>
<td>string</td>
<td>CVODE</td>
</tr>
</tbody>
</table>
Summary  This command selects the ODE solver to use for a segregated solve. For available options, refer to the ‘ODE Solvers’ section of this chapter.

38.4.5 Relative Tolerance

Scope:  Chemistry Solver Parameters For

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>relTol</td>
<td>real</td>
<td>1e-6</td>
</tr>
</tbody>
</table>

Summary  Specifies the relative solution tolerance for the ODE solver
Chapter 39

Pressurization Model Reference

39.1 Pressurization Zones

Pressure-dependent chemical reactions may produce gas species in a confined space, so the pressure of that confined space must be tracked to resolve the pressure dependency properly. This pressure is often uniform enough that solving the continuity equation is unnecessarily expensive. To accomplish this without continuity, you can specify pressurization zones in the region block in the input file.

For each pressurization zone, you specify a list of source blocks (ChemEq blocks providing gas to that zone) and pressurization blocks (block in which the gas is accumulating) or pressurized bulk nodes (or both). The available gas volume for the zone is the integration of the gas volume fraction over all the pressurization blocks and bulk nodes you have listed. The temperature of the zone is calculated using a weighted average over the pressurization blocks, which you can select as either a volume average, ideal gas average, mass average, or thermal mass average ($\rho C_p$). The blocks do not have to be contiguous, and the source block does not have to be included in the pressurization blocks.

You can also enter an excess volume in the pressurization zone input block, which is added to the total volume of the blocks and bulk nodes for the pressure calculation. The temperature of this excess volume is assumed to be the same as the one calculated for the listed blocks and bulk nodes unless you specify a different temperature. Depending on what temperature averaging method you select, you may also need to provide a weight to use in conjunction with the specified temperature.

For each pressurization zone, a global variable for pressure, temperature, gas volume, and total moles will be created and shown in the log file. These can be output to heartbeat, included in the Exodus file, and used in Encore or user functions along with other global variables.

39.1.1 Temperature Averaging Methods

This section describes the different averaging methods that can be selected to get an effective temperature ($T_{eff}$). In all cases, $\phi$ is the volume fraction of gas. This effective temperature is what is used in the equation of state to calculate the pressure in the pressurization zone.

Volume

When using the volume averaging method, the effective temperature is calculated from

$$T_{eff} = \frac{\int \phi T dV}{\int \phi dV} \quad (39.1)$$
Ideal Gas

When using the ideal gas averaging method, the effective temperature is calculated based on the ideal gas law. Starting with

\[ n_{tot} = \frac{PV_{tot}}{RT_{eff}} = \sum n_i = \sum \frac{PV_i}{RT_i} \]  

(39.2)

By re-arranging these relationships to solve for \( T_{eff} \) and converting to integral form we get

\[ T_{eff} = \frac{\int \phi dV}{\int \phi / T dV} \]  

(39.3)

Density

When using the density (\( \rho \)) averaging method, the effective temperature is calculated from

\[ T_{eff} = \frac{\int \phi \rho T dV}{\int \phi \rho dV} \]  

(39.4)

Thermal Mass

When using the thermal mass (\( \rho C_p \)) averaging method, the effective temperature is calculated from

\[ T_{eff} = \frac{\int \phi \rho C_p T dV}{\int \phi \rho C_p dV} \]  

(39.5)

39.1.2 Equations of State

Given the volume, temperature, and number of moles of gas produced by all sources, an appropriate equation of state is still required to calculate pressure. This is specified in the pressurization model block, and can be one of the options listed below. Any arguments required for the equation of state are given on the same line, after its name.

Ideal Gas

The ideal gas equation of state requires no coefficients and evaluates the pressure as

\[ p = \frac{nRT}{V}. \]  

(39.6)

and is specified by "Equation of State = Ideal_Gas".

BKWS

The BKWS equation of state requires one parameter, the co-volume (\( V_{co} \)). The pressure is then evaluated using:
\[ x = \frac{nkV_{co}}{V\sqrt{T} + 6620} \quad (39.7) \]

\[ Z = 1 + x \exp(0.298x) \quad (39.8) \]

\[ p = \frac{ZnRT}{V}. \quad (39.9) \]

It is specified by "Equation of State = BKWS covol = VALUE".

**Van Der Waal**

The VanDerWaal equation of state requires two parameters, \( a \) and \( b \). The pressure is then evaluated using:

\[ p = \frac{nRT}{V - bn} - \frac{a}{V^2} \quad (39.10) \]

It is specified by "Equation of State = VanDerWaal a = VALUE b = VALUE".

### 39.1.3 Venting and Coupling of Multiple Pressurization Zones

As of version 4.40 Aria has some basic capabilities to model venting of a pressurization zone to the environment, as well as to couple the pressure of multiple pressurization zones to one another. Venting may be modeled by setting "VENTING MODEL = VENTED" in the pressurization model block. This also requires specifying a model for the venting flow rate as a function of the pressure in the zone. At present there are 2 supported models, one that accounts for choked flow and one that does not. Both models take the form:

\[ \dot{m} = K\rho \sqrt{(P - P_{ambient})/\rho} \quad (39.11) \]

where \( K \) is an empirical factor, \( \rho \) is the density of the gas in the pressurization zone, \( P \) is the pressure in the pressurization zone, and \( P_{ambient} \) is the pressure of of the environment the gas is venting to. The choked flow model switches to the form:

\[ \dot{m}_{choked} = K\rho \sqrt{P(1 - \frac{P_{ambient}}{P_{crit}})/\rho} \quad (39.12) \]

once \( P > \frac{P_{crit}}{P_{ambient}} \) where \( \frac{P_{crit}}{P_{ambient}} \) is the user-specified critical pressure ratio. The value of \( K \) can be set using any of the generic material model forms supported by Aria (i.e. constant, polynomial, user function, etc.). A sample vented pressurization zone input deck block is presented below:

```
Begin Pressurization Model pZone1
Initial Pressure = 13
Initial Mixture Molecular Weight = 32
Pressure Unit = psi
Pressurization Source Blocks = block_1
```
Pressurized Blocks = block_1

Venting Model = Vented
Venting Model Property Venting_Volumetric_Flow_Rate = K_FACTOR_WITH_CHOKING
critical_pressure_ratio = 2.
Venting Model Property ambient_pressure = constant value = 13
Venting Model Property K_factor = constant value = 1.e-2

Equation of State = Ideal_Gas

Temperature Averaging = rhoCp
End Pressurization Model pZone1

Multiple pressurization zones may be connected using the same model forms for the flow rate as are available for venting. This can be used to model multiple sealed volumes connected by a restricted flow path or feature that is initially sealed until it reaches a certain temperature or pressure for example. The coupling is turned on using the syntax:

Begin Bulk Node Coupling coupling
Bulk Nodes = pZone1 pZone2
Couple density model = k_factor_flow
Couple species model = k_factor_flow
additional parameter K_factor = user_function name = k_coupling X = time
End

where "pZone1" and "pZone2" are the names of 2 pressurization zones present in the model.

39.2 Pressurization Model
Scope: Aria Region

Begin Pressurization Model ModelName
Equation Of State {=} equationOfState...
Excess Volume {=} excessVolume
Excess Volume Temperature {=} excessVolumeTemperature
Excess Volume Weight {=} excessVolumeWeight
Initial Mixture Molecular Weight {=} initMMW
Initial Pressure {=} initPressure
Pressure Unit {=} pressureUnit
Pressurization Source Blocks {=} sourceBlocks...
Pressurized Blocks {=} pressurizedBlocks...
Pressurized Bulk Nodes {=} pressurizedBulkNodes...
Temperature Averaging {=} temperatureModel
Venting Model {=} ventingModel
Venting Model Property propertyName {=} propertyModelAndParams...
End

1028
Summary  Parameters for the pressurization model

### 39.2.1 Equation Of State

**Scope:** Pressurization Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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</thead>
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<td>undefined</td>
</tr>
</tbody>
</table>

**Summary** Specifies the equation of state to use (Ideal_Gas, VanDerWaal, or BKWS)

### 39.2.2 Excess Volume

**Scope:** Pressurization Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>excessVolume</td>
<td>real</td>
<td>0.0</td>
</tr>
</tbody>
</table>

**Summary** Specifies the volume of the excess volume. This volume is assumed to be at the average zone temperature unless you also specify an excess volume temperature.

### 39.2.3 Excess Volume Temperature

**Scope:** Pressurization Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>excessVolumeTemperature</td>
<td>real</td>
<td>Copied</td>
</tr>
</tbody>
</table>

**Summary** Specifies the temperature of the excess volume. If omitted, the excess volume is treated as the same temperature as the main averaged temperature. If provided, you must also provide a weighting factor to use for RHO and RHO_CP temperature averaging schemes.

### 39.2.4 Excess Volume Weight

**Scope:** Pressurization Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>excessVolumeWeight</td>
<td>real</td>
<td>1</td>
</tr>
</tbody>
</table>

1029
Summary Specifies the weighting factor to use for the excess volume. For RHO averaging, this will be the density. For rhoCp averaging, this will be rho times Cp. For volume and ideal gas averaging this should not be provided.

39.2.5 Initial Mixture Molecular Weight
Scope: Pressurization Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>initMMW</td>
<td>real</td>
<td>0.02987</td>
</tr>
</tbody>
</table>

Summary Specifies the mixture molecular weight of the initial gas present in the pressurization zone

Description Default is 0.02987 kg/mol for dry air. This should be specified in the appropriate units for the simulation. If no boundary conditions (e.g. venting or coupling to a gas transport region of the model) are applied to the pressurization zone then the value does not affect the results of the simulation.

39.2.6 Initial Pressure
Scope: Pressurization Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>initPressure</td>
<td>real</td>
<td>-1.0</td>
</tr>
</tbody>
</table>

Summary Specifies the initial pressure in the specified pressure units.

39.2.7 Pressure Unit
Scope: Pressurization Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>pressureUnit</td>
<td>string</td>
<td>Pa</td>
</tr>
</tbody>
</table>

Summary This command specifies the units of pressure that is returned. Options are PA, ATM, and PSI (Default is Pa)

39.2.8 Pressurization Source Blocks
Scope: Pressurization Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>sourceBlocks</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>
A list of ChemEq blocks that provide the gas source(s) for this pressurization model.

### 39.2.9 Pressurized Blocks

**Scope:** Pressurization Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>pressurizedBlocks</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: A list of blocks that the pressurization occurs in.

### 39.2.10 Pressurized Bulk Nodes

**Scope:** Pressurization Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>pressurizedBulkNodes</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: A list of bulk nodes that the pressurization occurs in.

### 39.2.11 Temperature Averaging

**Scope:** Pressurization Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>ventingModel</td>
<td>string</td>
<td>IDEAL_GAS</td>
</tr>
</tbody>
</table>

Summary: This command specifies the temperature averaging model to use (VOLUME, IDEAL_GAS, RHO, or RHOCP)

### 39.2.12 Venting Model

**Scope:** Pressurization Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>ventingModel</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: This command specifies the venting model to use (OPEN, CLOSED, or VENTED)
39.2.13 Venting Model Property

Scope: Pressurization Model

### Venting Model Property

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>propertyName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>propertyModelAndParams</td>
<td>string...</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**

Define a material property associated with the venting model. Required if the venting model is VENTED. At a minimum a model for the VENTING_VOLUMETRIC_FLOW_RATE must be provided, and depending on the choice of that model additional properties may be required.
Chapter 40

Solution Options

40.1 Solution Options

Scope: Aria Region

---

Begin Solution Options OptionsName

Assemble Tpetra {=|are|is} Value
Force Non Tale {=|are|is} Value
Free Stream Reynolds Number {=|are|is} input_Re
Ignore Coordinate Displacements {=|are|is} Value
Maximum Temperature Allowed From Temperature Extraction {=|are|is} MaxTemp
Minimum Temperature Allowed From Temperature Extraction {=|are|is} MinTemp
Omit Enthalpy Adjustment After Temperature Clipping
Omit Finite Difference Sensitivities For Cantera Properties
Use Inverse Density Continuity Scaling {=|are|is} Value
Begin Cvfem Algorithm Specification PStabName
End

Begin Edc Model Specification EdcSpecName
End

Begin Hdiff Model Specification HdiffOptionsName
End

Begin Porous Flow Options blockName
End

Begin Turbulence Model Specification TurbSpecName
End

---

Summary

Specify information regarding the governing equations to be solved.

40.1.1 Assemble Tpetra

Scope: Solution Options

---

Summary

EXPERIMENTAL: Enable (true) to assemble to TPETRA and compare matrix and RHS to FEI matrix.
40.1.2  Force Non Tale
Scope:  Solution Options
Summary  Enable (true) to force simulation not to use TALE.

40.1.3  Free Stream Reynolds Number
Scope:  Solution Options

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>input_Re</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary  Input Reynolds number is used for solving nondimensional viscous problems using the GasDyn equations.

40.1.4  Ignore Coordinate Displacements
Scope:  Solution Options
Summary  Enable (true) or disable (false) displacing the physical coordinates by either MESH_DISPLACEMENTS or SOLID_DISPLACEMENTS. By default, this is false, and physical_coordinates are displaced. However, this can be disabled by setting this to true, and physical_coordinates will be equal to model_coordinates.

40.1.5  Maximum Temperature Allowed From Temperature Extraction
Scope:  Solution Options
Summary  Specify a maximum cutoff temperature for extraction from enthalpy and composition

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>MaxTemp</td>
<td>real</td>
<td>2400.0 K</td>
</tr>
</tbody>
</table>

Description  This option specifies the maximum temperature to be allowed to be extracted from enthalpy, given the mixture composition. If a temperature is computed that is greater than this value, the temperature is reset to equal the maximum allowed value.

40.1.6  Minimum Temperature Allowed From Temperature Extraction
Scope:  Solution Options

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>MinTemp</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

1034
Summary: Specify a minimum cutoff temperature for extraction from enthalpy and composition.

Description: This option specifies the minimum temperature to be allowed to be extracted from enthalpy, given the mixture composition. If a temperature is computed that is less than this value, the temperature is reset to equal the minimum allowed value.

40.1.7 Omit Enthalpy Adjustment After Temperature Clipping
Scope: Solution Options

Summary: Do not adjust enthalpy to be consistent with clipped temperatures.

Description: When temperature extraction fails to produce a viable result, the temperature will usually be clipped at either the upper or lower limit temperature. By default, the enthalpy will then be adjusted up or down to match the clipped temperature so that the thermochemical state is internally consistent. This omits the enthalpy adjustment, leaving clipped nodes in an inconsistent state.

40.1.8 Omit Finite Difference Sensitivities For Cantera Properties
Scope: Solution Options

Summary: Do not populate any Cantera property sensitivities via finite difference.

40.1.9 Use Inverse Density Continuity Scaling
Scope: Solution Options

Summary: Enable (true) or disable (false) the inverse element scaling of the continuity equation.

40.2 Cvfem Algorithm Specification
Scope: Solution Options

Begin Cvfem Algorithm Specification $PStabName$

Activate Acoustic Compressibility Algorithm
Activate Ausm Plus Scheme
Activate Edge Based Diffusion Operator [ With CorrectionType ]
Activate Edge Based Fourth Order Advection
Activate Kexact Muscl Scheme For Equation EquationString [ {of|species} SpeciesName |{in|material_phase} MaterialPhaseName ][ Using K {=}|are|is} kappa ]

1035
Activate Mixture Fraction Clipping Utility At \textit{UtilClipLocation}

Activate Muscl Scheme For Equation \textit{EquationString} [ \{of\} \textit{species} \textit{SpeciesName} | \{in\} \textit{material\_phase} \textit{MaterialPhaseName} ]

Activate Pressure Projection Algorithm

Activate Projected Stress Stabilization

Activate Scv Nodal Gradient

Activate Shakib Scaling

Activate Vrtm In Mass Flux Vector

Ausm Alpha \{=|are|is\} \textit{alpha}

Ausm Beta \{=|are|is\} \textit{beta}

Clip Cvfem Level Set Dof

First Order Upwind Factor \{=|are|is\} \textit{RealValue} For Equation \textit{EquationString} [ \{of\} \textit{species} \textit{SpeciesName} | \{in\} \textit{material\_phase} \textit{MaterialPhaseName} ]

Flux Scheme \{=|are|is\} \textit{FluxScheme}

Freeze Muscl Limiter At Global Step \{=|are|is\} \textit{freezeStep}

Hybrid Upwind Factor \{=|are|is\} \textit{RealValue} For Equation \textit{EquationString} [ \{of\} \textit{species} \textit{SpeciesName} | \{in\} \textit{material\_phase} \textit{MaterialPhaseName} ]

Interpolate Density And Velocity Separately In Mass Flux Vector

Lag Nodal Pressure Gradient In Mass Flux Vector Expression

Lag Nodal Tau In Mass Flux Vector Expression

Muscl Limiter \{=|are|is\} \textit{MusclLimiter} For Equation \textit{EquationString} [ \{of\} \textit{species} \textit{SpeciesName} | \{in\} \textit{material\_phase} \textit{MaterialPhaseName} ]

Omit Diffusion Term From Sucv Tau

Omit Disting Bc Ip Sensitivities

Omit Dt Term From Sucv Tau

Pressure Stabilization Characteristic Length \{=|are|is\} \textit{CharLength}

Pressure Stabilization Order \{=|are|is\} \textit{Cufempstaborder}

Pressure Stabilization Parameter Scaling \{=|are|is\} \textit{TauScaling}

Pressure Stabilization Scaling \{=|are|is\} \textit{CufemPStabScaling} [ With Value \{=|are|is\} \textit{ConstantTau} ]

Roe Entropy Fix C \{=|are|is\} \textit{epsc}

Roe Entropy Fix U \{=|are|is\} \textit{epsu}

Upwind Method \{=|are|is\} \textit{UpwMethod} For Equation \textit{EquationString} [ \{of\} \textit{species} \textit{SpeciesName} | \{in\} \textit{material\_phase} \textit{MaterialPhaseName} ]

Use Approximate Roe Sensitivities

Use Opposing Subface In Open Mass Flux

Use Specified Pressure In Open Mass Flux

\textbf{End}

\textbf{Summary} Specify CVFEM algorithmic modeling options.
40.2.1 Activate Acoustic Compressibility Algorithm
Scope: Cvfem Algorithm Specification

Summary Variable thermodynamic pressure to allow for closed system pressurization

40.2.2 Activate Ausm Plus Scheme
Scope: Cvfem Algorithm Specification

Summary Activate AUSM plus with standard values for alpha and beta

Description Activate AUSM plus scheme.

40.2.3 Activate Edge Based Diffusion Operator
Scope: Cvfem Algorithm Specification

Summary Activate edge-based scheme for diffusion operator.

Description Only implemented now for tke and sdr - interior only

40.2.4 Activate Edge Based Fourth Order Advection
Scope: Cvfem Algorithm Specification

Summary Activate edge-based advection for fourth order LES scheme

Description Activate edge-based advection for fourth order LES scheme

40.2.5 Activate Kexact Muscl Scheme For Equation
Scope: Cvfem Algorithm Specification

---

**Activate Kexact Muscl Scheme For Equation**

**EquationString** [ {of|species} **SpeciesName** |{in |material_phase} **MaterialPhaseName** ][ Using K {=|are|is} **kappa** ]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>EquationString</strong></td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td><strong>SpeciesName</strong></td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td><strong>MaterialPhaseName</strong></td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Use kexact MUSCL variable extrapolation for the specified equation

Description Use kexact MUSCL variable extrapolation for the specified equation. The projected gradients for the necessary variables must be computed using a cvfem_lumped_muscl_projection equation.
40.2.6 Activate Mixture Fraction Clipping Utility At
Scope: Cvfem Algorithm Specification

Summary: Activate clipping utility for mixture fraction

Description: Sometimes it helps to clip the mixture fraction. This is a utility that provides this support.

40.2.7 Activate Muscl Scheme For Equation
Scope: Cvfem Algorithm Specification

```
Activate Muscl Scheme For Equation EquationString [ {of|species} SpeciesName |{in|material_phase} MaterialPhaseName ]
```

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>EquationString</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>SpeciesName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>MaterialPhaseName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Use MUSCL variable extrapolation for the specified equation

Description: Use MUSCL variable extrapolation for the specified equation. The projected gradients for the necessary variables must be computed using a cvfem_lumped_muscl_projection equation.

40.2.8 Activate Pressure Projection Algorithm
Scope: Cvfem Algorithm Specification

Summary: Activate PP alg.

Description: Rather than fully coupling uvwp, activate a pressure projection algorithm. This will require creating two EQ systems: uvw and p. However, once this is created, the code will handle supplemental utilities. This option can only be used in the context of a momentum lumped mass matrix.

40.2.9 Activate Projected Stress Stabilization
Scope: Cvfem Algorithm Specification

Summary: Activate full momentum stress term for residual stabilization

Description: Activate full momentum stress term for residual stabilization
40.2.10  **Activate Scv Nodal Gradient**  
**Scope:** Cvfem Algorithm Specification

**Summary**  Use scv nodal gradient for MUSCL and edge-based diffusion terms

**Description**  Use scv nodal gradient for MUSCL and edge-based diffusion terms; default is to use nodal grad based on Green Gauss over the duel mesh with edge based integration points.

40.2.11  **Activate Shakib Scaling**  
**Scope:** Cvfem Algorithm Specification

**Summary**  Specify Shakib scaling; temp line command

40.2.12  **Activate Vrtm In Mass Flux Vector**  
**Scope:** Cvfem Algorithm Specification

**Summary**  Activate velocity relative to mesh in mass flux vector

**Description**  In some cases, the volume may be constant, although there is mesh motion. This is the case in wind energy sliding mesh applications. This algorithm pays not attention to GCL and assumes that the time rate of volume is zero. Again, this is fine in solid rotation.

40.2.13  **Ausm Alpha**  
**Scope:** Cvfem Algorithm Specification

**Parameter**  Ausm Alpha {=|are|is} alpha  
**Value**  
- **alpha**  
  - **real**  
**Default**  0.0

**Summary**  Set constant alpha in Mach splitting function.

**Description**  Default is 0, unless ausm+ is chosen, then the default is 3/16.

40.2.14  **Ausm Beta**  
**Scope:** Cvfem Algorithm Specification

**Parameter**  Ausm Beta {=|are|is} beta  
**Value**  
- **beta**  
  - **real**  
**Default**  0.0

1039
Summary: Set constant beta in Mach splitting function.

Description: Default is 0, unless ausm+ is chosen, then the default is 1/8.

**40.2.15 Clip Cvfem Level Set Dof**

Scope: Cvfem Algorithm Specification

Summary: Clip level set; only germane for conserved level set formulation that has bounds between zero and unity

**40.2.16 First Order Upwind Factor**

Scope: Cvfem Algorithm Specification

Parameter | Value | Default
--- | --- | ---
EquationString | string | undefined
SpeciesName | string | undefined
MaterialPhaseName | string | undefined

Summary: First-order upwind factor, $0 < x \leq 1$.

Description: This value specifies the explicit upwind blending between pure upwind and the chosen convection operator, e.g., $UPW \times (firstOrderUpwind) + (1-firstOrderUpwind) \times (blendedUpwindCentral)$.

where $UPW$ is the pure first order upwind value and $blendedUpwindCentral$ is a blend between the selected upwind method and central difference operator based on the local cell Peclet number (see Hybrid Upwind Factor line command).

For now, the $blendedUpwindCentral$ is pure central.

**40.2.17 Flux Scheme**

Scope: Cvfem Algorithm Specification

Summary: Set flux scheme used by all equations in the high-Mach CVFEM formulation.

Description: Set flux scheme used by all equations in the high-Mach CVFEM formulation.

**40.2.18 Freeze Muscl Limiter At Global Step**

Scope: Cvfem Algorithm Specification

Parameter: freezeStep
Parameter | Value | Default
---|---|---
freezeStep | real | undefined

Summary: Freeze MUSCL limiter at a given global step.

Description: Freeze the MUSCL limiter.

### 40.2.19 Hybrid Upwind Factor

**Scope:** Cvfem Algorithm Specification

**Summary:** Hybrid upwind factor dials in blending between user specified operator and central.

**Description:** Allows blending of the Peclet factor. A value of zero produces pure central. Higher values blend more user specified upwind (currently, pure first order upwind).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>RealValue</td>
<td>real</td>
<td>1.0</td>
</tr>
<tr>
<td>EquationString</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>SpeciesName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>MaterialPhaseName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

### 40.2.20 Interpolate Density And Velocity Separately In Mass Flux Vector

**Scope:** Cvfem Algorithm Specification

**Summary:** use (rho)ip*(uj)ip rather than default (rho*uj)ip

### 40.2.21 Lag Nodal Pressure Gradient In Mass Flux Vector Expression

**Scope:** Cvfem Algorithm Specification

**Summary:** Compute the nodal pressure gradient calculation.

**Description:** Lag the nodal pressure gradient assembly for use in the stabilized mass flux vector expression. This command will place the calculation at the top of the non-linear iteration. In general, it will not be wise to use this option for steady simulations as there is one nonlinear loop.

### 40.2.22 Lag Nodal Tau In Mass Flux Vector Expression

**Scope:** Cvfem Algorithm Specification

**Summary:** Compute the nodal tau calculation.
Lag the nodal tau assembly for use in the stabilized mass flux vector expression. This command will place the calculation at the top of the non-linear iteration. In general, it will not be wise to use this option for steady simulations as there is one nonlinear loop.

### 40.2.23 Muscl Limiter

**Scope:** Cvfem Algorithm Specification

Muscl Limiter `{=|are|is} MusclLimiter For Equation EquationString [ {of|species} SpeciesName |{in|material_phase} MaterialPhaseName ]`

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>MusclLimiter</td>
<td>{muscl_minmod</td>
<td>muscl_none</td>
</tr>
<tr>
<td>EquationString</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>SpeciesName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>MaterialPhaseName</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Specify the MUSCL limiter method.

**Description**
TVD limiter.

### 40.2.24 Omit Diffusion Term From Sucv Tau

**Scope:** Cvfem Algorithm Specification

**Summary**
Do not use timestep in the expression for the SUCV stabilization coefficient

### 40.2.25 Omit Disting Bc Ip Sensitivities

**Scope:** Cvfem Algorithm Specification

**Summary**
Drop distinguishing bc sensitivities for ip values.

**Description**
Drop distinguishing bc sensitivities for ip values.

### 40.2.26 Omit Dt Term From Sucv Tau

**Scope:** Cvfem Algorithm Specification

**Summary**
Do not use timestep in the expression for the SUCV stabilization coefficient

### 40.2.27 Pressure Stabilization Characteristic Length

**Scope:** Cvfem Algorithm Specification

1042
Pressure Stabilization Characteristic Length \(\text{CharLength}\)

<table>
<thead>
<tr>
<th>Parameter (\text{CharLength})</th>
<th>Value (\text{real})</th>
<th>Default undefined</th>
</tr>
</thead>
</table>

**Summary** Specify a constant length scale to compute \(\tau\); only applicable for characteristic scaling.

### 40.2.28 Pressure Stabilization Order

**Scope:** Cvfem Algorithm Specification

Pressure Stabilization Order \(\text{Cvfempstaborder}\)

| Parameter \(\text{Cvfempstaborder}\) | Value \{fourth_order|second_order|zeroth_order\} | Default undefined |
|--------------------------------------|-----------------------------|------------------|

**Summary** Order of pressure stabilization.

### 40.2.29 Pressure Stabilization Parameter Scaling

**Scope:** Cvfem Algorithm Specification

Pressure Stabilization Parameter Scaling \(\text{TauScaling}\)

<table>
<thead>
<tr>
<th>Parameter (\text{TauScaling})</th>
<th>Value (\text{real})</th>
<th>Default undefined</th>
</tr>
</thead>
</table>

**Summary** Scaling of \(\tau\).

### 40.2.30 Pressure Stabilization Scaling

**Scope:** Cvfem Algorithm Specification

Pressure Stabilization Scaling \(\text{CvfemPStabScaling}\) [With Value \{characteristic|constant|shakib|stabilized|time_step\}]

| Parameter \(\text{CvfemPStabScaling}\) | Value \{characteristic|constant|shakib|stabilized|time_step\} | Default undefined |
|---------------------------------------|-----------------------------|------------------|

**Summary** Scaling coefficient for pressure stabilization.

### 40.2.31 Roe Entropy Fix C

**Scope:** Cvfem Algorithm Specification

Roe Entropy Fix C \(\text{epsc}\)

<table>
<thead>
<tr>
<th>Parameter (\text{epsc})</th>
<th>Value (\text{real})</th>
<th>Default 0.1</th>
</tr>
</thead>
</table>

1043
40.2.32 Roe Entropy Fix U
Scope: Cvfem Algorithm Specification

Summary: Small constant for entropy fix (modification of $|U +/ - c|$ eigenvalues in Roe scheme)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>epsu</td>
<td>real</td>
<td>0.1</td>
</tr>
</tbody>
</table>

40.2.33 Upwind Method
Scope: Cvfem Algorithm Specification

Summary: Specify method that is blended with pure second order.

Description: This user defined method will be blended with pure second order based on cell Peclet blending. In most cases, this operator is upwinded, however, in the case of 4th order a higher order centered scheme is used.

40.2.34 Use Approximate Roe Sensitivities
Scope: Cvfem Algorithm Specification

Summary: Use approximate Roe sensitivities for the LHS instead of Van Leer

Description: By default, the Roe flux uses the sensitivities for the Van Leer flux scheme form the LHS matrix. This command turns on a different approximation, based more closely on the Roe flux itself; this approximation converges more quickly for some problems, but is less robust for others. Currently this option only has an effect when the "NC_ADV" equation term is used.

40.2.35 Use Opposing Subface In Open Mass Flux
Scope: Cvfem Algorithm Specification

Summary: Open mass flux will be computed using the interpolated pressure at the opposing subface
40.2.36 Use Specified Pressure In Open Mass Flux
Scope: Cvfem Algorithm Specification

Summary User specified pressure in open BC will be used to compute gradp in open mass flux BC

40.3 Turbulence Model Specification
Scope: Solution Options

Begin Turbulence Model Specification TurbSpecName
Activate Cvfem Lumped Turbulent Source Term Algorithm
Activate Kepsilon Source Term Trickster Linearization
Activate New Komega Src Linearization
Activate Rhs Sdr Sensitivities To TurbSensParams
Activate Rhs Tke Sensitivities To TurbSensParams
Activate Turbulence Clipping Utility At UtilClipLocation
Implicit Les Filter Scale {=|are|is} Value
Limit Turbulent Ke Production To Value Times Dissipation
Omit Finite Difference Sensitivity Due To Utau
Omit Sensitivities In Turbulent Production Term
Omit Sensitivities To Turbulent Production From Ke Source Only
Omit Velocity Divergence In Turbulent Production Term
Time Filter {=|are|is} Value
Turbulence Model {=|are|is} TurbulenceModel
Turbulence Model Parameter TurbParams {=|are|is} Value
End

Summary Specify turbulence modeling options.

40.3.1 Activate Cvfem Lumped Turbulent Source Term Algorithm
Scope: Turbulence Model Specification

Summary Lump cvfem turbulent ke, tdr and ksgs rhs source terms

Description This is appropriate for CVFEM only as it uses a special flavor of lumping source terms.
40.3.2 Activate Kepsilon Source Term Trickster Linearization
Scope: Turbulence Model Specification

Summary
Follow classic Patankar approach to linearization of k rhs

Description
The dissipation rate in the k equation includes sensitivities to k by using the Prandtl Kolmogorov relationship to for the model for of the dissipation rate. This procedure provides additional diagonal dominance for the k rhs source term.

40.3.3 Activate New Komega Src Linearization
Scope: Turbulence Model Specification

Summary
Alternate komega src linearization

Description
Alt komega src linearization

40.3.4 Activate Rhs Sdr Sensitivities To
Scope: Turbulence Model Specification

Activate Rhs Sdr Sensitivities To TurbSensParams

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>TurbSensParams</td>
<td>{dens_sens</td>
<td>sdr_sens</td>
</tr>
</tbody>
</table>

Summary
Dial in rhs sens for sdr equation

40.3.5 Activate Rhs Tke Sensitivities To
Scope: Turbulence Model Specification

Activate Rhs Tke Sensitivities To TurbSensParams

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>TurbSensParams</td>
<td>{dens_sens</td>
<td>sdr_sens</td>
</tr>
</tbody>
</table>

Summary
Dial in rhs sens for tke equation

40.3.6 Activate Turbulence Clipping Utility At
Scope: Turbulence Model Specification

Summary
Activate clipping utility for turbulence quantities

Description
Sometimes it helps to clip the turbulence quantities. This is a utility that provides this support.
### 40.3.7 Implicit Les Filter Scale

**Scope:** Turbulence Model Specification

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Implicit Les Filter Scale</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**
Provide implicit filter length

**Description**
In constant coefficient LES models, allow the user to specify an implicit filter length. The default is to determine a length scale tied to the mesh.

### 40.3.8 Limit Turbulent Ke Production

**Scope:** Turbulence Model Specification

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Limit Turbulent Ke Production To Value Times Dissipation</td>
<td>real</td>
<td>1.0e-8</td>
</tr>
</tbody>
</table>

**Summary**
Choose to limit production source terms

**Description**
This option limits the turbulent ke production to a scale factor of dissipation, prod = min(prod, limit*den*en1). In practice, the ratio of production to dissipation is not very high. In some flows, it is useful to specify a value of approximately 1000. The ratio should be checked as part of the analysis to make sure that violation of the physical ratio has not been done. In general, this option is only activated in domain locations where dissipation rate is very small.

### 40.3.9 Omit Finite Difference Sensitivity Due To Utau

**Scope:** Turbulence Model Specification

**Summary**
Omit FD sensitivity from utau

**Description**
The wall friction velocity, utau, can be complex to evaluate in the context of the law of the wall. The default is to compute sensitivities via finite difference. Sometimes, this seems to add some overhead and spurious sensitivities.

### 40.3.10 Omit Sensitivities In Turbulent Production Term

**Scope:** Turbulence Model Specification

**Summary**
Do not include sensitivities for turbulence production

**Description**
This option removes the analytical sensitivities for the production of turbulent kinetic energy. It seems that most simulations are more stable if this option is activated.
40.3.11 Omit Sensitivities To Turbulent Production From Ke Source Only
Scope: Turbulence Model Specification

Summary Omit sensitivities to turbulence production from the KE source term

Description This option removes the analytical sensitivities for the production of turbulent kinetic energy in the KE source terms (where they may be destabilizing) but retains them elsewhere (where they may be helpful). Mutually exclusive w/ the previous option.

40.3.12 Omit Velocity Divergence In Turbulent Production Term
Scope: Turbulence Model Specification

Summary Do not include divergence term in turbulence production

Description This option removes the divergence term from the turbulent production of kinetic energy. The default is to include this term.

40.3.13 Time Filter
Scope: Turbulence Model Specification

<table>
<thead>
<tr>
<th>Time Filter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter</td>
<td>Value</td>
</tr>
<tr>
<td></td>
<td>real</td>
</tr>
<tr>
<td>Value</td>
<td></td>
</tr>
<tr>
<td>Default</td>
<td>1.0e32</td>
</tr>
</tbody>
</table>

Summary Time filter size for Time Filtered Navier-Stokes model

Description Turbulent viscosity is normally calculated based on a time scale given by k/epsilon (for k-epsilon models) or T (v2f model). The TFNS model substitutes the minimum of the normal computed value and the user-specified time filter size in the turbulent viscosity calculation. In general, this filter should be no less than twice the physical time step. A non-fatal warning is issued if this condition is violated.

40.3.14 Turbulence Model
Scope: Turbulence Model Specification

<table>
<thead>
<tr>
<th>Turbulence Model</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter</td>
<td>TurbulenceModel</td>
</tr>
<tr>
<td>Value</td>
<td>{dsmag</td>
</tr>
<tr>
<td>Default</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary Specify type of turbulence model to be used
40.3.15 Turbulence Model Parameter

Scope: Turbulence Model Specification

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>TurbParams</td>
<td>{a_1</td>
<td>beta</td>
</tr>
<tr>
<td></td>
<td>{c_\epsilon_1</td>
<td>c_\epsilon_2</td>
</tr>
<tr>
<td></td>
<td>{c_\mu_\text{cs}</td>
<td>c_\mu_\epsilon</td>
</tr>
<tr>
<td></td>
<td>{c_\gamma_\text{lam}</td>
<td>c_\gamma_\text{trb}}</td>
</tr>
<tr>
<td></td>
<td>{c_\mu_\epsilon_1</td>
<td>c_\mu_\epsilon_2</td>
</tr>
<tr>
<td></td>
<td>{c_\mu_\epsilon}</td>
<td>undefined</td>
</tr>
<tr>
<td></td>
<td>{edc_c_\gamma_\text{lam}</td>
<td>edc_c_\gamma_\text{trb}}</td>
</tr>
<tr>
<td></td>
<td>{edc_c_\gamma_\text{trb}</td>
<td>edc_c_\text{tau}_\text{lam}}</td>
</tr>
<tr>
<td></td>
<td>{edc_c_\text{tau}_\text{lam}</td>
<td>edc_c_\text{trans}}</td>
</tr>
<tr>
<td></td>
<td>{gamma</td>
<td>gamma_1</td>
</tr>
<tr>
<td></td>
<td>{gamma_1</td>
<td>gamma_2</td>
</tr>
<tr>
<td></td>
<td>{gamma_\star</td>
<td>gamma_\text{lam}}</td>
</tr>
<tr>
<td></td>
<td>{gamma_\text{lam}</td>
<td>gamma_\text{trb}}</td>
</tr>
<tr>
<td></td>
<td>{gamma_\text{trb}</td>
<td>gamma_\text{trans}}</td>
</tr>
<tr>
<td></td>
<td>{gamma_\text{trans}</td>
<td>gamma_\text{trb}}</td>
</tr>
<tr>
<td></td>
<td>{gamma_\text{trb}</td>
<td>gamma_\text{trans}}</td>
</tr>
<tr>
<td></td>
<td>{lr_\omega</td>
<td>lr_\omega_\star</td>
</tr>
<tr>
<td></td>
<td>{lr_\omega_\star</td>
<td>lr_\omega_\text{lam}}</td>
</tr>
<tr>
<td></td>
<td>{lr_\omega_\text{lam}</td>
<td>lr_\omega_\text{trb}}</td>
</tr>
<tr>
<td></td>
<td>{lr_\omega_\text{trb}</td>
<td>lr_\omega_\text{trans}}</td>
</tr>
<tr>
<td></td>
<td>{minimum_\text{turbulent}_\text{viscosity}</td>
<td>prt}</td>
</tr>
<tr>
<td></td>
<td>{minimum_\text{turbulent}_\text{viscosity}</td>
<td>prt}</td>
</tr>
<tr>
<td></td>
<td>{real undefined}</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Turbulence model parameters

40.4 Edc Model Specification

Scope: Solution Options

Begin Edc Model Specification EdcSpecName

Activate Co2 Dissociation Model
Activate Hydrogen Dissociation Model
Activate Ignition Model \{at|for|in|on|over\} IgnPart
Activate Laminar Limit Model
Activate Lumped Source Term Model
Activate Separate Co Irreversible Oxidation Pathway
Activation Time \{are|is\} Time
Fuel Name \{are|is\} Fuel
Ignition Threshold Temperature \{are|is\} IgnTemp
Minimum Product Fraction \{are|is\} Prmin
Reaction Time Scale \{are|is\} Tchem
Reference Property \{are|is\} Value
Reference Property Of Species \{are|is\} Value
Begin Oxidizer Mixture Specification OxMixName
End

Summary: Specify EDC combustion model options.
40.4.1 Activate CO2 Dissociation Model

**Scope:** Edc Model Specification

**Summary** Include effects of CO2 dissociation into CO and O2 at high temperatures

**Description** At high temperatures, the equilibrium between CO2, CO, and O2 shifts away from CO2, which can significantly decrease the flame temperature. Activating this model will add this effect to the standard EDC combustion model.

40.4.2 Activate Hydrogen Dissociation Model

**Scope:** Edc Model Specification

**Summary** Include effects of H2 dissociation into H

**Description** At temperatures greater than about 2000K, the equilibrium between H2 and H will yield non-negligible concentrations of H which can significantly decrease flame temperatures. Activating this model will add this effect to the standard EDC combustion model using the correlations of W.W. Erikson, which are derived from the NASA CEA code [40, 41].

Note that the H species must be included in the Cantera input XML file, and neither H nor H2 should be the "last" species in the list since this species is not independent of the rest (to enforce unity sum) and may be susceptible to more noise than the others. Since temperature and other properties are very sensitive to oscillations in the H and H2 equilibrium, this noise could be problematic.

40.4.3 Activate Ignition Model

**Scope:** Edc Model Specification

---

**Activate Ignition Model** `{@|at|for|in|on|over} IgnPart`

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>IgnPart</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary** MeshPart on which to use the EDC ignition model

**Description** Turn on the ignition model for this MeshPart. If fuel and oxidizer are present and the temperature is below the ignition threshold temperature everywhere in the part, reaction will begin. Volume block names and surface names are valid for this command, as well as the alias "all_blocks" and "all_surfaces". Multiple parts must be specified with multiple instances of this line command.

40.4.4 Activate Laminar Limit Model

**Scope:** Edc Model Specification
Turn on the EDC laminar limit model for low-turbulence situations

This model requires setting three model constants: CtauLam, CgammaLam, and ClamTrans. The model uses a time scale based on a velocity gradient rather than the turb_ke/turb_diss. This appropriate time scale permits the flame to anchor in laminar regions.

40.4.5 Activate Lumped Source Term Model
Scope: Edc Model Specification

Nodally lump the EDC source term

If set, the EDC source terms are nodally lumped, rather than using the default implementation of a consistent approach at the Gauss points.

40.4.6 Activate Separate Co Irreversible Oxidation Pathway
Scope: Edc Model Specification

Add CO oxidation pathway as a separate reaction pathway. This should really be used only in the context of a propellant fire in the presence of hydrogen combustion.

40.4.7 Activation Time
Scope: Edc Model Specification

The time at which the EDC combustion model is activated. No combustion will occur before this time.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time</td>
<td>real</td>
<td>0.0</td>
</tr>
</tbody>
</table>

40.4.8 Fuel Name
Scope: Edc Model Specification

The name of the EDC fuel species, typically either H2 or a major hydrocarbon

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fuel</td>
<td>string</td>
<td>undefined</td>
</tr>
</tbody>
</table>
40.4.9 Ignition Threshold Temperature
Scope: Edc Model Specification

**Ignition Threshold Temperature** \(=|are|is\) \(IgnTemp\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(IgnTemp)</td>
<td>real</td>
<td>1000 K</td>
</tr>
</tbody>
</table>

**Summary**  The temperature below which the ignition model is activated

**Description**  If the ignition model is requested (through the "ACTIVATE IGNITION MODEL" line command, then it will be activated if all temperatures in the corresponding block are below this temperature.

40.4.10 Minimum Product Fraction
Scope: Edc Model Specification

**Minimum Product Fraction** \(=|are|is\) \(Prmin\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Prmin)</td>
<td>real</td>
<td>1.0e-6</td>
</tr>
</tbody>
</table>

**Summary**  The minimum product fraction, below which the EDC model will be deactivated.

40.4.11 Reaction Time Scale
Scope: Edc Model Specification

**Reaction Time Scale** \(=|are|is\) \(Tchem\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Tchem)</td>
<td>real</td>
<td>7.0e-5</td>
</tr>
</tbody>
</table>

**Summary**  Reaction time scale to set extinction

**Description**  Characteristic time scale of the chemical kinetics. Residence time in the fine structure region will be compared to this to determine if extinction will result.

40.4.12 Reference
Scope: Edc Model Specification

**Reference Property** \(=|are|is\) \(Value\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Property)</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>(Value)</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

**Summary**  Reference property for initial diagnostic output
If all required properties are set, the EDC model will print the adiabatic flame temperature and heat of combustion for the mechanism defined by the specified fuel.

### 40.4.13 Reference

**Scope:** Edc Model Specification

| Reference Property Of Species {=|are|is} Value |
|-----------------------------------------------|
| **Parameter** | **Value** | **Default** |
| Property      | string    | undefined   |
| Species       | string    | undefined   |
| Value         | real      | undefined   |

**Summary**
Reference species-dimensioned property for initial diagnostic output

**Description**
If all required properties are set, the EDC model will print the adiabatic flame temperature and heat of combustion for the mechanism defined by the specified fuel.

### 40.5 Oxidizer Mixture Specification

**Scope:** Edc Model Specification

**Begin Oxidizer Mixture Specification OxMixName**

\[
\text{Mass\_Fraction } \text{SpeciesName} \{=|are|is\} \text{ Value}
\]

\[
\text{Mole\_Fraction } \text{SpeciesName} \{=|are|is\} \text{ Value}
\]

**End**

**Summary**
Specify either mass fractions or mole fractions for the oxidizer mixture. Only non-zero species need to be included, and the mass or mole fractions must sum to unity. The default is air, with a 0.2095:0.7905 molar ratio between O2 and N2.

### 40.5.1 Mass\_Fraction

**Scope:** Oxidizer Mixture Specification

| Mass\_Fraction SpeciesName {=|are|is} Value |
|---------------------------------------------|
| **Parameter** | **Value** | **Default** |
| SpeciesName   | string    | undefined   |
| Value         | real      | undefined   |

**Summary**
Oxidizer mixture mass fraction for the given species
40.5.2 Mole_Fraction
Scope: Oxidizer Mixture Specification

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>SpeciesName</td>
<td>string</td>
<td>undefined</td>
</tr>
<tr>
<td>Value</td>
<td>real</td>
<td>undefined</td>
</tr>
</tbody>
</table>

Summary: Oxidizer mixture mole fraction for the given species

40.5.3 Assemble Tpetra
Scope:

Summary: EXPERIMENTAL: Enable (true) to assemble to TPETRA and compare matrix and RHS to FEI matrix.
Chapter 41

Frequently Asked Questions

This chapter includes a compilation of frequently asked questions and problematic scenarios raised by users of the SIERRA Multimechanics Module: Aria. Here the questions are grouped into specific categories of interest but some of the items may appear in more than one category.

- General 41.1
- Capability 41.2
- Errors 41.3
- Solver 41.4
- Thermal 41.5
- Radiation 41.6
- Timestep 41.7
- Contact 41.8
- Species 41.9
- Porous Media 41.10
- Electrostatics 41.11
- Grid Refinement 41.12
- Postprocessing 41.13
41.1 General

Aria example/tutorial files.
SNL internal users can find an Aria training presentation, which includes two PDFs and multiple example files on the following website:
http://compsim.sandia.gov/compsim/
To navigate to the presentations, go to “Support & Services -> Documentation”. Once there, go to “Version of the Day -> General Release”. This will open a new page with an expandable tree. Go to “Training/Tutorials -> Thermal/Fluid”.

Methods for accessing VOTD code.
SNL internal users can access VOTD code on the various platforms by executing module load sierra/master.

Use of the VOTD code for production simulations is generally discouraged as stability of the code is subject to daily change. Whenever possible the analyst is encouraged to use the current released distribution executable.

What is the Aria command to specify a log file name.
If the input file is named base_input.i then by default the log file is written to base_input.log. One can specify an alternative log file by providing optional arguments, either -o other.log or -l other.log to the job execution command line.

What optional arguments can be used with the Aria command.
The list of arguments which can be used with the Aria command can be listed by:

    aria -h

Is the includefile command supported in Aria?
Yes, this feature is supported in all Sierra Mechanics applications. However, the aprepro include command is more robust and supports nested include files, while Sierra’s includefile command does not.

What is the difference between History and Heartbeat output?
The big difference between history and heartbeat output is the type of output. History output is binary format, like Exodus, while Heartbeat output is in ASCII format, like a .txt file. Thus the History output must be visualized with a graphical tool while Heartbeat output is simply inspected or used to generate x-y plots.

Why does restart output a file per processor?
The reason for this is that one typically continues a restart on the same number of processors as the original run. This is also the reason why restart files aren’t concatenated at the end of a successful run. Additionally, unless the intent is to overwrite a restart file, one must specify different names for the input and output restart files in the Restart Data block.

Recently the ability to restart with a different number of processors has been provided but requires that one first concatenate the original files and perform the decomposition on the new processor count.

Can one include a text file in an input deck?
Yes, Aprepro provides a way to do this. The syntax is as follows:

    {include "materials.dat"}

Aprepro must be executed on the input deck before running.
41.2 Capability

Can one use steady-state simulation results to specify initial conditions on a different geometry?
   There are two possible approaches to accomplish this:
   1. using the omit block option with the initial conditions via the IC read file option
   2. Interpolate the simulation results onto the new geometry using transfers from the Input_Output Region.

Can Aria solve a condensation problem?
   There is ongoing work in studies of evaporation, but condensation is not currently supported.

Can one use a FORTRAN user subroutine for heat flux boundary condition?
   Yes one can do this but FORTRAN subroutines are currently available only for heat transfer problems.

What can one model with shell elements in Aria.
   Typical usage of thermal shell elements in Aria is for materials, which have no through-thickness temperature variation but have thermal diffusion in the plane of the shell. The shell can also account for thermal capacitance. Cases where the through thickness resistance of heat flow through the material is more important will generally be modeled with resistance contact (no shell). The in-between case is a combination of the above. Limited support is also available for shells with through-thickness temperature variation but these models are restricted to having only shell elements.

Does Aria support 1D elements, such as bars?
   Aria does support 2D and 3D bar elements however no provision is made for application of flux boundary conditions at the end of the bar. Thus in many cases one may be better off using volume elements instead.

How does one implement a volumetric source on a 2D mesh.
   The fundamental assumption in 2D FEM Cartesian formulations is that of a unit depth cross-section in defining the corresponding volume. Hence, the command add volume block_name will work in both 2D and 3D cases.

Can Aria be run explicitly?
   Explicit methods do circumvent the complicated details of efficient preconditioners and general solver efficiencies based on both matrix condition number and potential high core counts with generally low work per core. However, depending on the time scales of interest of the problem, explicit methods can be painfully inefficient. In general, production Aria does not support explicit methods.

How to load a single displacement field.
   To load a previously computed displacement field and use it throughout the Aria thermal simulation, one needs to extract the displacement solution from a single time plane in the displacements file and assign it to the mesh_displacements Field in Aria. Internally Aria will update the current configuration, model_coordinates + mesh_displacements, and compute on that geometry.

Is it possible to write a plugin to create a boundary condition that prescribes an angular velocity to a block?
   Yes, one can write plugins for boundary conditions. However, in order to apply an angular velocity boundary condition to a block one would presumably be solving for the block motion.

Is there a way to apply a force on surface?
   In Aria forces cannot be applied directly at a node. Instead nodal forces are computed from integrated surface tractions:

   \[ BC \text{ Flux for solid on surface}_{10} = \text{Constant}_\text{Traction Y} = 1800.0 \]
What is the expression syntax for beam sections in Aria?
Syntax is similar to that of shells:

\[ \text{bar area = constant a = 0.000768} \]

How can one define a temporally-ramped velocity.
The syntax should be:

\[ \text{BC LINEAR_IN_TIME Dirichlet at surface_1 velocity_x coefs = 0.1} \]

If one wants to ramp up from \( \text{velocity}_x = C_0 \) to \( \text{velocity}_x = C_1 \) in time \( C_2 \) and then hold \( \text{velocity}_x \) at \( C_1 \) then try \( \text{BC RAMP_LINEAR_IN_TIME} \) instead:

\[ \text{BC RAMP_LINEAR_IN_TIME Dirichlet at surface_1 velocity_x coefs = C_0 C_1 C_2} \]

so that \( \text{velocity}_x = C_0 + (C_1-C_0) \cdot \min(t/C_2, 1.0) \). If the user manual is not handy, it is sometimes useful to extract the command syntax from the executable using:

\[ \text{sierra aria -i your.i -print-syntax | grep -i LINEAR_IN_TIME} \]

This produces a somewhat cryptic definition of the expected syntax but it sometimes helps.
41.3 Errors

**Cannot find a volume element on the surface error.**
This problem often occurs when attempting to use a nodeset in flux boundary condition definition. Aria applies surface sideset contributions as volume contributions and often uses volume information to lookup surface information. Thus Aria is unable to look up the volume element unless it has a sideset, even though the surface nodesets are specified.

**2D analysis fails due to incorrect element type used.**
Typically this occurs when the grid file uses some form of 3D elements. If the grid file was created in CUBIT, then one needs to change the element type before exporting the mesh. An example of how you do that in CUBIT is:

```
block 1 element type quad
```

Additionally one must make sure to use the `dimension 2` in the CUBIT export command in order to ensure that the z coordinate is not included in the output file.

**Unable to achieve convergence in anisotropic thermo-diffusion model.**
This class of problems is difficult to solve. First off, the matrix is non-symmetric, which is automatically harder to solve. Second, there are different sets of physics occurring in the problem (in this case, species and temperature), and each set of physics will have a different contribution. If their contributions aren’t within a couple orders of magnitude, then the problem also has great difficulty in converging. This has been viewed in thermoelectric models. In a few instances, the analyst was able to work around this difficulty by specifying a different system of units (e.g., changing from volts to millivolts). The idea here is to rescale the solution variables to be of comparable magnitude.

**Nonphysical temperature overshoot in 2-material conduction model.**
Typically, the temperature overshoot issue can be resolved by using `lumped_mass` instead of `mass` in the EQ specification. Use of the mass keyword invokes construction of a consistent mass matrix. While consistent mass is supported by a theoretical foundation for convergence, it oftentimes exhibits the behavior of overshooting. Most models which involve a large step change are better dealt with using a “lumped mass” matrix.

**Problem with enforcing species conservation.**
Usually a problem stems from inconsistent units being used in the models. Make sure the units for the reactions are consistent with everything else.

**TET10 interpolation for post-processing produces an error.**
Since the TET10 element is second-order, one needs to use Q2 in the EQ specifications and for post-processing, since in both cases there is interpolation involved.

**Contact resistance error: edges share more than two element faces.**
This can occur when portions of the specified contact surfaces are physically connected. Typically these type of problems are related to the mesh when unmerge command is used. Double checking the mesh would be the first place to start.
41.4 Solver

What do real residual check and time step fails mean?

Real Residual Check Failed means that the solve was unable to arrive at a solution that satisfied the solve tolerance specified in the radiosity solver command block. The Time Step Failed message in the log file means exactly that, the Newton step has failed to converge in the number of maximum number of nonlinear iterations specified. This will cause the time step to be reduced and the Newton step will be retried.

Unable to achieve convergence in anisotropic thermo-diffusion model.

This class of problems is difficult to solve. First off, the matrix is non-symmetric, which is automatically harder to solve. Second, there are different sets of physics occurring in the problem (in this case, species and temperature), and each set of physics will have a different contribution. If their contributions aren’t within a couple orders of magnitude, then the problem also has great difficulty in converging. This has been viewed in thermoelectric models. In a few instances, the analyst was able to work around this difficulty by specifying a different system of units (e.g., changing from volts to millivolts). The idea here is to rescale the solution variables to be of comparable magnitude.

Contact problem that is being solved is not converging.

Typically, this occurs when some blocks might not be constructing the appropriate contact constraints. By adjusting the interaction constraint tolerances might help to perform converged solves on a sub-model. The issue can be remedied by adding the following tolerances to the contact definition command block:

begin search options
  normal tolerance = 0.0001
  tangential tolerance = 0.00005
end search options

Also, the amount of contact present in a model will dictate the choice of linear solver. The recommended choice would be:

solution method = gmres
preconditioning method = dd-ilut

Aztec numerical breakdown error.

This specific problem occurs when the matrix near diagonal elements either go to 0 or approach a singularity. This is a common occurrence when fluids are involved.
41.5 Thermal

How to define function-based tensor thermal conductivity?
Assuming that functions are user defined functions (tabular functions). Letting the conductivity functions be func_kx, func_ky, func_kz respectively then the syntax for temperature dependent functions would be:

\[
\text{tensor thermal_conductivity} = \text{user_function Name_xx} = \text{func_kx Name_yy} = \text{func_kx Name_zz} = \text{func_ky X= temperature}
\]

If the functions are time dependent then only difference would be: X = time.

Does contact conductance contribute to the Joule heating term in the energy equation?
Joule heating is applied volumetrically. Contact conductance is modeled as an interface flux condition between opposing surfaces and has no intrinsic association with a volume. Hence contact conductance cannot contribute to Joule heating.

Can Aria solve a condensation problem?
There is ongoing work in studies of evaporation, but not yet condensation.

How to evaluate the time rate change in thermal energy.
The time rate change in thermal energy should be \( \rho \cdot C_p \cdot \frac{dT}{dt} \cdot \text{elem_volume} \). The results block should contain:

- nodal variables = time_derivative_at_time->TEMPERATURE as dtemp_dt
- nodal variables = pp->density as rho
- nodal variables = pp->specific_heat as cp
- element variables = current_element_volume as e_vol

Then within the EnSight calculator you can define a variable and evaluate the following expression:

\[
\rho \cdot C_p \cdot \frac{dT}{dt} \cdot e_{vol}
\]

How does Aria output the temperature of shell elements? Does it output the average or the outer or inner surface temperature?
For the standard shell there is only one nodal value of temperature. For the specialized implementations, linear and quadratic shells there are two nodal values of surface temperature, one associated with top and bottom of the shell.

Can one model a thermal symmetry boundary condition?
Yes, you can either specify no boundary condition on your symmetry surface, which will default to adiabatic, or specify a zero flux boundary condition, which will satisfy a symmetric solution.

How does one terminate a run based on variable rate of change (dT/dt)?
One can use the same approach for terminating a maximum rate of temperature change, dT/dt, in the same way as for using maximum temperature via Encore MIN MAX Postprocessor command block.

How does one specify variable density and specific heat?
One can specify both variable density and specific heat with the user_function.

What can one model with shell elements in Aria.
Typical usage of thermal shell elements in Aria is for materials, which have no through-thickness temperature variation but still model diffusion in the plane of the shell. The shell can also account for
thermal capacitance. Cases where the through thickness resistance of heat flow through the material is more important will generally be modeled with resistance contact (no shell). The in-between case is a combination of the above.

**How to link reference temperature to temperature of bulk fluid element used in convective flux boundary condition.**
Numerical modeling of heat transfer by enclosure radiation represents the transport between facets of the surface representation of a closed volume. Modeling this problem with enclosure radiation would require a surface representation of the body within the cavity where the closed volume is defined by the void region bounded by body and cavity. From a solution perspective, enclosure radiation heat transfer is de-coupled from the standard finite element solution. This means that one can solve the enclosure radiation problem and also include convective heat transfer from the surrounding cavity surface by computing an average body temperature and using that value as the reference temperature. Alternatively one could define a bulk element for the air cavity and setup the convective heat transfer problem to include both the cavity surface and body surface. Aria allows for superposition of heat flux contributions. In an approach with a bulk element representation of the body, the bulk element temperature would serve as the reference temperature for both convective and radiative heat transfer between the cavity surfaces and the body. If one wishes to treat the absorptive medium as being optically thin one would mesh the cavity interior and the body and use the Rosseland Mean approximation of thermal conductivity.

**How to save thermal conductivity as a nodal/element variable for output.**
This can be accomplished using the postprocess command. Use this command in the Region scope:

```
Postprocess Thermal_Conductivity on block_1 block_2 block_n
```

and in the Results Output, use this command:

```
Nodal Variables = pp->Thermal_Conductivity as K
```

**Is it possible to use radiation enclosure and heat loss by convection using the bulk fluid element approximation?**
In Aria flux contributions are cumulative. Both convection to a bulk node cavity temperature can be superposed with the enclosure radiation flux.

**How to represent thermal conductivity as a polynomial function of temperature.**
Since the independent variable is Temperature, specification would be:

```
something = Polynomial Variable=Temperature Order=2 C0=6.0789 C1=-0.0225 C2=4.0e5
```

**Element Death For Laser Melting Through Plate Array.**
Laser flux algorithms assume visibility of the source by the surfaces assigned in the flux definition. Hence, element death in each of the plates occurs simultaneously instead of in succession.

**How to calculate heat flux through a sideset.**
1. If it’s a global variable you want, then add the following block in the Region scope:

```
Begin Solution Options
  post process FLUX heat_conduction on surface_1 as Q_Surf
End Solution Options
```

Then you can use this as a global variable in your output blocks:
Begin Results OUTPUT AriaOutput
  global Variables = Q_surf
End Results OUTPUT AriaOutput

Or in heartbeat output:

Begin HEARTBEAT Heart1
  variable = global Q_Surf as Q_Surf
End HEARTBEAT Heart1

2. If you’d like the heat flux at each node point, then add this line to the Region scope of your input deck:

Postprocess Heat_Conduction on surface_1

Then you can use the nodal values in your output as such:

Begin Results OUTPUT AriaOutput
  Nodal Variables = pp->heat_conduction as Q
End Results OUTPUT AriaOutput

All of the post-processing commands can be found in the Postprocessor Operations chapter 21 of the Aria user manual.

**How to properly define contact to transfer heat.**

Contact requires that the mesh be discontiguously meshed at the contact interface with normals oriented into the contact interface. In the case that surface_1 and surface_2 have normals oriented in the same direction, the search will fail and no contact constraints will be applied, hence no heat transfer.

**Assigning a Dirichlet temperature BC to an entire block in aria.**

Try the following:

```
BC USER_FUNCTION_IN_TIME DIRICHLET ON BLOCK_3 TEMPERATURE = UserFunctionName
```

**Is there a way to apply a force on surface?**

Nodal forces are computed from integrated surface tractions:

```
BC Flux for solid on surface_10 = Constant_Traction Y = 1800.0
```

**Syntax for having a tensor thermal conductivity described by polynomials in both directions in Aria?**

The Aria code does not currently have an option to specify tensor thermal conductivity using polynomials. Alternative are to use either:

1. User Function tensor or
2. a Calore style user sub to compute and set the tensor.

**Estimating the heat flux.**

For the Aria application, there a couple of ways to estimate the heat flux at selected locations. The first method is to use the heat_flux postprocessor on all_blocks while the second is to use the calculator function in EnSight.
41.6 Radiation

**Viewfactor calculation doesn’t converge due to bad rowsum.**

Typically, this occurs when there is a problem with the mesh or the surfaces specified in the Enclosure Definition. The preferred method to debug enclosures is to add the following line to the Enclosure Definition Block:

```
Rowsum Database name = your_enclosure_file.e
```

Upon job execution Aria will output a file containing rowsums for each of the enclosure facets. One can then view the enclosure geometry colored by the rowsum value in order to spot the problem areas. For a good enclosure, each facet should have a viewfactor rowsum very near 1. Another option that one could try is to use the option Use Dash Enclosures at the Aria Region Scope. This option will “skin” all the blocks in the data set and will create all the enclosures that are contained therein.

**Is it possible to use radiation enclosure and heat loss by convection using the bulk fluid element approximation?**

In Aria flux contributions are cumulative. Both convection to a bulk node cavity temperature can be superposed with the enclosure radiation flux.

**Is there a way to rebalance aria viewfactors from a 64 node run to be used on a 16 node run?**

Currently, the view factors can be used only with the same number of processors for which they were originally computed.

**Is there a way to extract the incident heat flux seen by a component within an enclosure definition?**

Add these commands to the Begin Results Output block:

```
face variables = radiosity as J
face variables = irradiance as G
face variables = rad_flux as Q
```

**Problem with enclosure radiation calculations.**

Typically the root cause of this problem is having surface normals pointing in the wrong direction. Make sure that all sidesets have the correct surface normal.
41.7 Time Stepping

What do real residual check and time step fails mean?

Real Residual Check Failed means that the solve was unable to arrive at a solution that satisfied the solve tolerance specified in the radiosity solver command block. The Time Step Failed message in the log file means exactly that, the Newton step has failed to converge in the number of maximum number of nonlinear iterations specified. This will cause the time step to be reduced and the Newton step will be retried.

How does one terminate a run based on variable rate of change (dT/dt)?

One can use the same approach for terminating a maximum rate of temperature change, dT/dt, in the same way as for using maximum temperature via Encore MIN MAX Postprocessor command block.

Time step size not increasing.

Even after setting the maximum time step, the time step might be limited or constrained by the output increment in the Begin Results Output block.

Aria fails to exit when timestep refined to zero.

Simple workaround is after the Initial Time Step Size = 5 command, add the syntax Minimum Time Step Size = 5. This will prevent Aria from trying to half the time-step and will force it to exit.

Early-time results of steady-state run are nonphysical.

When one runs a linear to mildly non-linear model in a pseudo-transient manner, the intermediate results obtained will generally appear unreasonable and even nonphysical. Although it may appear that physical time steps are being taken, the only result that can be relied upon is the steady-state solution, as long as the solution has converged.

How can I transition a tabular material property from steady-state to transient?

The following input is sufficient for changing a material property between steady-state and transient analysis in the same simulation. Note that the steady-state material property goes from 0.0 to 1.0, and the transient portion comprises 1.0000001 and onward.

begin definition for function mat1_time
  type is piecewise linear
  abscissa = x_label
  ordinate = y_label
  begin values
    0.000000  0
    1.000000  0
    1.0000001 129
    1e9      129
  end values
end
41.8 Contact

What are the proper units for electrical contact conductance coefficient?
The units are equivalent to a heat transfer coefficient which would have the units of \(1/(\text{resistance} \times \text{area})\), which is equivalent to \((\text{current}/(\text{volts} \times \text{area}))\).

Can I define contact conductance as an analytic function?
No, this capability is not currently available.

Can one simulate a sliding block in Aria?
Yes, one can do this but the contact physics will be based strictly upon contact between the the two surfaces, without sliding friction effects.

How to model multiple thermal and electrical contact conductances.
Typically, attempts to use two contact definitions in the Aria Region fail due to conflicts between the conductance coefficient used in each of the contact definitions. This problem can be circumvented by using multiple Aria Regions or by defining multiple sidesets at the contact interface.

Contact problem that is being solved is not converging.
Typically, this occurs when some blocks might not be constructing the appropriate contact constraints. By adjusting the interaction constraint tolerances might help to perform converged solves on a sub-model. The issue can be remedied by adding the following tolerances to the contact definition command block:

```plaintext
begin search options
  normal tolerance = 0.0001
  tangential tolerance = 0.00005
end search options
```

These tolerances reflect physical distance which should represent the the geometry of the modeled discretization. Also, the amount of contact present in a model will dictate the choice of linear solver. For generalized contact, `tied_temperature` or `gap_conductance` the recommended solver is:

```plaintext
solution method = gmres
preconditioning method = dd-ilut
```

What are the proper units for electrical contact conductance coefficient?
The units are equivalent to a heat transfer coefficient which would have the units of \(1/(\text{resistance} \times \text{area})\), which is equivalent to \((\text{current}/(\text{volts} \times \text{area}))\).
41.9 Species

Does Aria support modeling a reaction that is dependent on temperature, pressure, and species concentration?
Yes, it is possible to implement a user plugin for quantities that are dependent on each of those.

Problem with enforcing species conservation.
Usually a problem stems from inconsistent units being used in the models. Make sure the units for the reactions are consistent with everything else.

Species diffusion subroutines cause non-physical results and residuals are high.
Try using a direct solver. At times, GMRES with Jacobi or DD-ILUT for preconditioning methods may not converge.
41.10 Porous Media

**How can one output the non-wetting phase pressure?**

Since the non-wetting phase pressure is needed in the wetting phase solution, it is available for output not as a solution variable but as a post-processed variable. In general, the pressure can be output provided one is able to specify its material phase. In this case the syntax would be:

```
postprocess non_wetting_phase_pressure on all_blocks
```

and the variable could be output to a results file as:

```
nodal variables = pp->non_wetting_phase_pressure as P_NW
```

**Can capillary pressure be added as postprocess variable?**

Yes, post-processing of capillary pressure is supported, which can later be added to an output.

**No expression for intrinsic permeability vs. time table.**

Intrinsic permeability is handled a bit differently than other porous media properties since the tensor character (ratio of coefficients) is normally preserved. Instead intrinsic permeability is modified by introducing a scaling factor:

```
Intrinsic Permeability = Constant XX=1.0 YY=1.0 ZZ=1.0
Intrinsic Permeability scaling = user_function name = Time_Permeability_Function x = time
```

**What is the difference between the options Porosity = Mesh_deforming and Porosity = Solid_deforming?**

Mesh_Deforming and Solid_Deforming are identical, except Mesh_Deforming works on the mesh equation and Solid_Deforming works on the solid equation.
41.11 Electrostatics

What are the proper units for electrical contact conductance coefficient?

The units are equivalent to a heat transfer coefficient which would have the units of \(1/(\text{resistance}\times\text{area})\),
which is equivalent to \((\text{current}/(\text{volts}\times\text{area}))\).

Is Aria capable of calculating and outputting an electromotive force resulting from a body in an electric field?

Currently, there is no native postprocessor in the Aria code to compute resultant electromotive force.
41.12 Grid Refinement

Can Aria do focused mesh refinement?
Yes, the manner in which the refinement occurs is controlled by the `marker_field`. Most of the adaptivity support is finite element based, single level of refinement and is driven by Encore. The necessary commands include:

```
Begin Sierra Aria
  Begin Uniform Marker uniform_mark
    Store in marker_field
  End Uniform Marker uniform_mark

Begin Procedure AriaProcedure
  Begin Solution Control Description
    Use System Main
    Begin System Main
      MarkAdapt AriaRegion using uniform_mark
      Begin Transient TransientBlock
        Advance AriaRegion
      End Transient TransientBlock
      Begin Parameters for Transient TransientBlock
        ...
  End System Main
End Solution Control Description
```

When using uniform mesh refinement, does the new mesh get saved somewhere?
The new mesh is on the output from the uniform mesh refinement run. The disadvantage with that is the file size with multiple time planes in it. A more economical option is to use an additional Results Output block that saves zero time plane with the refined mesh. The syntax would be something like that in the example where the `At Step 0 Interval = 10e10` being ridiculously high should cause only one time plane to appear in the output file. One may also use the Sierra executable `stk_adapt_exe` for uniform mesh refinement - documentation for this application may be found on Compsim.

Problem encountered with using 8-node quad.
In order to support mesh refinement in the same way for all elements, Aria does not support the 8-node quad. It only supports 9-node element.

How does one coarsen a mesh?
The input modifications consist of two new lines:

```
MarkAdapt AriaRegion Using total\_mark When "recover\_ind > 0.5"
MarkAdapt AriaRegion Using total\_mark2 When "recover\_ind < 0.25"
```

This syntax turns on refinement when the error indicator exceeds the value 0.5 and switches to coarsening when the error is less than 0.25. When the error is between these values, the mesh is left unchanged.
41.13 Post Processing

**How can one output the non-wetting phase pressure?**

Since the non-wetting phase pressure is needed in the wetting phase solution, it is available for output not as a solution variable but as a post-processed variable. In general, the pressure can be output provided one is able to specify its material phase. In this case the syntax would be:

```plaintext
postprocess non_wetting_phase_pressure on all_blocks
```

and the variable could be output to a results file as:

```plaintext
nodal variables = pp->non_wetting_phase_pressure as P_NW
```

**How to output the min and max temp on specified blocks.**

This can be accomplished using Encore:

```plaintext
$---------------------------------------------------
$ Specify Encore output to text file
$---------------------------------------------------
Begin Postprocessor Output Control Encore_Out
  Output Time
  Write To File TempMinMax.txt
  Floating Point Precision Is 3
  Floating Point Format Is Fixed
End   Postprocessor Output Control Encore_Out
$---------------------------------------------------
$ Create Encore field function
$---------------------------------------------------
Begin Field Function spos
  Use Nodal Field Solution->Temperature
End   Field Function spos
$---------------------------------------------------
$ Determine maximum value of field function
$---------------------------------------------------
Begin Min Max Postprocessor TMinMax_1
  Use Function spos
  Compute Min Max
  Volumes block_1
End   Min Max Postprocessor TMinMax_1

Begin Procedure AriaProcedure
  Begin Aria Region AriaRegion
    Evaluate Postprocessor TMinMax_1
  ...

How to output mass inventory.

Include the following commands at the Aria Region scope:
Model hangs when trying to output elem_temperature.
The proper syntax for postprocessing Temperature as an element variable is:

```
Postprocess TEMPERATURE on all_blocks using P0
```

This will interpolate the nodal temperature as a single element value, a zeroth-order polynomial (P0).

**How to evaluate the time rate change in thermal energy.**
The time rate change in thermal energy should be $\rho \cdot C_p \cdot (dT/dt) \cdot elem\_volume$. The results block should contain:

```
nodal variables = time_derivative_at_time->TEMPERATURE as dtemp_dt
nodal variables = pp->density as rho
nodal variables = pp->specific_heat as cp
element variables = current_element_volume as e_vol
```

Then within the EnSight calculator you can define a variable and evaluate the following expression:

```
rho*cp *dtemp_dt*e_vol
```

**How does Aria output the temperature of shell elements? Does it output the average or the outer or inner surface temperature?**
For the standard shell there is only one nodal value of temperature. For the specialized implementations, linear and quadratic shells there are two nodal values of surface temperature, one associated with top and bottom of the shell.

**How to save thermal conductivity as a nodal/element variable for output.**
This can be accomplished using the postprocess command. Use this command in the Region scope:

```
Postprocess Thermal_Conductivity on block_1 block_2 block_n
```

and in the Results Output, use this command:

```
Nodal Variables = pp->Thermal_Conductivity as K
```

**Can capillary pressure be added as postprocess variable?**
Yes, post-processing of capillary pressure is supported, which can later be added to an output.

**How to visualize the results continuously regardless of the time step selected by ARIA?**
To visualize the results of an adaptively time-stepped simulation in an equal-time-step fashion, one can use a ParaView filter, “Temporal Interpolator”. This filter allows one to view a time-dependent data set where the data’s field data between adjacent time steps is linearly interpolated.
How to output an ASCII history file for data at a selected location over time.
One way of getting the history output that one specifies is to use Encore to do the interpolation. Graphics packages such as EnSight and ParaView can then be used to generate time history of an output parameter at selected locations.

**TET10 interpolation for post-processing produces an error.**
Since the TET10 element is second-order, one needs to use Q2 in the EQ specifications and for post-processing, since in both cases there is interpolation involved.

How to output element block volumes for steady-state simulation.
In order to output volume and mass in Aria one uses the following command block in the Aria Region:

```plaintext
Begin Solution Options
  post process volume
End   Solution Options
```

Log file has unwanted postprocessor history at every time step.
Remember to comment out the following:

```plaintext
begin postprocessor output control test
  output to log file
end   postprocessor output control test
```

Is there a way to apply a force on surface?
Nodal forces are computed from integrated surface tractions:

```plaintext
BC Flux for solid on surface_10 = Constant_Traction Y = 1800.0
```

Using Tecplot to plotting/visualize Aria results.
One can use exotec2 - the syntax is:

```plaintext
exotec2 exo_in tec_out
```

where `exo_in` is the exodus file and `tec_out` is the Tecplot translation.

Is there a way to extract the incident heat flux seen by a component within an enclosure definition?
Add these commands to the Begin Results Output block:

```plaintext
face variables = radiosity as J
face variables = irradiance as G
face variables = rad_flux as Q
```

Estimating the heat flux.
For the Aria application, there a couple of ways to estimate the heat flux at selected locations. The first method is to use the `heat_flux` postprocessor on `all_blocks` while the second is to use the calculator function in EnSight.
Chapter 42

Developer Documentation

When working within the code base developers may wish to setup their environment to reference the development toolset.

42.1 Setting The Environment-Developers at Sandia Labs

The environment for using Aria is the same as for individual Sierra applications and can be configured by module files. The modules ensure that the look and feel of running Sierra applications is the same across a multitude of compute platforms. To obtain the proper environment for code execution one simply runs:

```
$ module load sierra-devel
```

42.1.1 Setting up for the csh and tcsh Shells

Add SNTools to your path. In either your `.cshrc` or `.tcshrc` file add the line

```
set path=(/home/sntools/devel/linux/install/sntools/engine $path)
```

42.1.2 Setting up for the bash Shell

Add SNTools to your path. In either your `.bashrc` or `.profile` file add the line

```
export PATH=/home/sntools/devel/linux/sntools/engine:$PATH
```

42.2 An Introduction to Aria’s Expression System

The following is the conference paper from the Coupled Problems 2005 conference, See [42].

This section provides a brief overview of a core element of Aria’s design called the Expression system. Understanding the Expression system is not essential for using Aria but it is useful in understanding how some of Aria’s features behave. It is also the simplest point of entry for developing and extending Aria with new material properties and boundary conditions. In short, the Expression system is the abstraction of low- to mid-level numerical calculations so as to make the code highly general, reusable, extensible and scalable.

In order to provide a low entry barrier for user extensions we have chosen a design where the essential numerical entities of boundary conditions, material properties, etc., are all implemented the same way.
Specifically, these are implemented as C++ classes called Expressions. In addition to supplying the functional evaluation of the numerical quantity, Expression implementers provide information regarding what the Expression provides, what other Expressions it may depend on, the tensorial order of the provided quantity, etc. A higher-level Expression Manager is then able to determine the minimal and sufficient set of Expressions required to perform the simulation and to ensure that they are evaluated in the proper order. In addition to providing a simple extension technique, this makes the core development of Aria very clean and abstracts nonessential details from many of the basic algorithms, while still employing the most efficient numerical kernels and data management techniques at the lowest levels.

To motivate our approach, we start with a simple example. In the FEM solution of the Navier-Stokes equations the following integral, among others, is computed.

$$
\int_V \rho v \cdot \nabla \phi^i \, dV = \sum_e \int_{V_e} \rho v \cdot \nabla \phi^i \, |j| \, dV_e
$$

(42.1)

Here \( \rho \) is the density, \( v \) is the fluid velocity and \( \phi^i \) is weight function associated with local node \( i \). \( V \) is the domain of integration in the physical space and \( V_e \) is that in the space of the element with \( |j| \) being the Jacobian of transformation from \( V_e \) to \( V \). Thus, the assembly kernel responsible for this calculation needs access to all of these quantities. We will show that the kernel, however, does not need to know the particulars of, say, the density model or of the weight function, especially regarding which other quantities they may depend on. Moreover, there are likely to be other assembly kernels that may also require these same quantities, such as the advection term of the energy transport equation,

$$
\int_V \rho C_p v \cdot \nabla T \phi^i \, dV = \sum_e \int_{V_e} \rho C_p v \cdot \nabla T \phi^i \, |j| \, dV_e
$$

(42.2)

where \( C_p \) is the specific heat and \( T \) is the temperature. Efficient evaluation of (42.2) should make use of \( \rho, |j|, v \) and any other quantities that might have previously been computed in the assembly of (42.1). Keeping track of which quantities are needed and which quantities have already been evaluated in a previous part of the assembly, regardless of the equation system that is being solved, is the purpose of Aria’s Expression Manager.

In Aria, all of the quantities presented above, \( \rho, \nabla v, |j|, \phi^i \), etc., are implemented as C++ classes derived from the base class Expression. Each Expression has these basic responsibilities:

- **Identification.** Each Expression identifies itself with a generic description, such as density and, when applicable, a specific model description, such as cubic_in_temperature.

- **Prerequisites.** Each Expression (and Expression user) provides a list of other Expressions (using the generic identifiers) that are required for its own calculations. To continue the example, the Expression which implements the cubic_in_temperature model would have temperature as a prerequisite. Arbitrarily complex terms may be built of successively simpler components which ultimately result in a dependency tree with the simplest ingredients being the leaves of the graph.

- **Tensor Properties.** Each Expression states its tensor order (scalar, vector, second order tensor, etc.) and dimension. With this information, Expression users can dynamically determine how to index the values provided by the expression. For example, some thermal conductivities may be scalars while others may be second order tensors as is the case with anisotropic materials.

- **Evaluation.** Each Expression implements a virtual function for computing its values. Depending upon the nonlinear solution strategy, additional methods may be required for computing, for example, Newton sensitivities.

When an Expression is created, it makes itself known to an Expression Manager. The Expression Manager is responsible for establishing the minimal but sufficient list of Expressions required for the calculation. This is
done utilizing the lists of prerequisites provided by Expression users and other Expressions. The relationships among Expressions, Expression users, and the Expression Manager is illustrated schematically in figure 42.1 with the use of UML. This figure illustrates two salient features of our Expression subsystem: Expressions are themselves Expression users and Expression users, such as assembly kernels, are the original source for all prerequisites.

The richness of the information provided in these lists of prerequisites quickly becomes apparent. For example, the Expression Manager is able to further utilize the prerequisite information to order the list of Expressions for evaluation. This is done so that when an Expression is evaluated, all of its prerequisite Expressions have already been evaluated. Regardless of how complicated the multiphysics problem happens to be, there are no duplicate evaluations and no overhead costs that would be incurred with lazy-evaluation. Another powerful use for the prerequisite information is in dynamically determining and computing sensitivities for Newton’s method. Since primitive variables, or degrees of freedom, are also represented as Expressions, the prerequisites for any given Expression are recursively checked to see if any of them represents a primitive variable. In combination with runtime-queried tensor properties (see above), an Expression developer can implement and evaluate the Expression’s sensitivities without ever having direct knowledge of the degrees of freedom in the system. Lastly, Aria utilizes this information for additional purposes such as sparse matrix allocation and automated memory management.

42.3 Nonlinear Coupling Strategies in Aria

One of the difficulties with writing broadly applicable computational mechanics software is that developers can’t take advantage of specific knowledge of the application domain in order to optimize the algorithm. Thus, in providing generality one sometimes sacrifices efficiency. One place this is evident in multiphysics modeling is in the choice of coupling strategies. While it is well understood that a fully coupled system solved with Newton’s method utilizing analytic sensitivities is formally the most robust and correct approach to solving multiphysics applications it is also computationally expensive and complex to implement. Furthermore, while Newton’s method has the fastest rate of asymptotic convergence it’s domain of convergence is often empirically observed to be smaller than other methods. Lastly, in some applications, certain subsets of the physics may be only weakly coupled so that a loosely coupled approach may be more computationally
efficient. To address these concerns while remaining general and flexible Aria offers a number of options for nonlinear solution strategies and physics coupling.

In defining a problem in Aria, users configure one or more Regions. Each Region consists of one or more PDEs to be solved on some or all of the input mesh. All of the PDEs in each Region are solved in a tightly coupled (i.e., single matrix) manner using one of several nonlinear solution strategies available. Users may then define loose couplings between two or more Regions. For example, some or all of a solution from one Region may be transferred to another Region where it is treated as a constant, external field. The aggregate nonlinear problem including the contributions from all of the Regions may be iterated to convergence. The particulars of which physics are solved in each Region and the nonlinear solution strategy used within and between Regions is completely specified through the input file. Furthermore, an Aria user may pick a simple, minimal algorithm without needing to fit it into an overly-generalized worst-case scenario that represents the union of all possible algorithms.

Dynamically-specified loose coupling has many potential advantages that users may leverage. First, the resulting linear system is considerably smaller and contains far fewer off-diagonal contributions which can significantly increase the performance of linear solvers. Also, a resulting linear system may have a more attractive form, such as symmetric positive-definite, that permits the use of tailored iterative solutions techniques. Other extensions to loose coupling include subcycling of transient simulations where each Region may advance in time with its own time step size and in-core coupling to other applications based upon the Sierra framework.

### 42.4 Developer Recipes

#### 42.4.1 Adding a New Flux Boundary Condition

1. Add a new name for your model
   (a) Add a new Identifier entry to Aria_Model_Names.h
   (b) Add the string in Aria_Model_Names.C
2. Add your new Expression class.
   (a) Copy-paste-rename a similar Expression definition. See, e.g., Aria_Material_Models.h.
   (b) Copy-paste-rename a similar Expression implementation. See, e.g., Aria_Material_Models.C.
3. Add an entry to the Expression_Factory::create_model_expression().
   (a) Copy-paste-rename a similar entry in Aria_Expression_Factory.C.

#### 42.4.2 Adding a New Material Model

1. Add a new name for your model
   (a) Add a new Identifier entry to Aria_Model_Names.h
   (b) Add the string in Aria_Model_Names.C
2. Add your new Expression class.
   (a) Copy-paste-rename a similar Expression definition. See, e.g., Aria_Material_Models.h.
   (b) Copy-paste-rename a similar Expression implementation. See, e.g., Aria_Material_Models.C.
3. Add an entry to the Expression_Factory::create_model_expression().
   (a) Copy-paste-rename a similar entry in Aria_Expression_Factory.C.
42.5 Expression Reference and API

This section provides a high-level view of how Expressions are run by Aria, from creation to computation. This section also provides a brief description of the Expression methods that a user either must or can optionally supply.

42.5.1 Execution Sequence

1. Initialization Phase (Input Parsing Time)

   (a) Expression::Expression() (See 42.5.2) : Constructor. Called at creation time. Defines identity, dependents and parameters. Required.

   (b) Expression_Manager::preprocess_expressions() : Called after all expressions have been created and all dependencies have been registered/requested. Unused expressions are destroyed. Dependency based ordering established.

   (c) Expression::set_nonlinear_method() : Sets the nonlinear method that will be used.

   (d) Expression::create_dynamic_storage() : Initializes storage for values and Newton sensitivities but does not actually allocate memory.

   (e) Expression::postregistration_setup() (See 42.5.3) : Called once, after the Expression_Manager has determined that the Expression will indeed be used but before simulation begins. Called in dependency order. Can get references to dependent’s storage (values, Newton sensitivities, etc.).

   (f) Kernel::postregistration_setup() : Assembly kernels can interrogate the Expressions available to them and get references to values and Newton sensitivities.

2. Simulation Phase

   (a) Expression::prepare_to_recompute() (See 42.5.4): Called once for each workset, immediately prior to compute_values(). Can be used to resize or reinitialize storage (usually done by base class), inquire about simulation time, etc.

   (b) Expression::compute_values() (See 42.5.5) : Computes the values of the Expression. Required to compile.

   (c) Expression::compute_sensitivities() (See 42.5.6) : Computes the Newton sensitivities of the Expression. Required to compile.

3. Shutdown Phase

   (a) Expression::Expression() (See 42.5.7) : Destructor. Can be used to free memory allocations (note: Real_MDArr storage is automatically de-allocated). Required but normally empty.
42.5.2 Constructor

42.5.3 postregistration_setup()

42.5.4 prepare_to_recompute()

42.5.5 compute_values()

42.5.6 compute_sensitivities()

42.5.7 Destructor

42.6 Newton Sensitivity Checking for Expressions

When you run Aria you can ask it to look for sensitivity errors in every Expression if you’re using Newton’s method. If activated, Aria will compare the analytic sensitivities provided by each expression with numerically computed values. This feature is enabled by adding the command line option -arialog sens_check to Aria. If you’re using the sierra tool to run Aria, then add -O "-arialog sens_check" (including quotes) to the sierra command line. The same holds if you’re running the Arpeggio application.

You can also run the entire test suite with Jacobian checking enabled by adding -u aria_args = "-arialog sens_check" to the runtest command line.

42.7 Profiling

It’s a good idea to profile the code before spending time trying to optimizing and performance tuning. Fortunately, profiling is pretty easy to do. On Linux, one approach is to use the gprof tool. To profile code with gprof, you’ll need to compile with the -pg option. The easiest way to do this is to set environment variables USER_CFLAGS and USER_LDFLAGS to -pg. In the bash/ksh/zsh shell, use, e.g., export USER_CFLAGS=-pg and in tcsh/csh use setenv USER_CFLAGS -pg. You’ll only get detailed profiling information for code compiled and linked with this flag so you may want to recompile some of the framework too. You can profile debug or optimized code.

With the instrumented executable, run Aria as you normally would. This will generate a file called gmon.out. Lastly, run gprof to analyze the data: gprof executable > gprof.txt where executable is the executable (probably including an absolute or relative path) used in the run. The output stored in gprof.txt will give you some real data telling where the real bottle necks are.

42.8 Purify: Memory Analysis and Debugging

1. Build Aria on Linux. This works for both optimized and debug modes but use debug if you want to see line numbers in Purify.

2. Copy the complete link line into a file.

3. Add -show option to mpiCC.

4. Remove the -static option from the link line arguments.
5. Adjust any path to Apps\aria.o if necessary.

6. Put which mpiCC on the previous line to verify you’re getting mpiCC from /usr/local/mpi/sierra/mpich/1.2.5.2/gcc-3.2.2/bin/.

7. You might also verify gcc is version 3.2.2 (gcc -v).

8. Source the script.

9. Copy the link line (start with g++ ...) and paste it into a file.

10. Preface g++ with purify, e.g., purify -always-use-cache-dir -cache-dir=’pwd’/ g++ ...

11. Source this script and it will instrument your executable. It takes a while...

12. Optionally verify that the executable is purifyed by running ldd aria.x

13. Copy executable to aria/bin directory, removing the .tmp extension.

14. Run it standalone (via sierra, runtest or by hand) or with TotalView.

42.9 Building Against Other Projects

Sometimes it can be very useful to build against changes that have been made in a different project. For example, if you are waiting for a project’s changes to be checked in but would like to start developing against those changes in another project. To do this with the SNTools build tool you can use the -a option to build. The argument to this option is the XML file for each product you want to include in your build. So, if you want to use the version of Krino found in a project located in /path/to/project/ then you would add -a /path/to/project/krino/krino.sn.xml to your normal build command line.

42.10 Interfacing with MATLAB

42.10.1 Reading Aria Matrices Into MATLAB

This is sort of an “old” way of getting matrices into MATLAB but it still works just fine.

From Matt Hopkins:

You can read a .mtx matrix into MATLAB via the following. Let my_matrix.mtx be the matrix sitting in the debug output path you’re interested in.

```matlab
>> A_ascii = load(’my_matrix.mtx’);
>> A = spconvert(A_ascii(2:end,:));
```

And now A is in MATLAB’s sparse matrix format. It even ignores the zeros in the .mtx file (does not allocate space for them).

From there you can do some pretty nifty things. spy(A) does sort of what matvis does (graphical view of matrix nonzeros). condest(A) estimates the 1-norm condition number of A (might take a while for large A). Etc.
42.10.2 Interfacing Aria from within MATLAB

Russell Hooper is developing a pretty sophisticated MATLAB interface that allows you to control Aria and probe nonlinear and linear solver data structures directly from within MATLAB. This is rapidly evolving so no details are given here... except for how to build and run Aria/MATLAB.

This recipe is taken from an email exchange between Matt Hopkins and Russell Hooper in early April 2006. It most likely only works on Linux.

% setenv LD_LIBRARY_PATH /usr/local/matlab/7.2/bin/glnx86:$LD_LIBRARY_PATH

% build aria -j8 -a /home/rhoope/Trilinos_6.0 \
    CXXFLAGS="-I/usr/local/matlab/7.2/extern/include/" \
    -a /home/rhoope/projects/noxMatlabStable/equationsolver \
    LDFLAGS="-L/usr/local/matlab/7.2/bin/glnx86 -leng"

If you want to build a debug version, add “-g” to both the CXXFLAGS and LDFLAGS.

To execute aria, first invoke sierra aria -i input.i -script-only. Then copy the file sierra.sh into another, ie opt.sh. Finally, remove the “-o logfilename” argument from opt.sh so that your run is truly interactive.

42.11 Error Handling

Traditional techniques for error handling of numerous and varied. Sometimes functions will return special values indicating success or failure of the call. Sometimes global variables or flags are set to indicate an error has occurred. Sometimes abort() is simply called (bad!).

C++ and the SIERRA Framework offer facilities to support error handling. The two primary error handling techniques are exception handling and assertions. Both of these can be combined to implement Design by Contract though this is not formally done in Aria.

Whether one should use an exception or an assertion is sometimes a subtle question. In general, assertions should be used for cases where the error is not likely to occur due merely to user input. However, assertions can also be useful for computationally expensive tests which would adversely affect production calculations.

42.11.1 Exception Handling

C++ offers exception handling and the SIERRA Framework utility library provides a parallel-safe layer for exception handling. Because exceptions in C++ are ordinary classes, applications may choose to specialize exceptions for handling and detecting specific failure modes. When SIERRA exceptions are used in conjunction with the SIERRA diagnostic tracing facility, it is possibly to get a stack trace (or sometimes a partial one) that illustrates the code path the where the exception was thrown.

In most cases, Aria developers do not need to worry about catching exceptions (they’re normally fatal errors). The top-level SIERRA Framework routine run_sierra() handles the responsibility of catching any uncaught exceptions, converting them into parallel exceptions if necessary, and propagating the exception to any other processors.
If the reader is not already familiar with exception handling in C++, a good place to start is with [43]. In its simplest form, SIERRA exceptions can be used as follows.

```cpp
#include <util/Exception.h>
// ...
if(something_bad_happened)
{
    sierra::Exception x("Nice descriptive error message.");
    throw x;
}
```

The `sierra::Exception` class inherits from `std::string` and so it supports all of `std::string`'s operations, most notably the `+` (concatenation) operator.

```cpp
#include <util/Exception.h>
// ...
if(something_bad_happened)
{
    sierra::Exception x("Nice descriptive error message.");
    x += " More info here.";
    throw x;
}
```

Additionally, the `sierra::Exception` class supports the put-to operator, `<<`, for stream-like handling.

```cpp
#include <util/Exception.h>
// ...
if(something_bad_happened)
{
    sierra::Exception x;
    x << "Nice descriptive error message."
        << " More info here.";
    throw x;
}
```

However, in order to keep the `sierra::Exception` class lightweight, it does not inherit from `std::ostringstream`. Instead, `operator<<` is only defined for the most common objects, such as strings, integers, floats/doubles, etc. The consequence of this is that if you define a class with `operator<<` for `ostream`, that operator will not work with `sierra::Exception` objects. In these cases, it is sometimes useful to use a `std::ostringstream` to construct the error message and then pass the resulting `std::string` to the exception.

```cpp
#include <util/Exception.h>
#include <sstream>
// ...
if(something_bad_happened)
{
    std::ostringstream os;
    os << "Nice descriptive error message."
       << " Object foo: ";
    throw os;
}
```
Finally, the construction and throwing of the exception can occur on the same line for brevity. In this case, all we need is a temporary object which is unnamed.

```cpp
#include <util/Exception.h>
// ...
if(something_bad_happened)
{
    throw sierra::Exception() << "Nice descriptive error message."
        << " Integer out of bounds: j = "
        << j;
}
```

### 42.11.2 Assertions

Assertions are a common way to ensure that certain conditions that are assumed by the code to be true are indeed true. For example, a function or piece of code may require that a pointer be non-NULL, an integer be greater than zero, etc. Using the C++ `assert()` macro is a good way to test such conditions that should hold under normal circumstances. By using `assert()`, the enclosed test will only be performed if the code is compiled in debug mode; in optimized mode, the macros expand to nothing. Thus, developers can use asserts extensively to test conditions assumed by their code without affecting production-mode performance. Assertions also provide an excellent way to document and enforce the assumptions that the developer made in writing the code.

The SIERRA Framework provides an alternate implementation of the `assert()` macro that utilizes the parallel-safe exception handling described above. The macro, `ThrowAssert()`, is defined in the same header file as the exception class.

```cpp
#include <util/Exception.h>
// ...
void
some_function(const MyClass * myclass_ptr)
{
    ThrowAssert(0 != myclass_ptr);
    // It should be OK to use the pointer.
    // ...
}
```

Because assertions expand to empty code in optimized mode, developers must be careful to never put variable assignments or other state-altering code inside an assert.

```cpp
#include <util/Exception.h>
// ...
// This is a bug:
ThrowAssert(j = i + 2 > 4);
```

A common trick to getting more useful messages from a failed assertion is to add a help string like this:
#include <util/Exception.h>
// ...

void
some_function(const Int & num)
{
    ThrowAssert(num > 10 &&
                "This function only works properly when num is larger than 10");
    // It should be OK to use the pointer.
    // ...

Lastly, another macro, ThrowRequire() is available. This macro is identical to the ThrowAssert() macro except this it is always enabled, even in optimized builds. This macro is currently not used in Aria but use it if you want. The concise and readable form of these macros can help keep code short while still being readable and expressive.

42.12 Outputting User Information (Logging)

The SIERRA Framework utility library supplies a flexible logging facility. The problem with normal printf and cout output functions is that they get really annoying in parallel. Further, these messages simply roll off the top of the user’s screen unless they are careful to redirect the standard output and error streams to a file. Lastly, it’s difficult, and may involve recompiling the application, to tailor these messages depending on the kind of information a user is interested in.

42.12.1 The DiagWriter Logging Facility

SIERRA’s logging facility, DiagWriter, is designed to address these three issues. The DiagWriter class, in the simplest sense, is a std::ostream class that supports the put-to («) operator for writing messages.

#include <Aria_DiagWriter.h>
// ...
namespace Aria
{
    arialog << "This is a message the user will always see." << std::endl;
}

Note the use of std::endl instead of std::endl (the standard and portable newline). This is an unfortunate but important point; using std::endl will result in ugly compiler errors.

The most interesting feature of the DiagWriter class is that the output can be tagged with a bit field called a message mask. These masks are defined in Aria_DiagWriter_fwd.h as enums with meaningful names, e.g., LOG_EXPRESSION and LOG_NONLINEAR. To mask your output message, you can use the m() method on the DiagWriter object.

#include <Aria_DiagWriter.h>
// ...
namespace Aria
{
    arialog.m(LOG_EXPRESSION)
    << "This message is supposedly related to Expressions."
Messages that contain a mask are only written if that is enabled via the print mask and messages that don’t have a specified mask are always written. The user can define a print mask using the “-arialog string” command line argument where the provided string maps to one (or more) of the bit masks. The file Aria_DiagWriter.C contains the mapping between the string names and the bit mask values. For example, “expression” is assigned to LOG_EXPRESSION. When the user supplies the command line argument “-arialog expression” then the messages written are those masked with LOG_EXPRESSION and those with no bit masks.

The DiagWriter class contains many more features that are beyond the scope of this section. Explore the header files and the code for more examples.

42.12.2 The Tracing Facility

Tracing is a common and extremely useful debugging utility. With tracing enabled, the code prints the name of each function as it enters and exits the function. C++ doesn’t provide any standard way to accomplish this so the Trace class is provided by the SIERRA Framework utility library.

The Trace class works by creating an instance (object) as the first line of a. The object constructor takes as an argument a string (char *) containing the name of the function. When tracing is enabled the constructor prints this name. When the application leaves this function, the local Trace object is automatically destroyed as it goes out of scope. When tracing is enabled, the automatically-called destructor re-prints the function name. The output of nested functions is nested so the output provides a hierarchical view of the flow of control. The most basic usage looks like this:

```
#include <Aria_DiagWriter.h>

namespace Aria
{
    void my_func(const Int & i)
    {
        Trace diag_trace("Aria::my_func(const Int & i)");
        //... normal code follows
    }
}
```

The Trace class utilizes the DiagWriter class for writing and for determining if tracing is enabled. So, tracing may be enabled using “-arialog trace” on the command line and Aria::Trace objects mask all output with LOG_TRACE.

Though it’s not discussed here, tracing can be turned on and off during the normal execution so that tracing information can be gathered only when it’s desired. Read through the header files and the code for examples of doing that. One example of where this is used is in the exception handling. When an exception is thrown, the tracing bit mask is automatically turned on. As the function stack unwinds during the throw, any functions instrumented with Trace objects will get destructed and hence they will print their owning function name. The result is a stack trace, or traceback, which can be extremely useful for debugging.

Adding all of the Trace objects to a code can be a daunting task, especially if you’re really anal and
want the function arguments and namespace to be a part of the function’s printed name (which can be important with polymorphic functions). Aria uses the traceString tool (http://tracestring.sourceforge.net/) to automatically instrument the code with the Trace objects. The traceString tool will enclose the Trace objects in a pair of special strings so that it can safely update or remove the inserted code later on. Typically, it looks like this:

```c
#include <Aria_DiagWriter.h>
//...
namespace Aria
{
    void my_func(const Int & i)
    {
        /* %TRACE[ON]% */ Trace diag_trace("Aria::my_func(const Int & i)"); /* %TRACE% */
        //... normal code follows
    }
}
```

To learn more about traceString, talk to Pat...

### 42.13 Catalogue of Assembly Kernel Expressions

Aria supplies several generic expressions that can be used for top-level assembly kernels for equation terms. When adding new equations or terms to existing equations, one of the follow generic expressions can often be used.

#### 42.13.1 Scalar _Source_ Kernel _Expression_

This Expression assembles a source term for a scalar transport equation. Note that the MASS and SRC terms can both be cast in this form. The general form is:

\[
-m \sum_{g=1}^{N_g} \left( \prod_{r=1}^{N_c} c_r(x_g) \right) \left( \sum_{s=1}^{N_s} f_s(x_g) \right) \phi^i(x_g)|j|w_g
\]  

(42.3)

Here, \( m \) is a multiplier that defaults to 1, \( N_g \) is the number of Gauss points in the quadrature rule, \( N_c \) is the number of coefficients and \( N_s \) is the number of sources provided. As an example, consider the term,

\[
\int_{\Omega_e} \rho C_p \frac{\partial T}{\partial t} |j| \phi^i \, d\Omega_e = \sum_{g=1}^{N_g} \rho C_p \frac{\partial T}{\partial t} |j| \phi^i w_g
\]  

(42.4)

In this example, \( N_r = 2 \) with \( c_1 = \rho \) and \( c_2 = C_p \); \( N_s = 1 \) with \( f_1 = \frac{\partial T}{\partial t} \); and \( m = -1 \). As a side note, in the code the quantities \( |j| \) and \( w_g \) are treated as additional coefficients for convenience.

#### 42.13.2 Scalar _Advection_ Kernel _Expression_

This Expression assembles an advection term for a scalar transport equation. The general form is:

\[
m \sum_{g=1}^{N_g} \left( \prod_{r=1}^{N_c} c_r(x_g) \right) v_a \cdot \nabla S(x_g) \phi^i(x_g)|j|w_g
\]  

(42.5)
Here, $m$ is a multiplier that defaults to 1, $N_g$ is the number of Gauss points in the quadrature rule, $N_c$ is the number of coefficients, $v_a$ is a configurable advection velocity and $S$ is a scalar field. As an example, consider the term,

$$\int_{\Omega} \rho C_p v \cdot \nabla T |j| \phi^i \, d\Omega_c = \sum_{g=1}^{N_g} \rho C_p v \cdot \nabla T |j| \phi^i w_g$$

(42.6)

In this example, $N_r = 2$ with $c_1 = \rho$ and $c_2 = C_p$, $S = T$; $v_a = v$; and $m = 1$. As a side note, in the code the quantities $|j|$ and $w_g$ are treated as additional coefficients for convenience.

### 42.13.3 Scalar_Diffusion_Kernel_Expression

This Expression assembles a diffusion term for a scalar transport equation. The general form is:

$$m \sum_{g=1}^{N_g} \left( \prod_{r=1}^{N_r} c_r(x_g) \right) \left( \sum_{k=1}^{N_F} F_k(x_g) \right) \cdot \nabla \phi^i(x_g) |j| w_g$$

(42.7)

Here, $m$ is a multiplier that defaults to 1, $N_g$ is the number of Gauss points in the quadrature rule, $N_c$ is the number of coefficients and $N_F$ is the number of flux models provided. As an example, consider the term,

$$\int_{\Omega_c} q \nabla \phi^i |j| \, d\Omega_c = \sum_{g=1}^{N_g} q \cdot \nabla \phi^i |j| w_g$$

(42.8)

In this example, $N_r = 0$; $N_F = 1$ with $F_1 = q$ where $q$ is the Fourier heat flux, $q = -\kappa \nabla T$; and $m = 1$. As a side note, in the code the quantities $|j|$ and $w_g$ are treated as additional coefficients for convenience.

### 42.13.4 Vector_Source_Kernel_Expression

This Expression assembles a source term for a vector transport equation. Note that the MASS and SRC terms can both be cast in this form. The general form is:

$$-m \sum_{g=1}^{N_g} \left( \prod_{r=1}^{N_r} c_r(x_g) \right) \left( \sum_{s=1}^{N_s} f_s(x_g) \right) \phi^i(x_g) |j| w_g$$

(42.9)

Here, $m$ is a multiplier that defaults to 1, $N_g$ is the number of Gauss points in the quadrature rule, $N_c$ is the number of coefficients and $N_s$ is the number of sources provided. In this case, the sources $f_s$ are vector quantities. As an example, consider the term,

$$\int_{\Omega_c} \rho \frac{\partial v}{\partial t} |j| \phi^i |j| w_g$$

(42.10)

In this example, $N_1 = 2$ with $c_1 = \rho$, $N_s = 1$ with $f_1 = \frac{\partial v}{\partial t}$; and $m = -1$. As a side note, in the code the quantities $|j|$ and $w_g$ are treated as additional coefficients for convenience.

### 42.13.5 Vector_Advection_Kernel_Expression

This Expression assembles an advection term for a vector transport equation. The general form is:

$$m \sum_{g=1}^{N_g} \left( \prod_{r=1}^{N_r} c_r(x_g) \right) v_a \cdot \nabla V(x_g) \phi^i(x_g) |j| w_g$$

(42.11)
Here, $m$ is a multiplier that defaults to 1, $N_g$ is the number of Gauss points in the quadrature rule, $N_c$ is the number of coefficients, $v_a$ is a configurable advection velocity and $V$ is a vector field. As an example, consider the term,

$$
\int_{\Omega} \rho v \cdot \nabla v |j| \phi^i \text{d}\Omega = \sum_{g=1}^{N_g} \rho v \cdot \nabla v |j| \phi^i w_g
$$

(42.12)

In this example, $N_r = 1$ with $c_1 = \rho$; $V = v$; $v_a = v$; and $m = 1$. As a side note, in the code the quantities $|j|$ and $w_g$ are treated as additional coefficients for convenience.

### 42.13.6 Vector_Diffusion_Kernel_Expression

This Expression assembles a diffusion term for a vector transport equation. The general form is:

$$
m \sum_{g=1}^{N_g} \left( \prod_{r=1}^{N_r} c_r(x_g) \right) \left( \sum_{k=1}^{N_F} F_k(x_g) \right) : \nabla (e_j \phi^i(x)) |j| w_g
$$

(42.13)

Here, $m$ is a multiplier that defaults to 1, $N_g$ is the number of Gauss points in the quadrature rule, $N_c$ is the number of coefficients and $N_F$ is the number of flux models provided. As an example, consider the term,

$$
\int_{\Omega} T^t : \nabla (e_j \phi^i) |j| \text{d}\Omega = \sum_{g=1}^{N_g} T^t : \nabla (e_j \phi^i) |j| w_g
$$

(42.14)

In this example, $N_r = 0$; $N_F = 1$ with $F_1 = T$ where $T$ is the Newtonian stress, $T = \mu (\nabla v + \nabla v^t) - pI$; and $m = 1$. As a side note, in the code the quantities $|j|$ and $w_g$ are treated as additional coefficients for convenience.

### 42.14 Defining and Solving ODEs in Aria Plugins

This section covers how to define and solve an ODE or system of ODEs in an Aria plugin. The equations to solve must be of the form,

$$
\frac{d\bar{y}}{dt} = f(t, \bar{y})
$$

(42.15)

The ODE to solve must be declared as a subclass of the Aria ODE class, and must define at least the derivatives function, and it may also be useful to define the initialization and Jacobian functions:

```cpp
def derivatives(double t, const FArray<double,1>& y, FArray<double,1>& dydt)
def initialize(double t, const FArray<double,1>& y)
def jacobian(double t, const FArray<double,1>& y, FArray<double,2>& dfdy)
```

The initialize() function is called by the ODE solver at construction to set the initial values of $\bar{y}$, and the derivatives() function is called within the solver during solution. The default Jacobian function implements a numerical Jacobian, but it can be overwritten to allow an analytical one.

The currently available solvers are described in the Chemistry Solver Reference chapter. All solvers use adaptive sub-stepping to meet the requested tolerances. The solver stores the problem state $(t, \bar{y}, \frac{d\bar{y}}{dt})$ throughout solution as well as any intermediate states required by the particular solution algorithm. Any of the ODE solvers can be used to solve an ODE that inherits from the Aria ODE class, although for stiff ODEs an appropriate solver should be selected.
42.15 Defining and Using Global Variables

Occasionally it is useful to define a global variable that is available to the user for output to the log or results files, or to be used in user subroutines. These variables may be created using the Region::new_global_variable or Region::new_global_variable2 functions. A variety of value types (e.g. Real, Int, etc.) are supported, as are a number of reduction operations (MPI::Sum, MPI::Max, etc.). Region::new_global_variable() returns a GlobalVariable<Op, T> and is the preferred version. GlobalVariable<Op, T> stores a single value that is globally consistent once it is synchronized using the Region::synchronize_global_variables() call. Region::new_global_variable2() returns a GlobalVariable2<Op, T> which stores 2 values, one local and one global. There are currently some user plugins that access both the local and global values separately so the type is still available for compatibility, but we would prefer to not have both types in the future.

42.16 Errors and Warnings How-To

Editorial note: This How-To is mostly taken from Dave Bauer’s collection of How-Tos. Minor changes include formatting, editorial corrections and possibly additional information related to Aria.

42.16.1 Reporting

There are several types of warnings and exceptions that occur in sierra. Warnings are informational messages to the user which may effect the results, but which will allow the execution to complete. Dooms are error conditions which will allow the program to continue to a point, but will not allow it to complete. Often these are within the parser when you want the parser to continue to plod on ahead but not actually execute the code. Exceptions occur when all bets are off.

The Warning and Doom classes send their assembled message to the sierra reporter at destruction. The Exception classes send their assembled message to the sierra reporter within the catch block.

The warning and exception classes will all accept the put-to operator (<<) with plain old data. Since it is easy to put useful information to the user/developer in the message, please do so. For warnings directed to developers, it is recommended that the message be terminated with a

```cpp
<< std::endl << WarnTrace;
```

For exceptions, always use

```cpp
<< std::endl << StackTrace;
```

WarnTrace and StackTrace generate a "pretty" source file and line number message. If the message is purely informational for the user the std::endl and WarnTrace for the message should be omitted.

These are the include files:

```cpp
#include <util/Exception.h> // throw sierra
#include <util/ExceptionReport.h> // sierra reporter and
#include <util/RuntimeWarning.h> // throw RuntimeWarning
```

The RuntimeUserError class should be thrown in place of RuntimeError when it was the user’s fault that the program died. This generally means that the input deck has an error that a RuntimeDoomed, the
preferred method since it allows the parser, etc to continue, cannot handle since a seg fault or the like is imminent. It kills the output of the traceback since the user really does care and will only cause confusion.

Also, don’t put the WarnTrace or StackTrace to the RuntimeUserError.

The following code snippets serve as examples for the warning, doomed and exception conditions.

```cpp
class_tag *
Parser::prsr_handler_x(
    const Prsr_CommandValues & value)
{

    if (runtime_warning_condition)
        sierra::RuntimeWarning() << "My useful message about " <<
        some_data << std::endl << WarnTrace;

    if (runtime_exception_condition)
        throw sierra::RuntimeError() << "My useful message about " <<
        some_data << std::endl << StackTrace;

    if (runtime_exception_condition)
        throw sierra::RuntimeUserError() << "My useful message about "
        << some_data;

    if (parse_handler_warning_condition)
        sierra::RunWarning() << "My useful message about " <<
        some_data << std::endl << WarnTrace;

    if (parse_handler_doomed_condition)
        sierra::RuntimeDoomed() << "My useful message about " <<
        some_data << std::endl << WarnTrace;

}
```

### 42.16.2 Throttling a Specific Warning or Doom

If a particular warnings is going to be repeated countless times, you can request a unique message id from sierra using

```cpp
int message_id = sierra::get_next_message_id();
int message_id = sierra::get_next_message_id(int max_messages_displayed);
```

and pass message_id to the message constructor. This value is often stored within the function that generates the message using a static variable. This technique may not be suitable for all situations. Only a limited number of messages of each id are sent to the sierra reporter.

```cpp
if (runtime_warning_condition) {
    static message_id = get_next_message_id();
```
sierra::RuntimeWarning(message_id) << "My useful message about " << some_data << std::endl << WarnTrace;
}

if (parse_handler_warning_condition) {
    static message_id = get_next_message_id();
    sierra::Prsr::ParseWarning(value, message_id) << "My useful message about " << some_data << std::endl << WarnTrace;
}

if (parse_handler_doomed_condition) {
    static message_id = get_next_message_id();
    sierra::Prsr::ParseDoomed(value, message_id) << "My useful message about " << some_data << std::endl << WarnTrace;
}

if (parse_handler_exception_condition) {
    static message_id = get_next_message_id();
    sierra::Prsr::ParseError(value, message_id) << "My useful message about " << some_data << std::endl << WarnTrace;
}

42.16.3 Setting Output Throttles

There are several functions which can throttle the amount of data which is displayed.

To set the maximum number of warnings or dooms before an exception is thrown:

    void sierra::set_max_warnings(int max_warnings);
    void sierra::set_max_dooms(int max_dooms);
    int sierra::get_max_warnings();
    int sierra::get_max_dooms();

To set the maximum number of warnings displayed. Each occurrence is still counted, but not displayed. Dooms are always displayed.

    void sierra::set_max_displayed_warnings(int max_display_warnings);
    int sierra::get_max_displayed_warnings();

To set the default maximum id messages to display, set get_next_message_id() below:

    int sierra::set_default_max_id_display(int default_max_id_displayed)
    int sierra::set_default_max_id_display(int default_max_id_displayed)
42.16.4 Sierra Exception Reporter

What is this sierra reporter thing you ask? It goes like this: When a Warning or a Doom class is destroyed, report_exception() is called with the message. report_exception() then calls the registered report_exception_handler. The report_exception_handler has a signature of

\((\ast)(\text{const char } *\text{message, int type})\)

and is set with set_report_exception_handler(). Sierra sets this to use the sierra_report_exception_handler() which writes the message to sierra::Env::output().

If you want special decorations around your messages during execution, you can change the reporter. The parse, instantiation and commit handlers are set by run_sierra().

42.16.5 The Output versus OutputP0 Dilemma

Unfortunately, determining what to do with the output from multiple processors can be an issue. If a warning is going to happen on all processors, you may want to wrapper it with a test for processor zero and only output it there. The default report_exception_handler sends the output to Env::output().

42.16.6 Getting Counts

To get the number of warnings issued, even if not displayed:

\(\text{get_warning_count()}\)

To get the number of dooms issued, even if not displayed:

\(\text{get_doomed_count()}\)

42.16.7 Traceback and Tracing

The traceback messages are generated by the Trace and Traceback classes upon there destruction. So, if the source code has had trace objects constructed, the stack trace will show them. If efficiency becomes an issue, the \texttt{#define app TRACE ENABLED} can be undefined which will cause the Trace and Traceback classes to be empty.

The trace object can be used to extract function signature information. The getFunctionSpec(), getFunctionName(), getFunctionShortName(), getFunctionClass(), getFunctionShortClass() and getFunctionNamespace() functions will all extract appropriate information from the function signature.

The program /usr/netpub/traceString/traceString (on Linux) can be used to quickly place the trace objects in your source code. I suggest editing the results and placing \texttt{[ON], [TRACEBACK] or [NONE]} after the first \texttt{%TRACE} as in \texttt{%TRACE[NONE]}. These will instruct traceString which object type to create. Then, rerun traceString. I set it to \texttt{NONE} for functions which will execute many many times or cannot throw an error or not have much value in a stack trace (accessors, etc). I set it to \texttt{TRACEBACK} if it will not be useful to see traced during a trace run, but be useful in a traceback. And \texttt{ON} otherwise which incurs some overhead that enables the conditional runtime trace.

Here is the .traceString file I use and suggest be used throughout sierra.
42.16.8 Deriving from a Sierra Exception

When deriving a new exception from the framework exceptions and using the put-to (\<<\>) operator, you must include the following template functions in your class or

\[
\text{throw MyError()} \ll "\text{My error message cause of } \ll x;\]

will certainly fail. It only took me a day to relearn this, but:

If \text{MyError} does not implement a put-to operator, it uses the base classes put to operator, say \text{RuntimeError}. Well, the \text{RuntimeError} put-to operator returns a reference to a \text{RuntimeError}, so now you are going to be throwing a \text{RuntimeError} exception not a \text{MyError} exception. This is only an issue when using the temporaries. By putting the \text{MyError()} construction on the same line as the throw. I.e.,

\[
\text{MyError }\text{x; }\text{x }\ll "\text{My error message cause of } \ll x;\text{throw } x;\]

works fine since we are actually throwing \text{x}, a \text{MyError} object.

So, add these if you derive from a sierra exception and wish to throw the temporary object and use the put operator all in one pretty line:

\[
\text{class MyError : public RuntimeError { }\}
\]
template <class T>
inline MyError &operator<<(const T &t) {
    RuntimeError::operator<<(t);
    return *this;
}
};

42.16.9 path_name()

When generating error messages with object names, use the path_name() form rather than just name. This generates a dot (.) separated list of names from the Procedure on down.

42.16.10 abort() – Don’t use it

abort() does not provide any useful information when the application dies. By replacing abort calls with Warnings, Dooms and Exceptions, the code will be easier to maintain and the user will get better diagnostic output.

42.16.11 Apub_Parser_Base – Useful for parsing, not needed for error reporting

This class stashes useful information from within a parser that is likely to be needed after parsing. The runtime error reporting routines handle error reporting making the line number and command value information redundant for error reporting.

42.17 Diagnostic Writer How-To

Editorial note: This How-To is mostly taken from Dave Bauer’s collection of How-Tos. Minor changes include formatting, editorial corrections and possibly additional information related to Aria.

This document briefly describes the implementation of the diagnostic writer and masked based output. The diagnostic writer is intended to replace debug level output with but masked based diagnostic output. By utilizing the diagnostic writer, your normal output can be separated or interleaved with the diagnostic output, the diagnostic output can the enabled/disabled at specific times during a run, and entire classes can be output by simply putting the object to the diagnostic writer.

42.17.1 Output

Application output falls into three classes. Normal execution output, warning and error output, and diagnostic output. Normal execution output is handled via the Env::outputP0() and Env::output() functions and by the application diagnostic writer. Warning and error output is handled by the RuntimeWarning, RuntimeDoomed and runtime exception classes (RuntimeError, LogicError, etc), which is eventually written to the env output stream and the diagnostic output stream. And diagnostic output is for selected operationally descriptive output.

In many cases, the diagnostic output is utilized as the primary output stream when selectable level of user output is desired. With that in mind, an InfoWriter class has been written. However, additional discussion
is required to complete the design and implementation of this class for primary application output.

### 42.17.2 Diagnostic Writer

The diagnostic writer allows you to write diagnostic information to the diagnostics stream by specifying the content from the command line or the input deck. It a debug level built on a bit mask.

Since there are many applications and libraries, there are several diagnostic writers. Each diagnostic writer has its own bit mask, command line parser and writer. However, they all share a common diagnostic stream. So, output from each diagnostic writer is properly interleaved.

Each application defines its own diagnostic writer. This is generally defined within the `app_DiagWriter_fwd.h`, `app_DiagWriter.h`, `app_DiagWriter.C` files. The `app_DiagWriter_fwd.h` file defines the LOG_xxx bit assignments. These values are used to specify the type of message to be written. The `app_DiagWriter.h` rarely needs to be modified. It declares the diagnostic writer for the application or library. The `app_DiagWriter.C` files defines the parser which provides names for the LOG_xxx bit masks.

### 42.17.3 Using The Diagnostic Writer

To have your program send output whenever a specified log bit is set, add the following to your code:

```cpp
dwout.m(LOG_xxx) << "description, " << value << std::endl;
```

or, if much computation or MPI is involved:

```cpp
if (dwout.shouldPrint(LOG_xxx)) {
    dout << "really_spendy_function(), " <<
        really_spendy_function() << std::endl;
}
```

Where `dwout` is the name of the diagnostic writer, LOG_xxx is the bit which describes the type of message, description is a description of the data, value is the value of interest. And, std::endl ends the message.

Note that nearly all (please let me know what’s missing) containers have output operators. So to write an entire vector, just write the vector variable, the diagnostic writer will take care of the rest.

### 42.17.4 Turning on the LOG_xxx Bits

Each application has its own command line option and line commands for flipping on the bits. The command line option has limited functionality in that the parameters cannot be switched on and off during an application execution. However, it is useful for a quick look.

For sierra framework, the command line option is `-m`, and each application has its own option name. Use the `-h` option to display a table of command line options and parameters.

The Diagnostic Control command block in the sierra block controls each the diagnostic writers:

```bash
Begin Diagnostic Control <diagwriter>
    From time t0 to t1 enable <parameters>
    From step s0 to s1 enable <parameters>
```

1096
<table>
<thead>
<tr>
<th>Option</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>error</td>
<td>Display error messages</td>
</tr>
<tr>
<td>warning</td>
<td>Display warning messages</td>
</tr>
<tr>
<td>members</td>
<td>Display data structure members</td>
</tr>
<tr>
<td>timer</td>
<td>Display execution time during trace</td>
</tr>
<tr>
<td>trace</td>
<td>Display execution trace</td>
</tr>
<tr>
<td>contact</td>
<td>Display contact diagnostic information</td>
</tr>
<tr>
<td>geometry</td>
<td>Display geometry diagnostic information</td>
</tr>
<tr>
<td>scontrol</td>
<td>Display solver control diagnostic information</td>
</tr>
<tr>
<td>search</td>
<td>Display search diagnostic information</td>
</tr>
<tr>
<td>transfer</td>
<td>Display transfer diagnostic information</td>
</tr>
<tr>
<td>parser</td>
<td>Display parser diagnostic information</td>
</tr>
<tr>
<td>parameters</td>
<td>Display parameter diagnostic information</td>
</tr>
<tr>
<td>io</td>
<td>Display contact I/O information</td>
</tr>
<tr>
<td>plugins</td>
<td>Display user function and plugin diagnostic information</td>
</tr>
</tbody>
</table>

Table 42.1. Diagnostic writer options for Framework (fmwkout).

```
On condition c enable <parameters>
Enable <parameters>
End Diagnostic Control <diagwriter>
```

Please refer to Diagnostic Control for implementation details.

The diagnostic output can be selectively enabled based on time, step or an application specified condition. During the application's procedure execution loop, the diagnostic controller evaluates the enclosed line commands in the order specified in the input deck. The diagnostic options specified in the first line command that meets its criteria are applied.

Since control parameters are only applied when the criteria is met, it is important to include an ENABLE line command with the base settings to be applied as a baseline.

Table 42.17.4 lists some of the options available for the framework diagnostic writer fmkout and options for the Aria diagnostic writer arialog are given in table 42.17.4. Other application will likely implement additional diagnostic writers.

### 42.17.5 Diagnostic Stream

The diagnostic stream specified the output destination for all the active diagnostic writers. The stream allows the output from the diagnostic writers to be interleaved.

The command line option

```
-dout <destination>
```

and the sierra block line command

```
diagnostic stream <destination>
```

determines the output. <destination> may be cout, cerr, outputp0, output or a path. If a path is given,
<table>
<thead>
<tr>
<th>Option</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>bc</td>
<td>Display boundary condition information</td>
</tr>
<tr>
<td>debug</td>
<td>Display extra debugging information</td>
</tr>
<tr>
<td>eq</td>
<td>Display equation information</td>
</tr>
<tr>
<td>expression</td>
<td>Display Expression information</td>
</tr>
<tr>
<td>hadapt</td>
<td>Display h-adaptivity information</td>
</tr>
<tr>
<td>linsolve</td>
<td>Display linear solver information</td>
</tr>
<tr>
<td>nonlinear</td>
<td>Display nonlinear solver information</td>
</tr>
<tr>
<td>pp</td>
<td>Display postprocessor information</td>
</tr>
<tr>
<td>sens_check</td>
<td>Enable the Expression Newton sensitivity checker</td>
</tr>
<tr>
<td>species</td>
<td>Display species information</td>
</tr>
<tr>
<td>transfer</td>
<td>Display transfer related information</td>
</tr>
<tr>
<td>plugin</td>
<td>Display plugin information</td>
</tr>
</tbody>
</table>

Table 42.2. Diagnostic writer options for Aria (arialog).

each processor creates its own file suffixing the path with .n.p where n is the number of processors and p is the rank of each processor. See Diagnostic Stream for specifying the output destination.

42.17.6 Coding Objects

To code your own objects to play with the diagnostic writer, you only need to add a verbose_print() member function to your class. And, a operator<<() function to your namespace. Naturally the implementation for verbose_print() is generally in the .C file, not the header.

```cpp
class MyClass : public ParentClass
{
public:
    DiagWriter &verbose_print(DiagWriter &dout) const {
        if (dout.shouldPrint()) {
            dout << "MyClass " << m_name << push << std::endl;
            ParentClass::verbose_print(dout).std::endl();
            dout << "m_var1, " << m_var1 << std::endl;
            dout << "m_var2, " << m_var2 << std::endl;
            dout << "m_ptr1, " << c_ptr(m_ptr1) << std::endl;
            dout << "m_ptr2, " << c_ptr_name(m_ptr2) << std::endl;
            dout << pop;
        }
        return dout;
    }
};

inline DiagWriter &operator<<(DiagWriter &dout, const MyClass &my_class) {
    return my_class.verbose_print(dout);
}
```

If your class is polymorphic, be sure to define verbose_print() as virtual.
When writing a subclass from your class, the put-to operator (\( \bullet \)) will by do what you expect, even when you cast. So you need to use the direct call form.

### 42.17.7 Writing Containers

You can write an STL container by simple putting it to the diagnostic writer. This is implemented using templates in `utility/include/DiagWriter.h`.

### 42.17.8 Writing Pointers

Writing of pointers is usually quite ugly since you need to check if the pointer is null first. Instead, use the `c_ptr()` function. If the pointer is null it writes "(pointer), <not created>". Or, you can use `c_ptr_name()` function which will call the `name()` function of the pointed to object if the pointer is not null.

You can have your own pointed object function called by replacing name with your member function name below.

```cpp
template <class T>
inline c_ptr_func_<T, const String &> c_ptr_name(const T *t) {
    return c_ptr_func_<T, const String &>(t, &T::name);
}
```

### 42.17.9 Diagnostic Control

The diagnostic control block in the sierra command block is handled by the `Fmwk::DiagControl::Control` class. Simply add

```cpp
Fmwk::DiagControl::Control diag_control(step_cntr(),
    time(Fmwk::STATE_OLD), 0);
```

at the beginning of your main procedure control loop. This line exists in the Solver Control procedure.

### 42.18 Timers and Timing How-To

*Editorial note: This How-To is mostly taken from Dave Bauer’s collection of How-Tos. Minor changes include formatting, editorial corrections and possibly additional information related to Aria.*

This document briefly describes the time metrics collection features available to sierra applications.

The `DiagTimer` and `Timer` classes provide runtime metric information for properly rigged objects.

The system has a root "System" timer. This timer is started when `run_sierra()` is called and stopped before the successful completion information is displayed.

### 42.18.1 DiagTimer and Timer

The `DiagTimer` class implements the basic developer interface to the timers. Generally, the framework form, `Timer`, will be used to implement timers within a framework derived class.
Timers are intended to be members of your classes or static objects created within a function or member function. Each timer has a name, a parent and a timer class. The name is used to find a child timer of a parent and to display the collected metrics. The parent is used to build the hierarchy of metrics gathered in an application. And the timer class or type categorizes the timers so they may be enabled, disabled and selected for display.

The timing information is actually maintained in a separate tree. So, during timer construction within a class, the timer is a reference to the real timing metric information in the tree. This design means that timers are not destroyed when an object is destroyed.

The timers are hierarchical by name. So, by giving a timer a unique name among its siblings, each object has its own timer and its children timers are also unique.

42.18.2 Add a Timer to a Class

To add a timer to a class, include the header file and add the timer as a member of the class. Then, during construction initialization, specify the timer’s name and parent timer. You can also specify a timer class if it is different from the parents.

Then, to start/stop the timer, use the TimeBlock and TimeBlockSynchronized to ensure that a started timer is stopped.

```cpp
#include <util/Timer.h>

class MyClass
{
  public:
    // etc.

    Timer &getMyTimer() {
      return m_myTimer;
    }

    Timer &getMySubTimer() {
      return m_mySubTimer;
    }

  private:

    Timer m_myTimer;
    Timer m_mySubTimer;
    // etc.
};

MyClass::MyClass(
  Region & region)
: m_myTimer("My Timer", region.getRegionTimer()),
  m_mySubTimer("My SubTimer", m_myTimer)
{
  // etc.
}
```
When controlling timers using the TimeBlocks, be sure to give the object a name. Some compilers destroy unnamed objects immediately, other destroy them at the end of the block. I usually use timer__

```cpp
MyClass::someFunction()
{
    Timer::TimeBlock timer__(m_myTimer); // Metrics in block are collected to m_myTimer;
}
```

```cpp
MyClass::someFunction()
{
    Timer::TimeBlockSynchronized timer__(m_myTimer);
    // MPI_Barrier then start timer
}
```

```cpp
MyClass::someOtherFunction()
{
    m_myTimer.start(); // Works, but is dangerous if stop() is not called.
    m_myTimer.stop();
}
```

### 42.18.3 Adding a Timer to a Function

To add a timer to a function, create a static Timer in the function, then start/stop it using the Timer::TimeBlock. Note that if there is no parent specified, the root timer "System" will be the timer’s parent.

```cpp
int myFunction()
{
    static Timer my_timer("My Timer");
    Timer::TimerBlock(my_timer);
    // etc.
}
```

### 42.18.4 Getting Information from a Timer

Each timer has an accumulation time/count, a checkpoint time/count and a recent lap time/count. The accumulation time is the overall time/count since the start of the application. The checkpoint time/count records the current values when set, then the difference from that time/count can be obtained. This is useful for displaying delta times for iterations. The lap time is the time accumulated during the last start()/stop() cycle for the timer.

The metrics available are `getCPUTime()`, `getWallTime()`, `getFlopCount()`, `getIOCount()`, `getMsgCount()`. Flop, IO, and Msg are not implemented on most platforms (any really).

```cpp
m_myTimer.getCpuTime().getStart(); // Timer most recent start time
```
m_myTimer.getCpuTime().getStop(); // Timer most recent stop time
m_myTimer.getCpuTime().getLap(); // Most recent stop - start
m_myTimer.getCpuTime().getTotal(); // Accumulated time
m_myTimer.getCpuTime().getTotal(false); // Accumulated time less checkpoint time

I could add getCheckpoint() and getCheckpointTotal() as variations on a theme, but...

### 42.18.5 Displaying the Timers

To display a table of the collected timing information, use the `Timer::printTable()` function. The function lets you specify the classes and the metrics to output. Note that only enabled timers and metrics are displayed. Use `TIMER_CHECKPOINT` to display checkpointed time. It also resets the checkpoint time/count after display.

```cpp
class Timer {
    // ... implementation ...

    static void printTable(int mask);
};
```

```cpp
std::cout << Timer::printTable(TIMER_CPU | TIMER_ALL);
```

### 42.18.6 Enabling/Disabling the Timers

Timers are enabled by timer class. They can be enabled using the `Timer::setEnabledTimers()` function, by the `-timer` command line option or by the `ENABLE TIMER` input deck line command.

### 42.18.7 Timers in the Framework

The framework creates several timers and starts/stops them when it has control of the operations.

- header timers are given the name associated with the name of the object automatic timers are handled within framework and required no additional coding otherwise, instructions are given on how to collect the data for the timer

#### Domain:

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Domain</td>
<td>header</td>
</tr>
<tr>
<td>LinearSystem</td>
<td>automatic</td>
</tr>
<tr>
<td>Initialize</td>
<td>automatic</td>
</tr>
<tr>
<td>Execute</td>
<td>automatic</td>
</tr>
<tr>
<td>Load</td>
<td>automatic</td>
</tr>
<tr>
<td>Solve</td>
<td>automatic</td>
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#### Procedure:

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<td>Restart</td>
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<td>Execute</td>
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#### Region:

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<tr>
<td>Region</td>
<td>header</td>
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<tr>
<td>Initialize</td>
<td>call Timer::TimeBlock timer__(getRegionInitializeTimer()); at beginning of initialization</td>
</tr>
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</table>

**1102**
Execute call Timer::TimeBlock timer__(getRegionExecuteTimer()); at beginning of execute

Mechanics:
  Mechanics header

Algorithm:
  Algorithm header
  Apply automatic

WorksetAlgorithm: (sub class of Algorithm)
  Gather automatic
  ScatterAssemble automatic

NonLinearCoupler:
  NonLinearSolver automatic
  Initialize automatic
  Scatter automatic
  Solve automatic

NonLinearSolver:
  NonLinearSolve automatic
  Initialize automatic
  Scatter automatic
  Solve automatic

UserInputFunction:
  UserInputFunction automatic
Glossary

coefficient: Each field that is represented by a basis function expansion has a set of coefficients that are used in that expansion. For example, the three dimensional velocity vector represented by an eight node, tri-linear hex element has eight coefficients. In this example, each coefficient is also a vector with three components.

This is consistent with the Sierra data model wherein vectors and tensors are single-entity data types. Although for certain basis function representations the coefficients may be the exact value of the field at a point this is not the case in general.

component: The number of values required to describe a field at a point. Equal to the tensor dimension raised to the power of the tensor order. For example, temperature has one component, velocity has 3 components (in 3 dimensions) and stress has 9 components.

dof: An entry in the vector of unknowns, i.e., in the linear solver solution vector.

field: The physical variable of interest, e.g., temperature or velocity.

multidof: A field is a multidof if it contains more than one component
References


[3] The SNTools Project. SNTools SourceForge Project. Online. 2.3


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