Report on Accelerated Corrosion Studies of Electrical Components

S. Jill Glass, Curtis D. Mowry, and N. Robert Sorensen

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

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Report on Accelerated Corrosion Studies

S. Jill Glass, Curtis D. Mowry, and N. Rob Sorensen
Sandia National Laboratories
P.O. Box 5800
Albuquerque, New Mexico  87185-MS0886

Abstract

Sandia National Laboratories (SNL) conducted accelerated atmospheric corrosion testing for the U.S. Consumer Product Safety Commission (CPSC) to help further the understanding of the development of corrosion products on conductor materials in household electrical components exposed to environmental conditions representative of homes constructed with problem drywall. The conditions of the accelerated testing were chosen to produce corrosion product growth that would be consistent with long-term exposure to environments containing humidity and parts per billion (ppb) levels of hydrogen sulfide (H$_2$S) that are thought to have been the source of corrosion in electrical components from affected homes. This report documents the test set-up, monitoring of electrical performance of powered electrical components during the exposure, and the materials characterization conducted on wires, screws, and contact plates from selected electrical components. No degradation in electrical performance (measured via voltage drop) was measured during the course of the 8-week exposure, which was approximately equivalent to 40 years of exposure in a light industrial environment. Analyses show that corrosion products consisting of various phases of copper sulfide, copper sulfate, and copper oxide are found on exposed surfaces of the conductor materials including wires, screws, and contact plates. The morphology and the thickness of the corrosion products showed a range of character. In some of the copper wires that were observed, corrosion product had flaked or spalled off the surface, exposing fresh metal to the reaction with the contaminant gasses; however, there was no significant change in the wire cross-sectional area.
ACKNOWLEDGMENTS

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<tr>
<th>Acronym</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>AES</td>
<td>Auger–Electron Spectroscopy</td>
</tr>
<tr>
<td>AFCI</td>
<td>arc-fault circuit interrupter</td>
</tr>
<tr>
<td>Al</td>
<td>aluminum</td>
</tr>
<tr>
<td>Au</td>
<td>gold</td>
</tr>
<tr>
<td>CPSC</td>
<td>Consumer Product Safety Commission</td>
</tr>
<tr>
<td>Cr</td>
<td>chromium</td>
</tr>
<tr>
<td>ct</td>
<td>contact tab</td>
</tr>
<tr>
<td>Cu</td>
<td>copper</td>
</tr>
<tr>
<td>Cu2O</td>
<td>cuprite, copper oxide</td>
</tr>
<tr>
<td>CuS</td>
<td>covellite</td>
</tr>
<tr>
<td>CuSO4</td>
<td>chalcocyanite</td>
</tr>
<tr>
<td>Cu3(SO4)(OH)4</td>
<td>antlerite</td>
</tr>
<tr>
<td>CuO5S5</td>
<td>digenite, copper sulfide</td>
</tr>
<tr>
<td>(Cu2Zn)3(SO4)_{6.6}(H2O)</td>
<td>ktenasite</td>
</tr>
<tr>
<td>EDS</td>
<td>energy dispersive spectroscopy</td>
</tr>
<tr>
<td>FACT</td>
<td>Facility for Atmospheric Corrosion Testing</td>
</tr>
<tr>
<td>Fe</td>
<td>iron</td>
</tr>
<tr>
<td>FIB</td>
<td>focused ion beam</td>
</tr>
<tr>
<td>GFCI</td>
<td>ground fault circuit interrupter</td>
</tr>
<tr>
<td>GS</td>
<td>ground screw</td>
</tr>
<tr>
<td>GW</td>
<td>ground wire</td>
</tr>
<tr>
<td>HSP</td>
<td>hot screw plate</td>
</tr>
<tr>
<td>HS</td>
<td>hot screw</td>
</tr>
<tr>
<td>HW</td>
<td>hot wire</td>
</tr>
<tr>
<td>l</td>
<td>left</td>
</tr>
<tr>
<td>lpm</td>
<td>liters per minute</td>
</tr>
<tr>
<td>lcp</td>
<td>left contact plate</td>
</tr>
<tr>
<td>MFG</td>
<td>mixed flowing gas</td>
</tr>
<tr>
<td>m/z</td>
<td>mass to charge ratio (in mass spectrometry)</td>
</tr>
<tr>
<td>micrometer</td>
<td>1 millionth of a meter or 1x10^{-6} meters</td>
</tr>
<tr>
<td>micron (µm)</td>
<td>1 millionth of a meter or 1x10^{-6} meters</td>
</tr>
<tr>
<td>mm</td>
<td>millimeter</td>
</tr>
<tr>
<td>m-ohm</td>
<td>milli-ohm or one thousandth of an ohm</td>
</tr>
<tr>
<td>Ni</td>
<td>nickel</td>
</tr>
<tr>
<td>NS</td>
<td>neutral screw</td>
</tr>
<tr>
<td>O</td>
<td>oxygen</td>
</tr>
<tr>
<td>PIHW</td>
<td>push-in hot wire</td>
</tr>
<tr>
<td>ppb</td>
<td>parts per billion</td>
</tr>
<tr>
<td>ppm</td>
<td>parts per million</td>
</tr>
<tr>
<td>Pt</td>
<td>platinum</td>
</tr>
<tr>
<td>R</td>
<td>receptacle</td>
</tr>
<tr>
<td>r</td>
<td>right</td>
</tr>
<tr>
<td>rcp</td>
<td>right contact plate</td>
</tr>
<tr>
<td>S</td>
<td>sulfur</td>
</tr>
<tr>
<td>Acronym</td>
<td>Description</td>
</tr>
<tr>
<td>---------</td>
<td>-------------</td>
</tr>
<tr>
<td>SEM</td>
<td>scanning electron microscopy</td>
</tr>
<tr>
<td>SNL</td>
<td>Sandia National Laboratories</td>
</tr>
<tr>
<td>Ti</td>
<td>titanium</td>
</tr>
<tr>
<td>XRD</td>
<td>X-ray Diffraction</td>
</tr>
<tr>
<td>XRF</td>
<td>X-ray Fluorescence</td>
</tr>
<tr>
<td>Z</td>
<td>atomic number of element from the periodic table (e.g., Z for copper is 29)</td>
</tr>
<tr>
<td>Zn</td>
<td>zinc</td>
</tr>
</tbody>
</table>
1. INTRODUCTION

In September 2009, Sandia National Laboratories (SNL) was tasked by the U.S. Consumer Product Safety Commission (CPSC) staff with identifying the extent and nature of corrosion on conductor subcomponents of residential electrical components harvested from homes in several states. Residential electrical components, including receptacles, switches, ground-fault circuit-interrupters (GFCIs), arc-fault circuit interrupters (AFCIs), and circuit breakers were provided to SNL for analyses. That study examined electrical components that had been installed in Florida and Virginia homes for a period of two or more years prior to their removal by the CPSC. A set of six receptacles was the focus of Sandia’s study of harvested electrical components. Corrosion was observed on electrical conductors (wires, screws, and contact plates) from receptacles harvested from six homes. The major findings of that study were reported and are summarized as follows:1

- Wires showed the greatest degree of corrosion, with some areas showing a continuous layer of corrosion product.
- Scanning Electron Microscopy / Energy Dispersive Spectroscopy (SEM/EDS) elemental analyses showed that the corrosion product, which was up to 20 microns thick in samples analyzed, to date, consisted primarily of copper and sulfur.
- SEM analyses of FIB cross sectioned ground wires also showed that there was localized corrosion of the base copper that produced pits. This is consistent with our understanding of the copper sulfidation process. As the copper sulfide film forms, copper from the metal below the film migrates through the film, is sulfidized at the surface, and results in growth of the film. Thus, the voids in the copper surface are the result of copper migration during the sulfidation process. In some cases, the voids contained a spongy looking material. The voids observed, to date, were up to 20 microns in depth.
- X-ray diffraction (XRD) analyses identified Cu₂S₅ (digenite) and Cu₂O (cuprite) as the two major constituents of the surface corrosion layer on the ground wire.
- Similar results in terms of corrosion product thickness and composition were found in a parallel study conducted by the National Institute of Standards and Technology (NIST) on Cu hardware used in HVAC, gas distribution, and fire safety components.2
- Screws and contact plates also showed evidence of copper sulfide corrosion products, but observations showed that it was a thinner layer than that observed on wires.
- Wire insulation and coverage by other metallic surfaces provide some degree of protection against corrosion. Screw and contact plate surfaces that were in contact with other conductors also showed minimal or no corrosion compared to exposed conductor surfaces.

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The 2009 study was able to identify the nature and extent of the corrosion product that had occurred during the exposure to problem drywall in the six homes, but not how much corrosion could occur for the typical lifetime of household electrical components, and what effects this might have on the electrical function and safety of powered circuits. To answer these questions, SNL designed and conducted the experimental study described in this report, in which new electrical components, supplied by CPSC, were exposed to elevated temperature and humidity, and contaminant gasses, including H2S, NO2, and Cl2, in Sandia’s larger Facility for Atmospheric Corrosion Testing or FACTII. The conditions of the test and the duration of the exposure were chosen to be representative of a Battelle Class IV\(^3\) environment to stimulate the growth of corrosion product that would be expected to be generated by a 40-year exposure to the conditions that produced the corrosion product thickness measured on harvested components in the 2009 study.

Three powered electrical circuits, consisting of switches, receptacles (conventional and GFCI), and load centers were cycled electrically over the 8-week exposure in the FACTII environmentally controlled chamber. The total cycle time was 20 minutes, with 10 minutes on, followed by 10 minutes off. The purpose of the cycling was to provide thermally-driven stresses of the corrosion product (due to differences in the thermal expansion properties of the metal conductors and the corrosion products). Furthermore, assessment of corrosion-induced functional impacts was performed while components were undergoing exposure. At the end of the exposure, the electrical components were subjected to the same materials analyses used for the characterization of harvested electrical components from the 2009 study, including SEM, XRD, and focused ion beam (FIB). These analyses were used to determine the morphology, thickness, and chemical identity of the observed corrosion layers.

2. **EXPERIMENTAL DETAILS**

This experiment was designed to test multiple components of a powered electrical system in an environmental chamber in which Battelle Class IV conditions were maintained. The selection of the Class IV environment was based on results from components harvested from several homes. The components were analyzed, and a corrosion product thickness was determined. The exposure was estimated to be three years. Figure 1 presents the data from the harvested components and compares the result to published mixed flowing gas testing. Based on these data, the best fit corrosion class appears to be Class IV, the most extreme corrosive conditions of the Battelle exposure system.

2.1 Chamber and Conditions

The Battelle Class IV environment consists of exposure to a mixture of gases (H$_2$S, NO$_2$, and Cl$_2$), and elevated humidity. The specific conditions are:

- 200 ppb H$_2$S
- 200 ppb NO$_2$
- 50 ppb Cl$_2$
- 75% relative humidity (RH)
- 50 degrees (celsius)

The pollutant gasses used in the Battelle mixed flowing gas test are based on empirical correlations between field exposures and accelerated test exposures. That is, laboratory tests were selected based upon their ability to reproduce corrosion products seen in the field. There is no fundamental basis for the type or concentration of the pollutant gasses, so extrapolation to other conditions, or additional failure mechanisms is not possible. In the Class IV test, H$_2$S is the primary corrosive species, and NO$_2$ and Cl$_2$ are present primarily as accelerants. The elevated temperature also provides an acceleration factor. The primary corrosion product produced in a Class IV test is copper sulfide. The pollutant gasses were supplied using permeation tubes, and mass flow controllers were used to maintain flow. The chamber used is a 300 liter chamber, and the total flow rate was set at 12 liters/minute (lpm). The exposure was for a period of eight weeks, which roughly represents a field exposure in a light industrial environment of 40 years, according to the Battelle Class IV conditions. Due to the large area of exposed reactive metal (primarily copper), the system was not able to maintain the desired concentration of 200 ppb.
H$_2$S. To compensate for the decreased concentration, the test time was extended to maintain the equivalent exposure in ppb-hrs (concentration in ppb times time in hours). A schematic of the exposure system is shown in Figure 2. The three sets of powered electrical components were set up in the FACTII reaction chamber.

![Figure 2. FACT II schematic](image)

### 2.2 Component Hardware Setup/Wiring

A schematic of the entire electrical system contained within the environmental chamber is shown in Figure 3. Three separate electrical circuits were tested in the FACT at the same time. For each circuit, the electrical components were broken into two categories. The first included components found in the electrical service panel (circuit breakers). Due to size constraints, and to ensure access of the pollutant gas to the circuit breakers, the service panels did not include their steel enclosures. The mounts were attached directly to acrylic plates. Hardware consisted of a load center panel, one standard circuit breaker, and one arc-fault circuit interrupter (AFCI) breaker. Each of the three circuits contained unique hardware from separate manufacturers. The two circuit breakers were wired in series. The output from the circuit breaker provided the input to the AFCI. The neutral lead from the AFCI was connected to the neutral lead of the input. A schematic of the load center circuit is shown in Figure 4, as well as a photograph in Figure 5.
Figure 3. Schematic of electrical system (complete system). Each circuit had an electrical load of 12 A (via three 500W halogen lamps).

Figure 4. Schematic of load center circuit. Three separate load centers were wired in parallel. Each load center contains a circuit breaker, an arc-fault circuit breaker, and a junction bar. The two circuit breakers are wired in series.
The second group of electrical components was comprised of electrical devices that are typically found throughout the home. Each of the three circuits contained one rocker switch, one toggle switch, one standard duplex receptacle, and one GFCI receptacle. As with the load center hardware, three sets of unique devices were included (from various manufacturers), with no duplication of the components. A schematic of this circuit is shown in Figure 6, and a photograph of the circuit is shown in Figure 7. All four components were connected in series-parallel. This entire circuit was mounted in an acrylic box with a fan on the end to draw chamber gas into the box and through the individual components in order to approximate the manner in which gases would flow around and through components when mounted in consumer residences. All of the wires were copper. The contacts in the electrical devices contained a variety of metal, and were not all analyzed for composition.
Copper witness coupons provided by Sandia were degreased and then pickled in 10 percent HCl and rinsed in flowing de-ionized water to provide a reproducible surface. The coupons were then placed inside the chamber for later analysis. Coupons were removed and weighed at the conclusion of the test.

2.3 Disassembly Procedure
At the conclusion of the 8-week exposure in a Battelle Class IV environment, all three circuits were removed from the chamber. They were photographed while still electrically connected (unpowered). The inlet and outlet wires were cut to remove the circuit from the chamber. Twist-on connectors remained in place. The system was then disassembled according to the following procedure:

- Label all components (switches, circuit breakers, receptacles) to identify circuit used for testing.
- Photograph each system as-is, using digital camera.
- Remove the lid from the box containing the components, and photograph the back side (digital camera).
- Clip the wires near each of the components. Leave a pigtail attached to the components. Bag all of the wires (with twist on connectors) in Ziploc® bags (1 bag for each circuit). Label wires with circuit and hookup locations (e.g., 1RS to 1TS).
- Remove receptacles and switches from system.
- Photograph areas of interest (wires, attachment points, case, etc.) using low power optical capabilities.
- Remove circuit breakers from system.
- Photograph screw connections and bus (clip) connections using microscope.
- Determine twist on connectors to be analyzed and clip them from the wiring.
- Photograph twist on connectors using microscope.
- Assemble hardware and images and determine materials analyses to be performed.
  - SEM
  - XRD
  - FIB sectioning and analyses
Metallurgical cross section for SEM or other analyses

A sample identification convention was adopted to keep track of all of the parts to be analyzed for corrosion. The convention used was “Circuit-Component-Location-Sequential Number.” For example, the neutral wire from the GFCI receptacle in circuit 2 is identified as “2-GR-NW-2.” Table 1 gives the complete listing of the parts. This identification was used throughout the analyses performed, including optical microscopy, SEM, FIB, cross section, and XRD.

Table 1. Naming Convention for Parts Exposed to Class IV Environment

<table>
<thead>
<tr>
<th>Circuit</th>
<th>Component</th>
<th>Location</th>
<th>Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>RS Rocker Switch</td>
<td>HW Hot wire</td>
<td>1</td>
</tr>
<tr>
<td>2</td>
<td>TS Toggle Switch</td>
<td>NW Neutral wire</td>
<td>2</td>
</tr>
<tr>
<td>3</td>
<td>GR GFCI Receptacle</td>
<td>GW Ground wire</td>
<td>3</td>
</tr>
<tr>
<td>R Receptacle</td>
<td>HS Hot screw</td>
<td></td>
<td>...</td>
</tr>
<tr>
<td>ACB</td>
<td>AFCI circuit breaker</td>
<td>NS Neutral Screw</td>
<td></td>
</tr>
<tr>
<td>CB</td>
<td>Circuit breaker</td>
<td>GS Ground Screw</td>
<td></td>
</tr>
<tr>
<td>WN</td>
<td>Twist on connector</td>
<td>HSP Hot screw plate</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>NSP Neutral screw plate</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>GSP Ground screw plate</td>
<td></td>
</tr>
</tbody>
</table>
2.4 Sampling Plan

Once the parts were removed from the chamber and optically examined and photographed, specific components were selected for various analyses. In some cases, the selection was random, while in others, specific features observed on the surfaces drove the types of analyses to be performed, as shown below. Details of the instrumentation follow in the next section.

**XRD analysis**
- 2-R-GR-GW  Wire between the receptacle and the GFCI, unattached copper ground wire circuit
- 1-plug  Connector plug from circuit 1 that was in receptacle
- Copper coupon from chamber, randomly selected

**Photos only**
- 1-RS  Rocker switch
- 2-RS  Rocker switch
- 3-RS  Rocker switch
- 3-TS  Toggle switch
- 2-out  Yellow twist on connector

**SEM, cross section**
- 1-RS-HSP2  Rocker switch, hot screw plate
- 2-RS-HW1  Rocker switch, hot wire
- 1-GR-HW1  GFCI, hot wire–do 2 cross sections (by insulation, by “teeth” marks)
- 2-out  Red twist on connector

**SEM, corrosion morphology plan view**
- 1-R-HW1  Receptacle, hot wire
- 3-RS-HW1  Rocker switch, hot wire
- 3-TS-HW1  Toggle switch hot wire–pull back insulation (hot wire)
- 2-OUT  Long neutral connecting wire–sample in middle of long insulated length
- House  Sample from harvested parts–sample in middle of long insulated length

**Optical photos, then possible SEM morphology**
- 3-R-PIHW  Receptacle, push-in hot wire
- 3-R-PINW  Receptacle, push-in neutral wire
- 3-RS-PIHW  Rocker switch, push-in hot wire
- 2-RS-PIHW  Rocker switch, push-in hot wire
- 1-GR-GW  GFCI circuit breaker ground wire

**Dissassemble, determine analysis path based on observations**
- 1-ACB  AFCI circuit breaker
- 1-GR  GFCI receptacle
- 3-TS  Toggle switch
- 3-RS  Rocker switch
2.5 Materials Characterization Instrumentation and Methods

Corrosion products were analyzed using several materials characterization techniques. The following paragraphs provide basic details on the instrumentation and pertinent information related to the specific techniques.

Two scanning electron microscopes (SEM) (Hitachi S4500 and Zeiss Supra 55VP) with X-ray energy dispersive spectroscopy (EDS) capabilities were used. The SEM can produce high magnification images of parts and/or materials. Prior to analysis, samples sometimes were coated with a thin layer of platinum (Pt), gold-palladium (Au-Pd), or carbon (C) to enhance imaging and prevent charging. The EDS data collected here provides information about which elements are present at or near the surface. Qualitative information about differences in the relative amounts of various elements can also be obtained. Samples used for plan view SEM imaging and analysis were affixed to an aluminum stub with conductive carbon tape. Subcomponents or wires submitted for SEM cross-sectional imaging and analysis were mounted in epoxy, and then cut, ground, and polished, to provide a smooth surface, including a cutaway of the subcomponent.

A focused ion beam (FIB) instrument (FEI Helios Nanolab) was used to generate local cross sections, as well as remove material from the surface of interest, in order to measure corrosion layer thickness. A platinum coating was used to protect the top surface of the sample during the FIB sectioning process. The exposed cross sections then were analyzed using SEM techniques including analysis by Hyperspectral Imaging, which allows the entire elemental spectrum at each point to be measured.

The µXRD (microsampling X-ray diffraction) technique allows the phase identification to be made for the material in a region of interest. Micro XRD data were collected using a Bruker D8 system with GADDS software (area detector frame buffer software for data collection, display and data processing). Cu K alpha radiation was employed from a sealed tube source. The X-ray beam was conditioned via an incident beam mirror for removal of K beta radiation. A 300 micron pinhole snout was used for beam collimation. The detector used was a Hi-Star area detector. Sample to detector distance was 15 centimeters. Data reduction was performed using GADDS software. Phase identification was performed using Jade v 9.0 and the ICDD (International Centre for Diffraction Data) database. Analyses were performed either on powder scraped physically from the corroded surface of the component of interest, or in situ directly on the component. The approximate sampling area for each technique is a circle about 500 microns in diameter. Data are plotted as signal intensity versus two-theta (2θ) in degrees.

Analyses designated as obtained by XRD (i.e., not “micro”) were collected using a Siemens D500 q/q powder diffractometer. The data were collected over a scan range of 10-80° 2θ at a step size of 0.04° 2θ and a dwell times ranging from 1 to 20 sec. Monochromatic Cu Kα (0.15406nm) radiation was produced using a diffracted beam curved graphite monochromator. Fixed slits of 1.0, 1.0, 1.0, 0.15, and 0.15 degrees were used. The instrument power was 40kV and 30mA. Alignment and calibration were checked using a LaB₆ (NIST SRM660) external standard. XRD analysis was performed on corrosion product scraped physically from the component of interest. Data were plotted as signal intensity versus two-theta in degrees.
X-ray fluorescence (XRF) data were collected using a Bruker M4 micro-XRF system employing a Rh X-ray source, a polycapillary beam optic (resolution ~30 micron spot size), and a silicon-drift detector. Elemental mapping of specimens was performed by rastering the specimen over various X–Y coordinates while simultaneously collecting XRF spectra from each map coordinate. These spectra were employed to generate the false color images displaying the concentration of a given elemental species as a function of map location.

3. RESULTS AND DISCUSSION

This section includes detailed descriptions of several of the components and provides an overview of the questions that the accelerated corrosion studies were intended to answer and resulting observations applicable to each question. A summary of the electrical measurements performed during exposure to the corrosive atmosphere and observations consistent among the three assembled electrical test circuits are also presented. The results of the characterization work follows, organized by test circuit, then by component. Because a subset of wires and components was characterized, not every component will have observations and data associated with it.

3.1 System Measurements

3.1.1 Electrical Measurements

To determine if corrosion products on contacts could lead to resistive heating, measurement points were established to allow the voltage drop across the entire electrical circuit to be measured. Measurements were made several times a week and were taken while the circuit was activated under a 12 A load. Measurements were made with a Keithley 2000 multimeter. Dedicated test points were wired into the circuit (external to the exposure chamber) to allow measurement on a regular basis. Voltage drop data are shown in Figure 8 (hot leg) and Figure 9 (neutral leg). All three of the circuits exhibited essentially identical behavior. The hot leg of the circuit registered about 40 milli-volts and remained constant throughout the test. The neutral leg of the circuit registered about 25 milli-volts and remained constant. The difference is due likely to the smaller number of components with neutral wire connections (both switches and the standard circuit breaker used only hot wire connections). The small magnitude of the voltage drop suggests that resistive heating—due to the presence of corrosion products—should not be a concern.
Figure 8. Voltage measurements made along the length of the ungrounded conductor (hot leg) for each of the three circuits, which included the load center and all the electrical components.

Figure 9. Voltage measurements made along the length of the grounded conductor (neutral leg) for each of the three circuits. Circuits included the load center and all of the electrical components.
3.2 Observations Across All Circuits

Several materials characterization tests were performed on a variety of components to measure and observe the corrosion product across different metals and components. Following disassembly, all three circuits had the following similarities:

- Extensive black corrosion product on bare wires;
- Variability in extent of corrosion (thicknesses, colors, morphology) on contact screws;
- Lack of corrosion observed on galvanized metal plates;
- A variety of microgrowth morphologies;
- Apparent “layering” of growth;
- Variations in corrosion product thickness between the inner (less gas access) and outer (exposed) areas;
- Flaking, blistering, and spalling of corrosion product films. This type of behavior is often observed with thick corrosion product films on metal substrates.
- Bare metal at contact points (pinch-points or screw contacts, contact plates) on wires, screws, and contact plates;
- Evidence of oxidation on the outer surface of the sulfide films. This is consistent with long-time exposure in an oxidizing environment.
- Multiple phases of copper sulfide on various surfaces.

Several of these observations have precedent in the literature of corroded copper, even though much of the literature is for tests with shorter timeframes and less aggressive conditions than those described in this report. These references illustrate the complexity observed in the products. For example, a variety of microgrowth morphologies have been observed in atmospheric monitoring of corrosion and controlled laboratory testing. Similar “layering” of growth has been observed using SEM and other analysis techniques. The flaking, blistering, and spalling of corrosion product films observed here is often observed with thick corrosion product films on metal substrates. The presence of multiple phases of copper sulfide, and in some cases, the oxidized form antlerite, also has precedent. In many studies, the goal did not

include identifying a specific crystal phase or stoichiometry. Therefore, only a generic Cu$_2$S assignment is made in these published reports.$^{10}$

Some of the specific crystalline phases of copper sulfide observed here are different than the digenite phase observed in the harvested components reported earlier. This is not unexpected because the accelerated conditions used here were not meant to mimic exactly conditions in the field. The sulfidation corrosion of copper is a complex event in which electrochemical, chemical, thermodynamic, and diffusion processes are all occurring. Each process is dependent upon factors such as temperature, concentration of gases present, and composition of local material at the surface. For atmospheric corrosion, or corrosion in mixed flowing gas experiments such as those in which oxygen is present, both copper oxides and sulfides form. In the case of copper sulfides, a variety of crystalline phases (elemental arrangements), each with a different stoichiometry (ratio of copper to sulfur), is possible and also can be found in nature. Some natural minerals, their stoichiometries, and weight percent of copper, include chalcocite (Cu$_2$S – 79.85% Cu), djurleite (Cu$_3$S$_16$ – 79.34% Cu), digenite (Cu$_9$S$_5$ – 78.10% Cu), and covellite (CuS – 66.46% Cu). In nature, and in laboratory tests, these copper sulfides can be oxidized and/or hydrated to form additional species, such as antlerite (Cu$_3$(SO$_4$)(OH)$_4$) and posnajakite (Cu$_4$(SO$_4$)(OH)$_6$·H$_2$O). For these reasons, the specific phase detected is less important than where the corrosion is (or is not) occurring.

The following tables contain comparisons of corrosion product thicknesses, using measurements from SEM cross section images of bare copper wires and a brass contact pad (1-RS-HSP2). The Figure number in this table documents where some of the SEM images used for these measurements can be observed. These measurements are not meant as a statistical sampling, and in each case, the thickest area or interesting feature was selected for measurement. These thicknesses are greater than those observed in the harvested components reported previously, where the maximum thickness found in any of the homes analyzed was 18 microns.

Table 2. Table of Corrosion Product Thickness Observed in SEM Cross Sections of Bare Copper Wires.

<table>
<thead>
<tr>
<th>Part</th>
<th>Figure Number</th>
<th>Comments</th>
<th>Thickness (microns)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-ACB-NW-BUS-1</td>
<td>Figure 26</td>
<td>Stranded wire Large pit</td>
<td>26.7</td>
</tr>
<tr>
<td>1-ACB-NW-BUS-1</td>
<td>Not shown</td>
<td>Stranded wire Flaked off layer + tight layer</td>
<td>6.3</td>
</tr>
<tr>
<td>1-ACB-NW-BUS-1</td>
<td>Not shown</td>
<td>Stranded wire “divot” growth</td>
<td>6</td>
</tr>
<tr>
<td>1-ACB-NW-BUS-1</td>
<td>Not shown</td>
<td>Stranded wire Wide pit with upper layer that appears to have growth under it</td>
<td>21.7</td>
</tr>
<tr>
<td>1-ACB-NW-BUS-1</td>
<td>Figure 27</td>
<td>Stranded wire Wavy layer with wide pit</td>
<td>8.5</td>
</tr>
<tr>
<td>1-GR-NW-1</td>
<td>Figure 18</td>
<td>Outer layer with what appears to be growth underneath</td>
<td>11.4</td>
</tr>
<tr>
<td>2T-RS-HW1</td>
<td>Not shown</td>
<td>Blister growth</td>
<td>16.4</td>
</tr>
<tr>
<td>2T-RS-HW1</td>
<td>Not shown</td>
<td>Blister growth</td>
<td>25.7</td>
</tr>
</tbody>
</table>

Table 3. Table of Corrosion Product Thickness Observed in SEM Cross Sections of Brass Contact Pad HSP2.

<table>
<thead>
<tr>
<th>Part</th>
<th>Figure Number</th>
<th>Comments</th>
<th>Thickness (microns)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-RS-HSP2</td>
<td>Figure 12, Figure 13</td>
<td>Void below corrosion film. Measure depth from perpendicular</td>
<td>15.3</td>
</tr>
<tr>
<td>1-RS-HSP2</td>
<td>Not shown</td>
<td>At inside curvature of part</td>
<td>12</td>
</tr>
<tr>
<td>1-RS-HSP2</td>
<td>Not shown</td>
<td>Separation of outer layer, possible fill underneath</td>
<td>36.6</td>
</tr>
<tr>
<td>1-RS-HSP2</td>
<td>Not shown</td>
<td>Wide view with layers and pits</td>
<td>52</td>
</tr>
</tbody>
</table>

27
3.3 Results for Circuit No. 1

3.3.1 Summary of Work

Figure 10 shows the components for Circuit 1 prior to disassembly. Once they were removed from the box and examined, several of the parts were selected for various analyses. Table 4 presents the individual components and associated analyses for Circuit 1.

Figure 10. Photographs of the electrical components tested in Circuit 1. The view from the bottom (C) shows how the system was wired electrically.
### Table 4. Circuit No. 1 Components Submitted for Analysis

<table>
<thead>
<tr>
<th>Part</th>
<th>Sample ID</th>
<th>Optical</th>
<th>SEM</th>
<th>FIB</th>
<th>Metallography (SEM)</th>
<th>X-ray</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rocker Switch</td>
<td>1-RS-HW-1</td>
<td></td>
<td>✔</td>
<td></td>
<td></td>
<td>✔</td>
</tr>
<tr>
<td>Rocker Switch</td>
<td>1-RS-HSP-2</td>
<td>✔</td>
<td></td>
<td>✔</td>
<td></td>
<td>✔</td>
</tr>
<tr>
<td>Toggle Switch</td>
<td></td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
<td>✔</td>
</tr>
<tr>
<td>GFCI Receptacle</td>
<td>1-GR-HW-1</td>
<td>✔</td>
<td></td>
<td>✔</td>
<td></td>
<td>✔</td>
</tr>
<tr>
<td>GFCI Receptacle</td>
<td>1-GR-GW-1</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
<td>✔</td>
</tr>
<tr>
<td>Receptacle</td>
<td>1-R-HW1</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
<td>✔</td>
</tr>
<tr>
<td>AFCI Circuit Breaker</td>
<td>1-ACB-NW-BUS-1</td>
<td>✔</td>
<td></td>
<td>✔</td>
<td></td>
<td>✔</td>
</tr>
<tr>
<td>Grounded Plug</td>
<td>1-PLUG-1</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
<td>✔</td>
</tr>
<tr>
<td>Cu coupon</td>
<td>Cu-5</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
<td>✔</td>
</tr>
</tbody>
</table>

#### 3.3.2 Rocker Switch

Figure 11 shows macro images for the rocker switch. The image in (A) shows one set of electrical connections to the switch. The image in (B) is of the contact plate under one of the screws. The image in (C) shows a close-up of one of the connections, and (D) shows both sides of the hot wire that was connected to the switch. The black color of the metal parts is the result of the sulfidation process. The contact plate is tarnished completely. Importantly, a row of shiny contact points is visible where the wire contacted the plate. There is a corresponding set of similar features on the wire, where the wire was in intimate contact with the screw. Note that bare metal is visible, indicating that the interface was shielded from the active species in the gas. The screws exhibit less attack. The wire shows considerable corrosion, with “furry” growth observed next to the insulation. Additionally, there is a band of bare metal adjacent to the insulation that appears to have suffered less attack. That is likely the result of slight movement of the wire, exposing fresh surface that had been protected previously by the insulation. This does point out the fact that the insulation shields the underlying metal from exposure to the gas. However, it is also important to note that this accelerated test is designed for boldly exposed metals. It is not designed to accelerate transport of aggressive species through the insulation. Thus, it does not represent 40 years relative to sulfide diffusion and cannot be used to assess the ability of the insulation to protect the wire over a 40-year life.
Figure 11. Optical images from sample 1-RS (rocker switch). The contact plate is shown in B, and one of the hot wires is shown in D (both sides of the same wire are shown). Note the presence of extensive corrosion product on the copper hot wire and contact plate.

Figure 12 shows SEM images of a cross section of the hot screw plate from the rocker switch. Figure 13 and 14 present the elemental data obtained from EDS analysis of the cross section shown in Figure 12. In Figure 13A, a thin layer of corrosion product is visible on the surface of the contact plate. There is evidence of voids below the corrosion product layer, consistent with the sulfidation process. The elemental maps also show Cu and Zn (Figure 13C-D), elemental constituents of the substrate (brass). The sulfide layer is clearly visible on the surface. As seen in Figure 14, the corrosion product layer also contains chlorine and oxygen.
Figure 12. SEM cross section of the hot screw plate (HSP2) from the rocker switch in Circuit No. 1.

Figure 13. SEM image (top left) and elemental maps using EDS of HSP2: sulfur (green), Cu (red), and Zn (purple).
3.3.3 Toggle Switch

Figure 15 shows optical images of the toggle switch from Circuit 1. The metal contact surfaces exhibit sulfidation. Tarnishing of the wires, contact plates, and screws was observed.

(A) 1-TS-1  (B) 1-TS-HW1-1

3.3.4 GFCI Receptacle (push in)

Optical images of the GFCI receptacle from Circuit 1 are given in Figure 16 and Figure 17. Corrosion of the contact surfaces was observed. The screws presented in (B) do not exhibit significant attack. Their color indicates that they are not brass screws, so they are considerably more resistant to sulfidation. Interestingly, the push-in wire (D) exhibits the presence of an extensive layer of corrosion product. This indicates that the gas permeated the receptacle itself. This is supported by the images in Figure 17, which show the insides of the GFCI. Clearly, the receptacle case presents little or no impediment to the corrosion gas. It is interesting to note that many of the internal components were corroded during this test (Figure 17D). Note that the majority of the contact is tarnished, with the real contact area (the spot near the center) exhibiting considerably less corrosion. During the test, one of the load lamps arced, resulting in tripping of
the AFCI for that circuit. This demonstrated that the AFCI component continued to function. The GFCI receptacles were functioned at the conclusion of the test using the “test” button. All functioned properly.

(A) 1-GR-1

(B) 1-GR-2

(C) 1-GR-HS1-1

(D) 1-GR-NW-1

Figure 16. Photos of outside of GFCI receptacle from Circuit 1.
Figure 17. Photos of inside of GFCI receptacle from Circuit 1.
The neutral wire from the GFCI receptacle was cross sectioned for analysis. The SEM images are shown in Figure 18. A thin corrosion product layer can be seen covering the wire surface. A very conservative estimate of the decrease in cross section of the wire can be calculated based on the SEM images. As given in Table 2, the corrosion product thickness was 11 microns. If we assume that the thickness is uniform, and that the corrosion product thickness equals the loss of metal, the cross sectional area of the wire at the conclusion of the test is 98% of the original cross section. In reality, the area decrease is considerably less (the corrosion product thickness is less than the thickness of lost). Thus, attack of the wire did not result in a significant decrease in cross sectional area and does not affect the current carrying capabilities of the wire.

![SEM images of cross section of neutral wire 1-GR-NW1.](image)

### 3.3.5 Receptacle

Analysis of the receptacle from Circuit 1 included optical images and SEM analysis of the hot wire. Optical images are presented in Figure 19 and Figure 20. They show corrosion of the Cu base materials. Very little attack of the neutral screws or the ground screw was observed. The wires exhibited significant attack, but were essentially untouched in two locations: where the wire was in intimate contact with screws, and where the wire contacted the pressure plate.
Figure 19. Optical photographs of receptacle from Circuit 1. The image in (A) shows the side view of hot wires; (B) hot screw and contact plate; (C) side view of neutral wires; and (D) neutral screw and contact plate.

Figure 20. Optical photographs of hot wire removed from receptacle from Circuit 1.
SEM analyses of the hot wire are shown in Figure 21. The location of these analyses is the bend in the wire. Note that the corrosion is significantly more extensive on the outside of the bend. The elemental maps clearly show the difference in corrosion product thickness, clearly a result of shielding the inner portion of the bend from the corrosive gasses.

Figure 21. SEM image (A) of hot wire 1-R-HW1; (B) SEM image of screw contact surface with zoomed area (top right); (C) EDS elemental map showing copper (red) and sulfur (green) distribution; and (D) SEM image ~300 µm from contact (all with 100 µm measurement bars).
Figure 22 presents SEM images from various locations on the hot wire. Several corrosion product morphologies can be seen. In addition, there is evidence of spalling of the film. Once the film ruptures, it reforms below the previous layer.

Figure 22. SEM images showing the different morphologies of corrosion products on the hot wire from the receptacle in Circuit 1.
In addition to differences in film morphology across the surface, there is also evidence of differences in film composition. An example is shown in Figure 23. Clearly, certain features are oxygen rich, while others are sulfur-rich. The oxygen-rich features are on the outer surface of the film, and are likely due to oxidation of the surface of the copper sulfide.

Figure 23. SEM images and elemental maps for two areas on wire 1-R-HW-1, showing copper (red), oxygen (blue), carbon (purple), and sulfur (green) distributions.
Figure 24 shows optical images of the AFCI from Circuit 1. The output from the standard circuit breaker provides the input to the AFCI. In addition, the neutral feed is connected to the neutral input, and the neutral sense wire is connected to the load center bus. Sulfidation of the contact surfaces can be seen. Both the neutral and hot wires exhibit corrosion, as well as the stranded neutral sense wire (where it is connected to the load center bus–lower left image).

Figure 24. Optical images of the load center and AFCI for Circuit 1.
Figure 25 presents SEM images of the cross section of the neutral wire. A multilayered structure is observed. There is little or no adhesion between some of the layers, suggesting that spalling and/or delamination of the individual layers occurred during the corrosion process.

Figure 25. SEM image of the cross section of the neutral wire from the arc-fault circuit breaker from Circuit 1.
Elemental maps for the neutral wire are shown in Figure 26 and Figure 27. A continuous sulfide film can be seen, with subsurface attack of the base metal which is typical of copper sulfidation reactions. In the void formed in the base metal, oxidation has occurred, filling the cavity with copper oxide and/or copper sulfate.

Figure 26. EDS map of wire cross section showing oxygen in “pocket. The elemental maps for Cu (red), S (green), and oxygen (blue) are shown.
Figure 27. Elemental maps taken on cross section of the neutral wire from the arc-fault circuit breaker from Circuit 1. The image at the top left is an SEM image of the area being analyzed. The elemental maps for Cu (red), S (green), and oxygen (blue) are shown.
Figure 28 shows elemental maps taken on the stranded wire from the AFCI that was attached to the load center bus. All of the strands of wire are covered with a sulfide layer.

Figure 28. Elemental maps taken on cross section of the neutral wire from the arc-fault circuit breaker from Circuit 1. The image on the left is an SEM image of the area being analyzed. The elemental maps for Cu (red), S (green), and oxygen (blue) are shown.
3.3.7 Circuit Breaker

Figure 29 shows optical images of the contact surfaces of the standard circuit breaker from Circuit 1. Corrosion of the upper contact is visible. The lower contact appears to be plated, or constructed of a different material. It did not exhibit extensive corrosion.

(A) 1-cb-contact1-1  (B) 1-cb-contact1-2

Figure 29. Optical images of the contacts in the standard circuit breaker in Circuit 1.
3.3.8 Grounded Plug

Figure 30 shows images of the plug used to connect the standard receptacle to the GFCI receptacle in Circuit 1. Note that the contact surfaces exhibit corrosion, even when inserted into the socket. Also, note that there are areas of contact that prevented or reduced the attack of the base metal.

(A) 1-gr-r-plug-l

(B) 1-gr-r-plug-gs-1

(C) 1-gr-r-plug-hs-1

(D) 1-gr-r-plug-ns-1

Figure 30. Optical images of the plug used to connect the standard receptacle to the GFCI receptacle in Circuit 1. The prongs and screws are blackened relative to those on a new, as-received, unexposed plug.

Corrosion product was scraped from the “hot” prong of the plug and analyzed by XRD, and an in situ spot was analyzed by µXRD. These tests detected that the base metal is a copper-zinc alloy and revealed the presence of a covellite (CuS) copper sulfide phase. Additional phases detected only by XRD include: copper sulfate hydroxide hydrate and copper zinc sulfate hydroxide hydrate (Ktenasite), whereas, only cuprite Cu2O was detected by µXRD. Lack of full cross-validation of XRD and µXRD is expected due to the differences in counting statistics and sampling area. The µXRD spectrum is shown in Figure 31 and illustrates the peaks attributable to the covellite, cuprite, and copper-zinc alloy. The figure is also labeled with the approximate locations where a digenite copper sulfide phase would be detected.
A copper-zinc alloy was detected for the ground prong of the plug by XRF analysis, and the elemental map, Figure 32, shows that the sulfur coverage is uniform within the tested length (approximately 2.5 mm) near the prong end.
Figure 32. Photo of one side of ground prong and XRF element maps of ground prong of grounded plug. XRF images taken “end-on” of full prong.
3.3.9 Copper Exposure Coupon

XRD of scraped powder and in situ µXRD both detected the copper sulfide phase known as “djurleite” (Cu$_{31}$S$_{16}$) as the predominant corrosion product observed on copper witness coupon No. 5. Data for the µXRD, along with the image indicating the location of the analysis, are shown in Figure 33. The image shows the copper witness coupon (which has a small hole for mounting, which was not used in these tests) on top of the metal instrument sample holder. The grey to bluish-black appearance of the sample is consistent with the appearance of djurleite. Both XRD and µXRD techniques verify also that the witness coupon is pure copper.

XRF analysis and mapping confirms sulfur signal in specimen and indicates uniform spatial distribution of sulfur (data not shown).

Figure 33. Video image (inset) showing location of µXRD analysis and µXRD data for witness coupon No. 5, showing labeled copper and djurleite peaks (green).
### 3.4 Results for Circuit No. 2

Table 5. Circuit No. 2 Components Submitted for Analysis.

<table>
<thead>
<tr>
<th>Part</th>
<th>Sample ID</th>
<th>Optical</th>
<th>SEM</th>
<th>FIB</th>
<th>Metallography (SEM)</th>
<th>X-RAY</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rocker Switch</td>
<td>2-RS-hsp-2</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rocker Switch</td>
<td>2-RS-HW-1</td>
<td>✓</td>
<td></td>
<td>✓</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rocker Switch</td>
<td>2-RS-PIHW-1</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(push-in hot wire)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GFCI Receptacle</td>
<td>2-R-GR-GW-1</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
<td>✓</td>
</tr>
</tbody>
</table>

#### 3.4.1 Summary of Work

Figure 34 shows optical images of the load center and the back side of the components for Circuit 2. Parts from the rocker switch and the GFCI receptacle were selected for cross section SEM and X-Ray analysis (see Table 5).

(A) CKT-2_6929

(B) CKT-2_6928

Figure 34. Photographs of the electrical components tested in Circuit 2. The view from the bottom (left image) shows how the system was wired electrically. The image on the right shows the load center.
3.4.2 Rocker Switch

Optical images of the hot wires from the rocker switch in Circuit 2 are shown in Figure 35. The wire on the left was connected using a screw contact, while the wire on the right is from the push-in contact. Note that both exhibit corrosion across the majority of the surface, with contact areas where the surface was shielded, showing little or no attack.

Figure 35. Optical images of wires from the rocker switch in Circuit 2. Both the standard wire connection and the push-in connection are shown.

Figure 36 shows SEM images of the hot wire (screw connected) from the rocker switch. As with other components, there is evidence of film growth and rupture. It is important to note that the thickness of the corrosion product film is very small relative to the diameter of the wire. In other words, copper sulfidation under these conditions is a surface phenomenon and does not result in significant reduction in the cross sectional area of the conductor. As such, the corrosion process will have no effect on the current carrying capabilities of the wire.

Figure 36. SEM images of the cross section of the rocker switch hot wire from Circuit 2. The image on the left shows a low magnification image of the wire, with a high magnification image showing details of the corrosion product layer presented in the figure on the right.
Figure 37 shows the elemental maps of the wire shown in Figure 36. The corrosion product consists of a layer of copper sulfide, with a thinner layer of either copper oxide or possibly copper sulfate.

Figure 37. Elemental maps from the rocker switch hot wire (cross sections) in Circuit 2. The upper left image is an SEM image identifying the area being analyzed. Three maps are shown for Cu (red), S (green), and oxygen (blue).
XRD analysis of scraped powder from this contact plate detected the same phases that were detected in the powder scraped from the Circuit 1 plug: covellite, copper sulfate hydroxide hydrate, Cu-Zn sulfate hydroxide hydrate (Ktenasite), and a copper-zinc alloy (as the Cu$_{0.67}$ Zn$_{0.33}$). µXRD analysis was also consistent with that of the Circuit 1 plug, detecting cuprite, covellite, and copper-zinc alloy. Additionally a chalcocyanite phase (CuSO$_4$) was detected in situ on this part. The location of sampling and µXRD data are shown in Figure 38. The large metal piece behind the sulfidated (black) part is the instrument sample holder.

Figure 38. Video image (inset) and µ-XRD data for 2-RS-HSP-2 (contact plate), showing cuprite (orange), covellite (green), and copper/zinc (purple), and chalcocyanite (Cu+2SO$_4$-light blue) peaks observed.
3.4.3 GFCI Receptacle

XRD analysis of black powder scraped from the wire surface (top, Figure 39) and μXRD *in situ* of the ground wire (bottom, Figure 39) of the GFCI component of Circuit 2 detected covellite (CuS) and the antlerite copper sulfate hydroxide phase (Cu₃(SO₄)(OH)₄). In addition, the μXRD data shows copper oxide (cuprite) and copper metal.

Figure 39. XRD data (top) for powder scraped from wire 2-R-GR-GW showing covellite (green) and antlerite (blue), and μ-XRD data (bottom) showing antlerite (orange), cuprite (light blue), covellite (dark blue), and copper (green) peaks.
3.5 Results for Circuit No. 3

Table 6 indicates which components from Circuit 3 were selected for analyses. In this case, both optical and SEM examinations were conducted for several of the parts.

Table 6. Circuit No. 3 Components Submitted for Analysis

<table>
<thead>
<tr>
<th>Part</th>
<th>Sample ID</th>
<th>Optical</th>
<th>SEM</th>
<th>FIB</th>
<th>Metallography (SEM)</th>
<th>X-Ray</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rocker Switch</td>
<td>3-RS-PIHW-1</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rocker Switch</td>
<td>3-RS-HW-1</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Toggle Switch</td>
<td>3-TS-HW-1</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Look under insulation as well.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GFCI Receptacle</td>
<td>3-GR</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Receptacle</td>
<td>3-R-PIHW-1</td>
<td>✓</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Receptacle</td>
<td>3-R-PINW-1</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AFCI Circuit Breaker</td>
<td>3-ACB-1</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Circuit Breaker</td>
<td>3-CB-1</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Twist on connector</td>
<td>3-WN-1</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
3.5.1 Rocker Switch

Figure 40 shows optical images for the rocker switch from Circuit 3. The case was removed to reveal the interior components. As with other components, it appears that the case provides little or no protection against the corrosive gasses. Both interior and exterior components exhibit corrosion. The bottom two images show both sides of the hot wire. In this case, the wire was connected to the push-in contact. Corrosion of the wire surface was less nonuniform, but it essentially covered the entire surface other than the contact point.

(A) 3-rs-insides-3  
(B) 3-rs-insides-2  
(C) 3-rs-pihw-1  
(D) 3-rs-pihw-2

Figure 40. Optical images of the rocker switch and hot wires from Circuit 3.

Figure 41 shows SEM images of the hot wire from the rocker switch. The figures on the left are secondary electron images that show topography. In this case, the insulation was removed next to the corroded area. The images on the right are back-scattered electron images. In these images, brightness is a function of the atomic number of the element on the surface. The bright area in the top of the image represents bare copper exposed by the removal of the insulation; the lower portion of the picture is darker and represents the sulfide layer (i.e., copper has a higher/larger atomic number than sulfur). These images show that sulfidation of the copper stops where the insulation protects the underlying copper. Figure 42 shows the elemental maps associated with this location. Clearly, the insulation protected the underlying copper from sulfidation because the top of the elemental maps showing the region of the wire that was covered with insulation shows no sulfur.
Figure 41. SEM images of the hot wire from the rocker switch in Circuit 3. The figures on the left are secondary electron images, which show topography. The images on the right are back-scatter images. In these images, brightness is a function of the atomic number, with higher brightness indicating elements of higher atomic number.
3.5.2 **Toggle Switch**

Optical images of the toggle switch from Circuit 3 are shown in Figure 43. As with previous components, sulfidation was universal, except where the surface was shielded (contact points).

Figure 44 shows SEM images of the hot wire from this component. The top two images are back-scattered electron images and show the presence of bare Cu at a contact point (light region). The corrosion product film structure/morphology is similar to that found on other components, exhibiting layering, buckling, and spalling.

Figure 45 shows EDS elemental maps of two areas of the hot wire from the toggle switch. Copper is shown in red, and sulfur is shown in green. These images are consistent with the SEM images and show the presence of bare Cu at the contact points.
Figure 43. Optical images of the toggle switch in Circuit 3. The figure on the left shows the inside of the switch. Note the extensive corrosion product on some of the components. The image on the right is of the hot wire connected to the toggle switch (screw side).

Figure 44. SEM images of the hot wire from the toggle switch in Circuit 3. Note the spalling of the corrosion product film. The contact areas can be seen in the upper two images (bright areas) where the contact shielded the surface from the corrosive gasses.
Figure 45. SEM images and EDS elemental maps of the hot wire from the toggle switch in Circuit 3. Sulfur is shown in green, and copper is shown in red.

The following figures show different SEM magnification views of a FIB cut into areas of the hot wire from the toggle switch, near where the toggle switch screw was in contact with the wire. The FIB technique uses a thin layer of deposited platinum to keep surface features intact so there is a thin layer that appears white in the images—this is the platinum overcoat. Using a FIB cut is
potentially less damaging to the surface layers; however, it is time consuming; therefore, only limited FIB analyses were performed. Figure 46 shows a low magnification view with circles indicating locations of two FIB cuts. Since this is a very low magnification, the instrument aperture is observed as the wider circle. The bright area in the upper right, is the area where the screw made contact with the wire and appears to be free of corrosion product.

Figure 46. Low magnification FIB image of hot wire from toggle switch (3TS-HW-1), showing location of FIB cuts (red designated as “Section 1” and blue designated as “Section 2”) into corrosion product film.

Figure 47 shows SEM images of the FIB cut designated as “Section 1,” indicated by the red circle in the previous figure, at increasing magnifications (350 to 12,000X magnification). This FIB cut is approximately 50 microns from the nearest portion of the wire contacted by the screw. The middle top image illustrates wavy features of the corrosion product observed in SEM cross sections of other samples. The right top and lower left images illustrate an abrupt edge of the wavy growth that appears to have smaller features, suggesting flaking and regrowth of the corrosion product. The middle lower and right lower images show closer views of the corrosion layer and the varying gap between the wire and corrosion product. This is consistent with crosssection observations of other wires.
Figure 47. FIB cuts into 3-TS-HW-1 “Section 1” (see Figure 46) with scale bars at 200 (350X), 50, 20, 5, and 5 (12,000X) microns, respectively.

Figure 48 shows three images at the same magnification (20,000X) of the edge of the FIB cut. The copper wire is at the bottom, the varying contrast areas are consistent with solid copper. Some pitted areas are observed along with roughened product growth underneath the upper layer of corrosion. The bright white top layer is the platinum layer deposited on the sample to preserve features.

Figure 48. FIB cuts into 3-TS-HW-1 “Section 1” (see Figure 46) with scale bars at 4 microns each.

Interesting growth morphologies were seen in all samples, which included needles and other features. One example, with needle and spiral-like growths, observed near the FIB cut is shown
in Figure 49 at various magnifications. On the left is a wider view of “Section 2,” indicated with the blue circle. The orange box indicates the approximate area of the middle image, while the orange box in the middle image indicates the approximate area of the image on the right.

A closer look at the cross section of corrosion growth revealed by the FIB cut designated as “Section 2” is shown in Figure 50. In the top row, a bare copper area is observed to the upper right of the images, the area underneath the screw. The lower row shows higher magnification views of the change in the corrosion growth thickness from left to right.
Figure 51 contains a series of SEM images across the FIB cut at 20,000X magnification. Layering and pitting is observed; moving from left to right and closer to the contact point of the screw, the overall corrosion thickness transitions from approximately nine microns to significantly less than one.

![Figure 51. FIB cuts into 3-TS-HW-1 “Section 2” (see Figure 46) with scale bars at 4 microns each (20,000X).](image)

### 3.5.3 GFCI Receptacle

Figure 52 shows optical images of the GFCI receptacle from Circuit 3. The contact screws showed evidence of minor corrosion on the surface. The contact plates below the screws were corroded extensively. As seen in the figure, they were black at the conclusion of the test.

![Figure 52. Optical images of the GFCI receptacle. Note the darkening of the normally shiny contact pads and slight corrosion (discoloration) of the screws.](image)
3.5.4 Receptacle

Figure 53 shows optical images of the hot wires (push-in contacts) from the standard receptacle from Circuit 3. These results are consistent with previous samples, showing corrosion product across the surface except where the contact was made.

The SEM images shown in Figure 54 show similar features to other samples: multiple layers of the film, spalling, voiding of the metal, and bare metal at the contact point.

(A) 3-R-PIHW-2  (B) 3-R-PINW-1

Figure 53. Optical images of wires from the standard receptacle from Circuit 3. Both the (A) hot and (B) neutral wire are shown. Both are “push-in” type contacts.
Figure 54. SEM images of the surface of the neutral wire from the receptacle in Circuit 3. The connection was a push-in connection. (B) shows the contact point, which is brighter due to the presence of Cu without S. The end of the wire is shown in the lower images at two magnifications. Note the spalling of the corrosion product film. It is believed that the spalling occurred during the test, but is it possible that it occurred after exposure and prior to analysis.
SEM images of two FIB cuts (designated as “Section 3” and “Section 4”) on the neutral wire from the push-in connector of the receptacle are shown below at magnifications of 128x and 127x. The aperture of the instrument is observed as the outer view circle due to wide view and low magnification. “Section 3” (left) was cut at the edge of the corrosion product film. Section 4 (right) was cut into an area surrounded by “wavy” corrosion growth.

Figure 55. FIB cuts into corrosion product on surface of neutral wire 3-R-PINW-1 with “Section 3” (left) and “Section 4” (right) with scale bar of 500 micron.

Higher magnification images of “Section 3”, ranging from 350X to 12,000X are presented next (Figure 56) and display varying thickness as well as voiding of the copper below the corrosion product film. The fracture line down the center of the images in the top row shows an area of corrosion product that flaked off.
Figure 56. FIB cuts into neutral wire 3-R-PINW-1 “Section 3” (see Figure 55) with scale bars of 200, 50, 20, 10, 5, and 5 microns.

Higher magnification images of the pitting observed in “Section 3” are below (Figure 57); with the thinnest corrosion growth, the platinum FIB coating, and pure copper grains prevalent in the image on the right.

Figure 57. FIB cuts into neutral wire 3-R-PINW-1 “Section 3” (see Figure 55) with scale bars of 4 micron each.

Higher magnification views of “Section 4” are shown in Figure 58 and illustrate an additional morphology seen in some samples—that of a “bubbling” or blistering of the corrosion product layer. The magnification ranges from 350X to 25,000X. In the lower right image, three different kinds of growth layer are observed—a layer with few voids, a middle layer with multiple voids,
and a third solid layer with what appears to be striated growth. The top-most white layer in the image is the platinum FIB overcoat.

Figure 58. FIB cuts into neutral wire 3-R-PINW-1 “Section 4” (see Figure 55) with scale bars of 200, 50, 20, 10, 5, and 3 microns, respectively.
Figure 59 shows SEM images of the push-in hot wire from the receptacle in Circuit 3. The image on the left shows the contact point. It is clear that the intimate contact between the wire and the receptacle contact prevented sulfidation from occurring in the contact area. The image on the right shows the wire at the interface with the insulation. In this case, the insulation was removed to reveal the wire surface directly beneath the insulation. The fact that bare copper is visible in the region shielded by the insulation suggests that the insulation prevents the pollutant gas from contacting the copper wire.

(A) 3-R-PIHW-1_08

(B) 3-R-PIHW-1_21

Figure 59. SEM images of the hot wire from the receptacle in Circuit 3. The connection was a push-in connection. The contact point is shown in the image on the left. The image on the right shows a region of a hot wire that was covered by insulation on the bottom.
Figure 60 shows the EDS elemental maps from the wire in Figure 59. Copper corrosion products (primarily sulfide with some oxide and/or sulfate) are present on the wire surface, except where the contact was made. In that location, bare copper is visible.

Figure 60. EDS elemental maps of the wire shown in Figure 59. Copper (red), sulfur (green), and oxygen (blue) are shown.
3.5.5 Twist-on Connectors

Figure 61 shows optical images of twist-on connectors and the associated wires used to connect component assemblies inside the chamber. Corrosion of both the wires and the twist-on connectors was observed. The extent of corrosion decreased with depth into the twist-on connector. A thick corrosion product layer was found next to the insulation. The ends of the wires, where protected by the twist-on connector, exhibit bare copper.

(A) 2-output-WN1-2  
(B) 2-output-WN1-1

(C) 2-output-WN2-2  
(D) 2-output-WN2-1

Figure 61. Optical images of twist-on connectors with hot and neutral wires used on the output section of Circuit 2.
3.6 Questions to Answer

Based upon observations and measurements of corrosion products in our previous analyses of harvested components, the accelerated testing and analyses were intended to investigate a variety of questions.

1. A specific phase (composition) of copper sulfide product known as digenite \( \text{Cu}_9\text{S}_5 \) was observed in harvested electrical components—Is copper sulfide observed in accelerated aging tests?
   a. Several copper sulfide phases were observed, including covellite \( \text{CuS} \) and djurleite \( \text{Cu}_{31}\text{S}_{16} \). A sulfated hydrated phase known as antlerite \( [\text{Cu}_3(\text{SO}_4)(\text{OH})_4] \) was also observed. In the Battelle MFG test, the corrosion products are identified simply as copper sulfide. No distinction is made between the various forms (i.e., covelite, djurleite, diginite).

2. In a sampling of harvested components previously analyzed, corrosion products were observed in a variety of thicknesses up to 20 microns. In these tests, chosen to simulate aging for 40 years—what range of thicknesses is observed? Note: For ground wires from components harvested from homes (data below)—values normally were less than 10 microns, and typically, were less than 3 microns.

   | 01R–1-7 | 14R–2.5 |
   | 33R–2.4 | 48R–2-6 |
   | 50R–8   | 52R–3-18 |

In the accelerated aging exposure paradigm, on bare exposed Cu wires, corrosion product thickness was observed from approximately 6 microns up to 27 microns (see
Table 2). A corrosion product thickness of up to 52 microns was observed on a brass contact plate (see Table 3). Additionally, the corrosion product layer was more consistent (less range in thickness from location to location) and generally thicker on parts from the accelerated test than on samples collected from homes.

- For areas inside components, such as wires or contacts within circuit breakers, receptacles, and GFCIs, cross sections were not measured. The following observations were made optically and by SEM:
  - Plug-in type wires have a range of corrosion product thickness. In contact areas (e.g., screw to wire contact) metallic copper is visible, indicating that the contact surfaces were shielded from the corrosive gasses.
  - Wrap-on wires have bare copper area where contact was made.
  - Inside twist-on wire connectors exhibit various thicknesses of corrosion product. One connector had a completely bare area (teeth and grooves), whereas another only has teeth (where contact was made).
  - Corrosion product layer thickness on metal surfaces inside circuit breakers/receptacles/GFCIs has not been measured.
3. Is there corrosion product under wire insulation?
   a. No. Visual and SEM analyses of several samples of Cu wire beneath the insulation showed no tarnishing or evidence of corrosion. However, this test does not accelerate H$_2$S diffusion through the insulation, so it is not appropriate for analysis of this failure mechanism. A better understanding of H$_2$S diffusion through PVC insulation would be required to address this issue. The fact still remains that the insulation protected the underlying wire from corrosion.

4. In the previous study of harvested electrical components (a limited sampling of six homes), corrosion product was NOT observed between wires and other fixed contact surfaces. Under the accelerated conditions of this work, is corrosion product observed underneath or between wire and contact surface?
   a. Based on visual and SEM inspection, corrosion product did not grow between surfaces in contact.

5. Different levels of corrosion were observed previously in harvested components where contact is made/broken (such as switches and GFCIs). Does corrosion grow between or on contact surfaces in switches and GFCIs?
   a. Corrosion product was observed on these contact surfaces. Components with moveable parts were not exercised during the accelerated aging tests. In general, these parts were tested in the closed condition. The switches and circuit breakers were all closed so that a continuous circuit was established. The sulfide film forms wherever bare metal is exposed to the environment.

6. What are the physical/electrical properties of the corrosion products?
   a. The copper sulfide that forms on the metal surface starts out as a thin, coherent film. It is strongly attached to the metal. It is not easily wiped off. It is, however, brittle, and is removed easily through mechanical abrasion (such as through inserting/removing a plug). In that case, only the material directly in contact with the sliding member is removed. The remainder of the corrosion film remains attached. As the film thickens, stresses can build up in the film, resulting in buckling and/or rupturing of the film. This process exposes fresh surface below the film, and the corrosion process continues.
   b. The electrical conductivity, as reported in the literature, ranges from 0.07 ohm$^{-1}$cm$^{-1}$ to 2700 ohm$^{-1}$cm$^{-1}$ depending on composition of the sulfide (Cu$_2$S to Cu$_{1.8}$S).$^{11}$

7. In the analysis of harvested electrical components, a cauliflower-like growth morphology was observed with some fracturing of the corrosion product layer and apparent “layering” similar to growth rings of a tree. Under the conditions of this test, is fracturing observed that could alter the access of corrosion gas to the underlying copper?

---

a. Yes, as the film thickens, rupture of the film (often identified as flaking, spalling, fracturing) was observed. This process certainly provides greater access of the pollutants to the metal surface. Despite constant ambient temperature conditions, “layering” was also observed.

8. In harvested electrical components, the nature of the corrosion product varied widely on wires, screws, and contact plates. We used a simple rating system (1 to 5, with 1 representing little or no corrosion, and 5 representing the most severe corrosion) to categorize the observed corrosion products in an effort to correlate measurements with harvested components (see Table 7). Using the same rating system, what range of corrosion is observed under accelerated conditions?

a. Exposed copper wire and brass contact plates were all category 5, with the visual extent (thickness) exceeding any observed in the harvested electrical components. Exposed screws were observed across the whole range and could be categorized between 1 and 5. Across the board, the extent of corrosion was much more severe in this test than in the harvested components.

9. While our lab did not perform electrical measurements on the harvested electrical components investigated in our earlier study, an important question in these current tests is: does corrosion cause any change in the resistance of the electrical circuit that could cause tripping or heating of the circuit?

a. Circuit resistance values did not show any detectable changes during measurements, which were made several times a week during the 8-week exposure. There were several trip events, but they were all the result of arcing at the lamp contacts, which were all exterior to the test chamber, and were not part of the hardware being tested.

Table 7. Table of Corrosion Ratings for Electrical Components from Homes.

<table>
<thead>
<tr>
<th>Category</th>
<th>Number of components (% of total)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0 (0%)</td>
</tr>
<tr>
<td>2</td>
<td>14 (16.1%)</td>
</tr>
<tr>
<td>3</td>
<td>25 (28.7%)</td>
</tr>
<tr>
<td>4</td>
<td>35 (40.2%)</td>
</tr>
<tr>
<td>5</td>
<td>9 (10.3%)</td>
</tr>
<tr>
<td>unknown</td>
<td>4 (4.6%)</td>
</tr>
</tbody>
</table>

Mean 3.4
Std. Dev. 0.9

NOTE: A total of 87 individual electrical components from six different homes are included. The components listed as unknown were circuit breakers for which the extent of corrosion could not be determined.
4. CONCLUSIONS

In a previous study, we characterized corrosion products on wires and electrical components harvested from homes by CPSC staff. One result of that study was the conclusion that some components were exposed to environmental conditions that could be classified (in the field of corrosion) as a Battelle Class III or IV. This study, a follow-on to the domestic analysis, was designed to simulate a long duration installation in a household and observe the effects with respect to failure modes, corrosion generated, and any other issues that might affect electrical performance.

A variety of commercial, off-the-shelf electrical components (switches, outlets, AFCIs, GFCIs) were enclosed in a chamber for eight weeks under Class IV Battelle environmental conditions and thus, simulated corrosion aging of ~40 years. This type of testing is commonly referred to as “accelerated aging.” Because actual field conditions have not been measured, this is an estimate meant to exceed conditions that might occur over that timeframe in an actual house.

Copper sulfide corrosion product characterized in this work exhibited:
- growth on copper-containing components;
- growth inside non-hermetically-sealed components (receptacles, breakers, twist-on connectors);
- reduced growth in areas of reduced availability to gas diffusion;
- many crystallographic phases;
- a wide variety of morphologies;
- a range of oxygen content within portions of the product; and
- physical features due to flaking, spalling.

Exposed areas of contact screws also exhibited corrosion; however, the copper content or exact composition of corrosion was not measured. The range of corrosion observed, as was observed in the study of harvested components, could be categorized from one to five (representing little or no corrosion to representing the most severe corrosion).

The insulation appeared to be very effective in protecting the underlying copper wire. In several instances, the insulation was removed, and no evidence of wire corrosion was seen. These observations cannot be extrapolated to a 40-year lifetime, as the accelerated test would not be expected to accelerate diffusion of sulfur-containing gasses through the PVC insulation. However, the fact remains that the insulated wire, when exposed to a very aggressive environment, did not suffer corrosion beneath the wire insulation.

With respect to electrical characteristics and possible hazardous conditions, our testing and materials characterization measurements revealed:
- no growth between surfaces that retained physical contact during testing;
- no manifestations of fire or overheating;
- no significant detectable changes in circuit resistance (measured via voltage drop); and
- no significant degradation of wire cross sectional area.
5. **IMPLICATIONS**

The 2009 study was able to identify the nature and extent of the corrosion product that had occurred during the exposure to problem drywall in the six homes, but not how much corrosion could occur for the typical lifetime of household electrical components, and what effects this might have on the electrical function and safety of powered circuits.

At the conclusion of this test, copper-containing elements (wires or contact plates) of the electrical components exhibited more extensive (with respect to coverage and thickness) corrosion than any of the components harvested from homes. The corrosion product thickness is small, however, relative to the thickness of any component element. Because the components from this test exhibited no significant loss in cross section, the current carrying capability of the conductors is not expected to be diminished. All of the exposed components functioned throughout the test, with no indication of performance degradation, even though significant corrosion was observed on the copper-based metals. Additionally, no voltage drop (across the entire circuit) was seen, indicating that the presence of the corrosion product film had not resulted in an increase in resistance. There was no intentional activation of the GFCI or AFCI assemblies during the course of the tests. One of the AFCI circuit breakers tripped several times during the test, due to arcing at one of the lamp contacts (external to the exposure chamber), indicating that the functional capabilities had not been compromised. All of these results suggest that the electrical performance and safety of static components or AFCI and GFCI assemblies have not been compromised by the corrosion process.

Electrical components, such as rocker switches and contact plates in receptacles that would normally receive both intermittent exposure and intermittent mechanical function in a household installation, were not exercised during the accelerated corrosion test. It is possible that moving metal surfaces relative to one another would remove fragile corrosion layers, reexposing bare metal conductor to the corrosive agent, leading to a higher rate of corrosion and loss of more of the underlying copper as it diffuses to corrosion product layer. Conversely intermittent use and incomplete removal of corrosion could potentially cause a higher resistivity, thus, affecting performance. Further testing that includes functioning of switches and inserting and removing plugs from receptacles is recommended to determine the effect on performance by these specific contact processes.

While metal-to-metal contact did not appear to be degraded by the corrosion, exposed metallic parts were corroded, and this likely would render them marginally fit for service (such as in relation to a remediation, they should be replaced).
6. DISTRIBUTION

U.S. Consumer Product Safety Commission
4330 East West Highway
Bethesda, MD 20814
Attn: Andrew Trotta
Mark Gill

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