Detection of Corrosion in Thick Film Resistors by X-Ray Imaging

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ABSTRACT In this article, a non-destructive method is presented, using 2D X-ray imaging, to investigate corrosion defects in thick film resistors stressed by two different corrosion experiments, a single gas experiment and a flowers of sulphur experiment. In total, 370 devices under test (DUTs) were investigated, using the 2D X-ray imaging technique, of which 10 were imaged in a sequence of 2D X-ray images to evaluate the corrosion propagation. The observed underlying corrosion product was verified by using focused ion beam analysis combined with an energy dispersive X-ray spectroscopy. The presented 2D X-ray imaging technique enables a fast and non-destructive method for accurately identifying corrosion and corrosion progression in thick film resistors. The presented imaging method was found to be particularly suitable for supplier benchmarking purposes, as well as for the failure analysis of field returns as it allows for a proper corrosion evaluation of surface mount thick film resistors, while keeping the analysed assembly intact.

INDEX TERMS Corrosion, resistors, failure analysis, scanning electron microscopy, silver, sulphur, X-rays.

I. INTRODUCTION

Frequency converters are used in vastly different fields of industries and applications, such as solar, wind, pulp and paper production, rubber vulcanization and mining. In solar applications, one of the biggest stressors for a frequency converter is temperature cycling, which has a big impact on the ageing phenomena of the insulated gate bipolar transistor (IGBT). In wind applications, humidity plays a significant role on the drive’s reliability. Other harsh environments, such that contains corrosive pollutants, can be found in pulp and paper production, as well as in rubber vulcanization and mining applications. In these industries, the amount of atmospheric sulphur in the environment has a big impact on the lifetime of the frequency converter. Critical components, such as power modules [1], electrical contacts and thick film resistors are affected by corrosion due to sulphur in the environment. In these electrical components, corrosion phenomena, such as electrochemical migration [1]–[3] and open connection [4], [5], have been reported.

Surface mount thick film resistors are used in almost all industrial electronic devices. Although the resistor is one of the simplest and most inexpensive amongst all of the other components, it may be responsible for the complete functional breakdown of a device operating in a harsh environment. In these environments, corrosion failures such as resistive shorts, open connections or changes in resistance indicating intermittent failure are typical. Usually, in sulphuric atmospheres, the weakest spot in a thick film resistor is the silver (Ag) termination. It is well known that silver can rapidly corrode in harsh atmospheres, particularly in those containing sulphides [6]–[8]. Silver reacts with sulphur (S), forming silver sulphide (Ag2S). Silver sulphide is known to be non-conductive and leads to changes in resistance or, eventually, an open circuit [4]. This silver termination is the most underlying layer in the resistor structure. It is protected by two different metallic terminations. The first one is a protective barrier, typically made of nickel (Ni), and the second one is an outer solder termination of tin (Sn). Additionally, a thin conformal organic epoxy layer is applied to the resistor to limit the gas permeation of water vapour and pollutant gases to reach the silver termination. A typical cross-section
image of a thick film resistor is shown in Fig. 1a) with the corresponding areas of interest visualized in Fig. 1b).

This study demonstrates an effective non-destructive X-ray imaging method for analysing corrosion in thick film resistors. Multiple studies [16]–[20] have shown successful results of analysing corrosion by X-ray imaging, where stainless steel and steel used in concrete have been analysed. In our study, standard commercial thick film resistors from different vendors were subjected to sulphuric atmosphere in two different corrosion experiment setups: single gas and flowers of sulphur. In each of the experiment setups, 90 resistors were electrically stressed, and 90 resistors were unbiased. A data acquisition (DAQ) unit was used to continuously measure the voltages and resistances of the DUTs during the experiments in order to observe any corrosion propagation. 2D X-ray imaging was implemented to provide deeper insight for assessing the propagation of the corrosion. Large differences in the resistors’ corrosion behaviour were observed between the vendors and between the experiment systems. It was also found that with X-ray imaging it was possible to detect the corrosion before it was observed in the electrical measurements.

II. METHODS

A. EXPERIMENT SETUPS

Two experiment setups, a flowers of sulphur (FOS) experiment, and a single gas experiment, were designed and built to induce corrosion in standard commercial thick film resistors. For both corrosions experiments, standards related to corrosion testing were followed. For the flowers of sulphur experiment, the EIA-977 [21] standard was followed. The single gas experiment was carried out based on IEC 60068-2-43 [14], with increased temperature and humidity to accelerate the corrosion.

Thick film resistors used as DUTs were soldered onto printed circuit boards (PCBs) specifically designed for the experiment. Experiments were carried out in both biased (Fig. 2a) and unbiased (Fig. 2b) configurations. Each PCB can hold up to 100 thick film resistors of which 90 (R1 to R90 in Fig. 2) in 9 groups of 10 resistors each were connected in parallel to its own PCB power supply terminals. Thus, 9 different vendors could be easily tested per test board. All thick film resistors were mounted with a 96.5 Sn - 3.0 Ag - 0.5 Cu solder and all had resistance values of 4700 $\Omega$ with a rated voltage of 150 V. The package size used was a standard 0805 package in all experiments and the electrical and structural specifications of the analysed thick film resistors are given in Table 1. The measured values in Table 1 were obtained with SEM and EDX.

For the FOS experiment, 2 boards were used with both having resistors from 9 different vendors. For each vendor, 10 thick film resistors were connected in parallel resulting in 90 DUTs in total per board. To assess the influence of voltage bias on the corrosion of the resistors, the first board was unbiased, and the second board was biased with a voltage of 10 V resulting in 2.5 V over the reference resistors and 7.5 V over the DUTs. For the unbiased board, the parallel resistance values of the 9 resistor groups were measured.
directly with an Agilent 34970A data acquisition unit equipped with Agilent 34901A multiplexers. For the biased board, the voltage over an external 160 Ω reference resistor outside the chamber, one per resistor group, was measured to electrically verify the propagation of the corrosion in the DUTs. In an event of a resistor open circuit, the change in voltage and/or resistance was recorded by the DAQ unit to enable a time-to-failure comparison between the different vendors. An additional unbiased FOS experiment was performed by using 10 DUTs of only one supplier where each resistance value of each single resistor was measured directly. In this experiment, X-ray images of the DUTs were obtained every 12h to analyse the corrosion propagation.

In the flowers of sulphur experiment, sulphuric powder CAS:7704-34-9 was used. The powder was placed in a petri dish and set at the bottom of a desiccator. Two desiccators were used in this experiment, one for the unbiased DUTs and one for the biased. Both desiccators were kept in a heat chamber at 105 °C for 750 hours, as specified in [21].

The single gas experiment was performed similarly as described for the FOS experiment, except no additional unbiased experiment with single resistors were performed. In the single gas experiment, the test PCBs were mounted inside a glass vessel placed inside a heat chamber. The temperature in the experiment (inside the heat chamber) was set to 60 °C, the relative humidity was set to 90 % and controlled with a separate vessel by a mixture of air and 68 °C water. The \( \text{H}_2\text{S} \)-concentration of 15 ppm was adjusted by a mixing of dry air and a bottled gas of nitrogen with 500 ppm \( \text{H}_2\text{S} \) using a mass balance meter and a rotameter. The total gas flow was adjusted to 3 l/min and a Dräger X-am 5000 gas detector was used to measure the \( \text{H}_2\text{S} \) concentration during the experiment. An illustrative drawing of the single gas experiment setup is shown in Fig. 3 and Table 2 presents the corrosion experiment specifications of each experiment.
In the single gas experiment the temperature influences the diffusion of water vapor and H$_2$S gas through the conformal organic epoxy layer. The corrosion does not commence before the gas mixture encounters the underlying silver metallization. Therefore, to define a suitable temperature for the single gas experiment, a diffusion model in a theoretical setting is described in Fig. 5. Here, the surface of the epoxy at $x = L$ is exposed to a well-stirred water-vapor-H$_2$S-mixture imposing one-dimensional mass diffusion in the $x$-direction, through the epoxy to the silver metallization at $x = 0$. The following assumptions apply:

- no chemical interactions occur,
- the temperature is uniform in all layers and
- the mass transfer between the epoxy and the silver metallization at $x = 0$ is negligible,

whereby the mass transfer within the epoxy is expressed as [22, p. 889]

$$\frac{\partial^2 C}{\partial x^2} = \frac{1}{D} \frac{\partial C}{\partial t}$$ (1)
where \( C \) is the mass concentration in consistent units (ppm), \( x \) is the distance (m), \( D \) is the coefficient of diffusion (m\(^2\)/s) and \( t \) is the time (s). For the initial uniform concentration, the following assumption applies [22, p. 890]

\[
C(x, 0) = C_i. \tag{2}
\]

To solve the equation (1), the following boundary conditions are needed [22, p. 890]

\[
\left. \frac{\partial C}{\partial x} \right|_{x=0} = 0 \quad \tag{3}
\]

\[
C(L, t) = C_a \quad \tag{4}
\]

where \( L \) is the thickness of the conformal epoxy layer (m). An exact solution to the gas concentration at the metallic silver layer \((x = 0)\) is of the form [22, p.257]

\[
\frac{C(0, t) - C_a}{C_i - C_a} = \sum_{n=1,3,5} \sin \left( \frac{n\pi}{2} \right) \exp \left[ -\left( \frac{n\pi}{2} \right)^2 \left( \frac{Dt}{L} \right) \right]. \tag{5}
\]

The effect of temperature was modelled as the Arrhenius relationship in the diffusion coefficient as [10]

\[
D(T) = D_{REF} \exp \left[ -\frac{E_a}{k \left( \frac{1}{T} - \frac{1}{T_{REF}} \right)} \right], \quad \tag{6}
\]

where \( D_{REF} \) is the diffusion coefficient (m\(^2\)/s) at the reference temperature, \( E_a \) is the activation energy (eV), \( k \) is the Boltzmann’s constant \( 8.617 \times 10^{-5} \text{ eV/K} \), \( T \) is the absolute temperature (K) and \( T_{REF} \) is the reference temperature (K). Table 3 presents the physical properties for H\(_2\)S and H\(_2\)O.

Table 3: Physical properties for H\(_2\)S and H\(_2\)O [10].

| Concentration | \( D_{REF} \) (m\(^2\)/s) | \( T_{REF} \) (K) | \( E_a \) (eV) |
|---------------|-----------------|--------|---------|
| H\(_2\)S       | \( 8.00 \times 10^{-13} \) | 340    | 1.060   |
| H\(_2\)O       | \( 1.53 \times 10^{-15} \) | 295    | 0.414   |

The temperature in the standard [14] for single gas experiments has been specified as 25 °C ± 2 °C. As can be seen
in Fig. 5 for the H₂S to reach the inner silver layer in the thick film resistors, it would take approximately 111 days to reach the level of 15 ppm H₂S. On the contrary, for a temperature of 60 °C it would take around 1.5 days. Therefore, it was not useful to perform a single gas experiment at the specified temperature as given in the standard [14]. It was also calculated that it takes four days for the H₂O to reach equilibrium at 60 °C. Wassermann et al. [1] have shown that increased humidity has an impact on the corrosion behaviour of electronics in a single gas testing, where a higher humidity generated larger dendrite structures under, otherwise, the same conditions. Therefore, a 90% relative humidity was chosen instead of 70% as specified in the standard [14].

C. ANALYSIS METHODS

X-ray images of the thick film resistors were obtained with an industrial Phoenix Nanomex X-ray machine, using both 2D and µ-computerized tomography (µ-CT) X-ray imaging technologies. For the scans, the maximum detail detectability of the open nanofocus x-ray tube was 200 nm [23].

In 2D X-ray imaging, a beam current of 150 μA and a voltage of 130 kV were used. All thick film resistors were imaged before and after the experiments to investigate the propagation of the corrosion. In µ-computerized tomography X-ray imaging, the DUTs were only imaged after the experiment since the DUTs needed to be cut out from the test PCBs to obtain high resolution images. In the µ-computer...
FIGURE 8. 2D X-ray images of both unbiased and biased DUTs (vendor D) taken before and after the FOS experiment. a) and c) images depicts the initial state of silver electrodes of the DUTs for both the unbiased and biased experiments, respectively. b) and d) depicts the corroded electrodes of two of the failed DUTs in the unbiased and biased FOS experiments, respectively.

To analyse the materials used in the end termination for each supplier, cross-section preparation techniques were utilized. For the cross-sectioning, the DUTs were encapsulated into epoxy for structural support during the grinding. The exposed surface was ground gradually by using silicon carbide grinding papers with grain sizes ranging from P320 to P4000. The final polishing was made by using woven acetate polishing cloths and aluminium oxide polishing films combined with polycrystalline diamond lubricants at grain sizes of 9 µm, 3 µm and 1 µm. Furthermore, the DUTs were polished with a JEOL cross section polisher to obtain a contaminant free DUT after the grinding process by using argon gas for the polishing. The flow rate for the argon gas was adjusted to 6.2 and an acceleration voltage of 7.0 kV was used in this study. The polishing process for each DUT was adjusted to 45 minutes. After polishing, the results of each thick film resistor were captured with a JEOL JSM-IT300 scanning electron microscopy combined with a Thermofischer energy dispersive X-ray spectroscopy (EDX).
III. RESULTS
A. STRUCTURAL INVESTIGATION

The thick film resistor end termination structure was investigated by cross-sectioning. All DUTs were ground and polished to the middle of the package. Fig. 6 shows an EDX and scanning electron microscopy image of each suppliers’ end termination structure after the polishing process. In all DUTs, tin and nickel were found in the top metallization and the protective barrier, respectively. Underneath the nickel layer, a silver layer was observed in all thick film resistors. The only difference between the suppliers was the thickness of the conformal organic epoxy layer.

To verify silver electrode corrosion in the thick film resistor DUTs, a JEOL JIB-4700F focused ion beam (FIB) with a gallium ion source was used. This technique allows for a contaminant free material identification of the inner structure of the analysed DUT. The final corrosion product was analysed by a JEOL energy dispersive X-ray spectroscopy.

B. TEST RESULTS

The time-to-failure (TTF) probability plot for the FOS experiment is shown in Fig. 7. The top row shows the TTF corresponding to the in-situ resistance measurements of the un-biased DUTs, whereas the bottom row shows the time-to-failure corresponding to the in-situ voltage measurements of the biased DUTs. Each step increase/decrease in the measurements indicates a DUT failure. As seen from the measurement results, the step changes (i.e. the failures), whether biased or unbiased, occur at similar times for each DUT type. Therefore, no differences in the time-to-failures probability could be observed, indicating that voltage biasing has little to no influence on the failure mechanism.

It is noted that in the biased experiment, once a DUT fails into an open circuit, the applied stress voltage over the remaining parallel connected DUTs increases slightly (and, as a result, the voltage over the reference resistor drops slightly). The maximum added stress voltage remains within 1.5% of the rated voltage of the DUTs and is assumed to have a negligible effect on the resistors.

Table 4 presents the obtained values from the Weibull probability distribution. The parameter $\beta$ for all vendors shows that the failure rate is increasing ($\beta > 1$) and results in a wear out failure due to corrosion for the DUTs. In Weibull analysis, parameter $\eta$ describes the time at which 63.2% of the thick film resistors have failed. $N$ presents the amount of failures in each experiment and $AD$ is the Anderson-Darling (AD) statistic, which measures how well the data follows the Weibull distribution. The $p$-value determines whether the data follows a normal distribution. Because the $p$-values for all vendors was greater than the significance level of 0.05, the decision was not to reject the null hypothesis and it cannot be concluded that the data does not follow a normal distribution.
Comparing DUT type H and E, both having similar end termination structures and conformal epoxy thicknesses, indicates an approximately ten-time higher $\eta$ for the H type. Likewise, comparing DUT type F with G and I, all having similar conformal epoxy thickness at the end termination, indicates an approximate a four-time higher $\eta$ for type F. Thus, based on the results, no direct correlation between the structure of the conformal epoxy layer and TTF in the corrosion experiments could be drawn. No failures were observed in the reference DUTs type B with anti-sulphur packaging.

X-ray images of an un-biased and a biased DUT taken before and after the FOS experiment are shown in Fig. 8. Fig. 8b) and d), both taken after the experiment of an un-biased and a biased DUT respectively, clearly shows severe corrosion of the underlying silver electrode resulting in open contacts. The corroded area is indicated by orange arrows. Shown in the obtained X-ray images, the unbiased and biased DUTs have failed due to the same corrosion failure mechanism. Together with the results from the time-to-failure data, it can be concluded that the voltage bias has negligible influence on the corrosion mechanism in thick film resistors failing in an open connection. Approximately 60% of the DUTs failed in an open connection and all were detected with 2D X-ray imaging. Furthermore, the progression of the corrosion was observed to be similar for all suppliers.

A sequence of X-ray images, taken at 0 hour until the open connection of the thick film resistor in the FOS experiment is shown in Fig. 9 and the corresponding online resistance measurements of the same DUT is shown in Fig. 10. The first sign of corrosion in the underlying silver layer is observed already at 24 hours in the X-ray image, even though no changes in the electrical measurements were observed. The corroded location is indicated by a red box in Fig. 9. The corrosion is seen as a small white area in the middle of the right-hand side end termination. Moreover, a denser material had appeared at the end of the end termination (seen as a black dot). After 36 hours, the propagation of the corrosion was easily observed as a bright stripe at the underlying silver layer, covering almost the full length of the end termination. Additionally, a dark line of denser material was now observed at the end of the top metallization. After 48 hours, only a small electrical connection remains, shown as a green arrow in Fig. 9. The only change in the electrical measurement was observed after 57 hours as the resistor failed in an open circuit indicated as a large step increase in the measured resistance shown in Fig. 10. In the following X-ray image
FIGURE 12. EDX point scan locations.

taken at 60 hours, a large area of the underlying silver layer had completely corroded.

To verify the results observed with the 2D X-ray imaging technique, a $\mu$-computerized tomography X-ray image of the failed DUT was taken and compared with results obtained with FIB of the same DUT. The FIB cut at the end of the top metallization of the end termination is indicated by the blue box in the $\mu$-CT X-ray image Fig. 11a) and b). As seen in the FIB image c) the underlying silver electrode has transformed into a porous corrosion product causing a change in the material density which can be detected with 2D X-ray imaging. The transition area of the porous corrosion product (red arrows) to the non-corroded silver layer (green arrow) is shown Fig. 11d) which corresponds to the red box in Fig. 11c). Additionally, multiple-point EDX scans were taken from the FIB cut with the corresponding atomic percentage of each element at the points shown in Fig. 12. Based on the EDX results, the elemental ratios at the corrosion location in point 1 is roughly 2:1, indicating that the underlying silver electrode has formed into silver sulphide by the reaction

\[
2\text{Ag} + \text{S}^{2-} \rightarrow \text{Ag}_2\text{S} + 2e^- (s). \tag{7}
\]

The protrusion at point 2 in the EDX measurements consists of silver sulphide which has originated from the underlying silver electrode. This protrusion phenomenon is caused by forces created by volumetric expansion of the corrosion product, which “pushes” the growth out from the interface between the conformal organic epoxy layer and the top metallization. This same protrusion was observed as a dark line of denser material at the end of the top metallization in the 2D X-ray images. Finally, point 3 shows the elemental analysis results at the un-corroded silver electrode.

In the single gas experiment, no increase or decrease in the electrical measurements were observed as shown in Fig. 13. Fig. 13a) presents the in-situ resistance measurements of the un-biased DUTs, whereas Fig. 13b) shows the in-situ voltage measurements of the biased DUTs. Although, no failures were observed in the electrical measurements, incipient corrosion was detected in multiple DUTs with 2D X-ray imaging. In Fig. 14, the red arrows depict the incipient corrosion in two thick film resistors from different vendors. The bright area at the end termination reveals the corrosion in the underlying silver electrode.
IV. DISCUSSION

Two different corrosion experiments were used to produce corrosion in thick film resistors. In both the FOS and single gas experiments, corrosion of the silver electrodes was observed. In the FOS experiment, the corrosion resulted in most DUTs failing in an open circuit during the 750-hour test time, whereas, in the single gas experiment, no open circuit failures could be observed during the experiment. Although no failures were recorded in the electrical measurements for the single gas experiment, incipient corrosion in the silver electrode was still detected with 2D X-ray imaging. The Ag$_2$S corrosion product observed in the DUTs was verified with FIB and the corrosion observed in the X-ray imaging corresponded well to the obtained results from the SEM image and the elemental analysis.

Both experiments are standardized, with only minor changes made to the temperature and humidity for the single gas experiment to accelerate the corrosion, only the FOS experiment was found to be applicable for a thick film resistor corrosion resistance evaluation in a practical testing time. The main reason for the different time-to-failure results is that the flowers of sulphur experiment is extremely aggressive, as shown in a comparison between single gas and flowers of sulphur setups in Ostermans et al. work [25]. In their work, it was shown that the corrosion thickness (nm/day) in copper coupons for a single gas experiment when using 1.8 ppm H$_2$S was 163.2 nm/day and in the flowers of sulphur experiment, 54094.2 nm/day. In our study, the amount of H$_2$S gas (15 ppm) in the single gas experiment was still too low and the test time was too short to obtain equivalent results as in the flowers of sulphur experiment. However, the single gas experiment should not be dismissed completely, as a biased single gas experiment with a higher humidity and H$_2$S gas concentration enables a reliable evaluation of thick film resistors in terms of resistance to electrochemical corrosion, as was shown in Fig. 15. In this case, X-ray imaging is not needed as the silver dendrite growth is optically observable.

Two major results were obtained in this study. Firstly, this paper clearly shows that 2D X-ray imaging is a powerful imaging technique for observing corrosion in thick film resistors. Contrary to most used methods for evaluating the
FIGURE 16. A failed device, where a corroded thick film resistor caused a critical process to stop. The corrosion can be observed in the 2D X-ray image shown with red dotted lines.

corrosion failures in thick film resistors, this technique can be used without destroying the printed circuit board assembly or single component. Furthermore, X-ray imaging was found to be a unique method for investigating the corrosion propagation in thick film resistors. From Fig. 9 and Fig. 10, it can be concluded that the change in resistance takes place instantaneously at the end of the corrosion propagation when the last contact point at either terminal finally corrodes open.

Secondly, the applied voltage bias of 7.5 V over the DUTs did not accelerate the corrosion. When comparing the results from the time-to-failure data, it can be concluded that the voltage bias has negligible influence on the corrosion mechanism in thick film resistors when failing in open connection. Furthermore, the failure signature from both biased and unbiased experiments showed identical corrosion progression, as shown in Fig. 8.

As an additional note, the failure mechanism produced in the corrosion experiments corresponds well to failures observed in the field where a device or a system has been exposed to a sulphuric atmosphere. Fig. 16 shows a device where a corroded thick film resistor caused a critical process to stop. As seen from Fig. 16, an open connection in the underlying silver electrode was easily observed with 2D X-ray imaging. The corroded area is shown with red dotted lines in the figure. Without X-ray imaging, it would not be possible to identify the root cause of the failure due to the black potting material on top of the surface mounted devices (SMD) of the failed device.

V. CONCLUSION

Two setups, a single gas and a flowers of sulphur experiment, were realized for evaluating corrosion progression in thick film resistors. The propagation of the corrosion was successfully examined by the implemented 2D X-ray imaging technique. FIB and EDX analysis were further utilized to verify the observed corrosion product.

In the single gas test, only incipient corrosion of the silver electrodes could be detected in the DUTs. However, optical analysis revealed an additional electrochemical corrosion failure mechanism in the form of dendrite growth on top of the conformal organic epoxy layer of the resistors.

In the FOS experiment, severe corrosion of the silver electrodes was observed in most of the DUTs. The corrosion occurred and progressed similarly, whether a voltage bias was applied over the DUTs or not. Hence, care must be taken when storing commercial thick film resistors, as exposure to a sulphuric environment may cause the eventual electrical device (end product) to fail already during commissioning.

In this study, the 2D X-ray imaging technique was successfully implemented to observe corrosion propagation in thick film resistors. This novel technique enables a fast and non-destructive investigation of corrosion in thick film resistors, as well as providing additional crucial information for further in-depth failure analysis. Moreover, for thick film resistor benchmarking and reliability assessment, this technique allows for a quick evaluation of the protective coating applied on thick film resistors. For thick film resistor benchmarking, only the flowers of sulphur experiment proved to be successful, and the resulting failure signature was found to correspond well with failed thick film resistors exposed to a sulphuric environment in the field.

REFERENCES

[1] T. N. Wassermann, O. Schilling, K. Müller, A. Rossin, and J. Uhlig, “A new high-voltage H2S single noxious gas reliability test for power modules,” Microelectron. Rel., vols. 100–101, Sep. 2019, Art. no. 113468, doi: 10.1016/j.microrel.2019.113468.

[2] C. F. Bayer, A. Diepgen, T. Filippi, C. Fuchs, S. Wustefeld, S. Kellner, U. Waltrich, and A. Schletz, “Electrochemical corrosion on ceramic substrates for power electronics—Causes, phenomenological description, and outlook,” in Proc. 10th Int. Conf. Integ. Power Electron. Syst. (CIPS), Stuttgart, Germany, 2018, pp. 161–167.
[3] D. Minzari, M. S. Jellesen, P. Moller, P. Wahlberg, and R. Ambat, “Electrochemical migration on electronic chip resistors in chloride environments,” *IEEE Trans. Device Mater. Rel.*, vol. 9, no. 3, pp. 392–402, Sep. 2009, doi: 10.1109/TDMR.2009.2022631.

[4] C. Hillman, J. Arnold, S. Binfield, and J. Seppi, “Silver and sulfur: Case studies, physics, and possible solutions,” in *Proc. SMTA Int. Conf.*, 2000, pp. 1–13.

[5] Rockwell Automation, Milwaukee, WI, USA. (2009). *Corrosion Resistance of Electric Wire Terminals Used in Harsh Industrial Environments.* Accessed: May 27, 2021. [Online]. Available: https://literature.rockwellautomation.com/idc/groups/literature/documents/wp/1492-44003_en-p.pdf.

[6] M. Watanabe, A. Hokazono, T. Handa, T. Ichino, and N. Kuwaki, “Corrosion of copper and silver plates by volcanic gases,” *Corrosion Sci.*, vol. 48, no. 11, pp. 3759–3766, Nov. 2006, doi: 10.1016/j.corsci.2005.12.009.

[7] H. Kim, “Corrosion process of silver in environments containing 0.1 ppm H2S and 1.2 ppm NO2,” *Mater. Corros.*, vol. 54, no. 4, pp. 243–250, Apr. 2003, doi: 10.1002/maco.200390053.

[8] T. E. Graedel, J. F. Franey, G. J. Gualtieri, G. W. Kammlott, and D. L. Malin, “On the mechanism of silver and copper sulfidation by atmospheric H2S and OCS,” *Corrosion Sci.*, vol. 25, no. 12, pp. 1163–1180, Dec. 1985, doi: 10.1016010-938X(85)90060-5.

[9] M. Reid, M. N. Collins, E. Dalton, J. Punch, and D. A. Tanner, “Testing method for measuring corrosion resistance of surface Mount chip resistors,” *Microelectron. Rel.*, vol. 52, no. 7, pp. 1420–1427, Jul. 2012, doi: 10.1016/j.microrel.2012.02.020.

[10] M. Reid, J. Punch, C. Ryan, J. Franey, G. J. G. Derkics, W. D. Reents, and L. F. Garfias, “The corrosion of electronic resistors,” *IEEE Trans. Compon. Packag. Technol.*, vol. 30, no. 4, pp. 666–672, Dec. 2007, doi: 10.1109/TPCAT.2007.917479.

[11] M. Cole, L. Hedlund, G. Hurt, T. Kiraly, L. Klein, S. Nickel, P. Singh, and T. Toft, “Harsh environment impact on resistor reliability,” in *Proc. SMTA Int. Conf.*, Orlando, FL, USA, 2010, pp. 1–9.

[12] M. Cole, J. Porter, J. Wertz, M. Coq, J. Wilcox, and M. Meilunas, “Effectiveness of conformal coatings in preventing resistor silver sulfide corrosion,” in *Proc. SMTA Int. Conf.*, Rosemont, IL, USA, 2016, pp. 937–944.

[13] P. Singh, L. Palmer, H. Fu, D. Lee, J. Lee, K. Guo, J. Li, S. Lee, G. Tong, and C. Xu, “A flowers of sulfur corrosion chamber for testing electronic hardware,” in *Proc. Pan Pacific Microelectron. Symp.*, Big Island, HI, USA, Feb. 2018, pp. 1–10.

[14] *Environmental Testing—Part 2–43: Tests—Test Kc: Hydrogen Sulphide Test for Contacts and Connections*, Standard IEC 60068-2-43, 2005.

[15] *Environmental Testing—Part 2–42: Tests—Test Kc: Sulphur Dioxide Test for Contacts and Connections*, Standard IEC 60068-2-42, 2005.

[16] K. Eguchi, T. L. Burnett, and D. L. Engelberg, “X-ray tomographic characterisation of pitting corrosion in lean duplex stainless steel,” *Corrosion Sci.*, vol. 165, Apr. 2020, Art. no. 108406, doi: 10.1016/j.corsci.2019.108406.

[17] S. M. Ghahari, A. J. Davenport, T. Rayment, T. Suter, J.-P. Tinnes, C. Padovani, J. A. Hammons, M. Stampanoni, F. Marone, and R. Mokso, “In situ synchrotron X-ray micro-tomography study of pitting corrosion in stainless steel,” *Corrosion Sci.*, vol. 53, no. 9, pp. 2684–2687, Sep. 2011, doi: 10.1016/j.corsci.2011.05.040.

[18] A. Česen, T. Kosec, and A. Legat, “Characterization of steel corrosion in mortar by various electrochemical and physical techniques,” *Corrosion Sci.*, vol. 75, pp. 47–57, Oct. 2013, doi: 10.1016/j.corsci.2013.05.015.

[19] P.-A. Itty, M. Serdar, C. Meral, D. Parkinson, A. A. MacDowell, D. Bjoqvist, and P. J. M. Monteiro, “In situ 3D monitoring of corrosion on carbon steel and ferritic stainless steel embedded in cement paste,” *Corrosion Sci.*, vol. 83, pp. 409–418, Jun. 2014, doi: 10.1016/j.corsci.2014.03.010.

[20] C. Wang, Y. Hua, S. Nadimi, W. Taleb, R. Barker, Y. Li, X. Chen, and A. Neville, “Determination of thickness and air-vold distribution within the iron carbonate layers using X-ray computed tomography,” *Corrosion Sci.*, vol. 179, Feb. 2021, Art. no. 109153, doi: 10.1016/j.corsci.2020.109153.

[21] *Test Method—Electronic Passive Components Exposure to Atmospheric Sulfur*, Standard EIA-977, 2017.

[22] F. P. Incropera and D. P. De Witt, *Fundamentals of Heat and Mass Transfer*, 3rd ed. New York, NY, USA: Wiley, 1990.

[23] Direct Industry. (2021). nanomez—The High-End 2D and 3D X-Ray Solution. Accessed: May 27, 2021. [Online]. Available: https://pdf.directindustry.com/pdf/inspection-technologies/nanomez/9257-365093.html.

[24] J. M. Ingman, J. P. A. Jormanainen, A. M. Vulli, J. D. Ingman, K. Maula, T. J. Kirkkainen, and P. Silventoinen, “Localization of dielectric breakdown defects in multilayer ceramic capacitors using 3D X-ray imaging,” *J. Eur. Ceram. Soc.*, vol. 39, no. 4, pp. 1178–1185, Apr. 2019, doi: 10.1016/j.jeurceramsoc.2018.10.030.

[25] M. Osterman. (Apr. 30, 2015). Corrosion concerns and testing for electronic equipment. University of Maryland, Collage Park, MD, USA. Accessed: May 27, 2021. [Online]. Available: http:// thor.inemi.org/webdownload/2015/Corrosion_Concerns_043015.pdf.

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