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On the effects of atmospheric particles contamination and humidity on tin corrosion

L. D’Angelo, V. Verdingovas, L. Ferrero, E. Bolzacchini, R. Ambat

Abstract—The effects of hygroscopic atmospheric particles are investigated in relation with corrosion of tin. Surface insulation resistance test boards were directly contaminated both with ambient particles sampled in the field at Milan (Italy) and with pure saline particles generated in the laboratory. An innovative particle deposition device was used to uniformly coat circular spots on to the test board surfaces. Deliquescence and crystallization of the water-soluble compounds were detected by observing the impedance response to varying relative humidity (RH) conditions with a gradual and continuous ramps. The effects of the adsorption/desorption kinetics and of the temperature on the deliquescence and crystallization relative humidity values were also investigated. Leakage current measurements at 5 V DC highlighted the ability of atmospheric particles to promote corrosion and electrochemical migration at RH levels far below condensing conditions (100% RH).

Index Terms—atmospheric corrosion, deliquescence, crystallization, leakage current, surface insulation resistance

I. INTRODUCTION

In order to satisfy the demand of smaller electronic devices, the component density on printed circuit boards has strongly grown in the last decades. Due to miniaturization and increase in the component density, the heat dissipation efficiency became ever important factor affecting the overall reliability of electronic devices. Elevated temperatures first have a significant influence on the reliability of semiconductor devices, secondly high temperatures of printed circuit board assembly (PCBA) can result in degassing of volatile corrosive compounds (fire retardants) from laminate material [1], [2], or cause a mechanical stress on the solder joints. Therefore, temperature of PCBA is often controlled by cooling, and one of the commonly used cooling methods is forced convection, causing increase of the airflow to the PCBA. On the other hand, this increases the exposure of the devices to moisture and contaminants [3], [4] and consequently the risk of metal corrosion.

Many researchers investigated the synergetic effects of relative humidity (RH) and contaminants on the PCBA. Warren et al. (1989) [5] observed electrochemical migration (ECM) at 76% RH on α-alumina substrates contaminated with CuCl2.

Frankenthal et al. (1993) [6] identified 75% RH (at 373 K) as the threshold condition above which the formation of Cu(\(SO_4\))\((OH)\)6 is observed for copper specimens contaminated with submicron-sized (NH4)\(SO_4\) particles. Verdingovas et al. (2014)[7] investigated the effect of NaCl contamination at different RH and temperature on surface insulation resistance (SIR) and reported a pronounced reduction in SIR above 75% RH at room temperature. Lately, Verdingovas et al. (2015)[8] pointed out the effects of five weak organic acids – namely adipic, succinic, DL-malic, glutaric and palmitic acids commonly used in no-clean fluxes – on the leakage current (LC), particularly on the RH thresholds above which LC increased abruptly. These studies highlighted the role of the deliquescence of the contaminants on the corrosion of metals. The deliquescence relative humidity (DRH) is the RH at which a solid-to-liquid phase-transition occurs; the reverse happens at the crystallization relative humidity (CRH), usually lower than DRH leading to a hysteresis behavior. This implies the formation of an electrolyte solution and the increase in the conductivity of the moisture film on the PCBA surface. As a consequence, the resulting increase in LC can compromise the function of electronic devices [9].

Pure contamination on the PCBA surface allows determination of threshold RH for safe operation; however, ambient contaminants present a more complex chemical composition for which the prediction of deliquescence RH is not easy. Considering the liquid or solid particles suspended within the troposphere (with a diameter between few nanometers up to hundreds of micrometers, i.e. atmospheric particles), their chemical composition varies in time and space because of the different sources and chemical reactivity [10–12]. The key role of these contaminants is due to the hygroscopic behavior of many compounds, mostly inorganic ionic ones, forming atmospheric particles, which enhances the formation of a thin water layer on the circuit surfaces. Concerning the fine fraction, i.e. PM2.5 (particles with aerodynamic equivalent diameter lower than 2.5 µm), many authors [13–15] showed that the water-soluble (WS) compounds represent an important mass fraction and the main species are NH4+, NO3-, SO4^2-, Cl-, Na+, Ca2+ and K+.

Recently, D’Angelo et al. (2016) [16] have studied the effect of PM2.5 chemical composition on the DRH and the reverse processes – the CRH – in Milan (Italy) during summer and...
winter conditions. The authors found that the wintertime’s nitrate-rich samples had lower DRH and CRH (55.2 ± 0.7% DRH and 46.9 ± 0.6% CRH) with respect to the summertime’s sulfate-rich samples (71.4 ± 1.0% DRH and 62.6 ± 1.2% CRH). Their results highlight the behavior of electrolytic solutions as a function of the RH history (at the atmospheric temperature) due to hysteresis behavior of atmospheric particles (CRH<DRH). In particular, during a humidification process (RH increase), particles are solid until RH reaches DRH; after that an aqueous solution is present. During dehumidification (RH reduction from a value above DRH) evaporation takes place until the CRH (<DRH) evaporation takes place until the CRH (<DRH) when crystallization occurs, leading once again to a dry state through a cycle of hysteresis.

These findings suggest that the electrolytic layer can form, and remain as liquid, far from condensing conditions. In this paper, the effect of atmospheric particles collected in Milan (Italy) during an intensive sampling campaign in March 2015 is investigated. The atmospheric contaminants were deposited on eleven hot air solder leveled SIR patterns and tested in a climatic chamber where impedance was measured in order to identify the DRH and CRH of the particles (Sect. III.A.1) similarly to the methods used elsewhere [17]–[19]. Moreover, the effects of the RH ramp rates and the temperature exposure on the impedance measurements are discussed (Sect. III.A.2 and III.A.3) and the LC at 5 V DC was measured in order to study the synergic effects of contaminants and RH on the corrosion of tin surface finishes (Sec. III.B and III.C).

II. METHODS

A. SIR contamination and particle’s sampling

The effects of atmospheric particles contamination on corrosion of tin was studied using test boards as shown in Fig. 1. The test board is made on a FR-4 laminate and contains surface mount components i.e. resistors and capacitors of various sizes, and 2 SIR patterns (13 mm x 25 mm area, 0.3 mm width of conductive lines and spacing in-between them) with hot air solder leveled surface finish using SN100C solder alloy.

The surface mount components on the test board are reflow soldered using ECO SOLDER M705-GRN360-K1-V solder paste (SAC305 alloy).

Prior to deposition of atmospheric or pure NaCl particles, the test board was cleaned using 2-propanol and rinsed with ultra-pure water (Milli-Q®, 18.2 Ω cm⁻¹ at 298 K).

A cascade impactor (Sioutas Personal Cascade Impactor, four impact stages, particle size cut-point: 2.5 µm, 1.0 µm, 0.5 µm, 0.25 µm, and a filtering stage) was modified in order to deposit a round uniform spot with a diameter of 2.0 cm, by means of a gear motor that allowed the complete rotation of the line-shaped nozzle in less than 6 min (Fig. 2). The airflow rate was set constant at 9 L min⁻¹ by means of a vacuum pump. The SIR patterns were housed at the lowest impaction stage (nominal particles size range: 0.25 – 0.50 µm) to deposit the finest fraction possible, which was shown to represent an important mass fraction within the atmospheric particles size mode [20], [21]. In addition, the finest particles are recognized being the most hazardous fraction due to the fact that filtering systems show low efficiency for them [3]. In order to avoid the deposition of liquid particles, the airflow was forced to pass through an airflow-tube dryer before entering the interior parts of the modified impactor, ensuring a complete dryness of the sampled particles.

Fig. 1 Test PCBA with SIR patterns and surface mount components used for testing. The insert of SIR pattern shows deposition of atmospheric particles using a cascade impactor, as shown in Fig. 2.

Fig. 2 Actual image of the cascade impactor followed by its detailed schematic. For atmospheric particle deposition, the SIR patterns were placed in Stage D.

At the same time, PM2.5 samples were collected (EN-14907) using the FAI-Hydra dual channel low-volume sampler (2.3 m² h⁻¹) on 47 mm PTFE filters (Millipore®, Ø=47 mm, PMP ring, 2 µm porosity). Filters were stored at low RH conditions (<15% RH) for 48 h, and weighed before and after the sampling campaign. Particle mass was determined using a precision weighting balance (Sartorius SE-2F, 0.1 µg precision).

Finally, the particle size distribution was determined using an optical particle counter (OPC Grimm, Model 1.107, “Environcheck”, time resolution: 5 min) which was able to count and classify airborne particles in 31 size classes from 0.25 to 32 µm. OPC data allowed to determine the degree of contamination on the SIR patterns (that were not weighted).

Moreover, the experimental approach described above (rotating impactor + gravimetric sampling) allowed at the same time to investigate the effect of ambient particles on: 1) SIR (via direct deposition), 2) capacitors (after WS extraction from
PM$_{2.5}$ filters, sect. II.B), 3) pure tin specimens (after WS extraction from PM$_{2.5}$ filters, sect. II.D), starting from particle samples having the same chemistry.

The sampling campaign was carried out between 19th March and 26th March 2015 in Milan; specifically, ambient particles were collected in a unique sampling site (urban background, with a full hemispherical sky-view) set up on the rooftop (10 m a.g.L) of the U9-building (45°30’38” N, 9°12’42” E) of the University of Milano-Bicocca, (Po Valley, Italy). The U9 site is equipped with a full-set of instrumentation for both particulate matter (see above) and meteorological measurements. With regard to particulate sampling, the instrumentation adopted here is made to collect particle samples on both test board with SIR patterns and Teflon filters simultaneously. In addition, it has allowed us to perform SIR, impedance, and electrochemical migration (ECM) tests and chemical analysis on the same ensemble of ambient particles. It is also noteworthy to mention that the atmospheric particle sampling activity has been a part of ten-year research programs to gather data related to DRH, CRH and chemical composition of the atmospheric particle contamination at wide range of field studies, namely: aerosol corrosion prevention, energy saving in data center [3] [16] [22], and preservation of cultural heritage places in Europe [23]. With these studies, we are able to understand the seasonal changes to the chemical composition of the atmospheric particulate matters (section IV).”

With regard to the meteorological data, RH was measured with a LSI-Lastem thermo-hygrometric sensor (DMA580 and DMA570 sensors, 5 min time-resolution, 10-98% RH range, accuracy ±2.5%, sensibility 0.2%). The acquired data was used to set the RH humidity ramp used in climatic chamber experiments (section II.B).

Finally, the effects of atmospheric particles were compared with pure NaCl particles. A 25000 ppm solution was prepared and a NaCl spray was generated by means of a Topas ATM 220 atomizer. The spray was introduced into a PTFE (poly-tetra-fluoro-ethylene) smog-chamber and collected on SIR patterns with the same set-up used to collect the ambient particles.

B. Climatic chamber and exposure conditions

Atmospheric particles and NaCl contaminated SIR patterns were tested in the ESPEC PL-3KPH climatic chamber, with specified temperature and humidity fluctuations below ±0.3°C±2.5%. The RH ramp was gradual and continuous in the range of 30-99% RH with a variation rate of 0.49% RH min$^{-1}$ at 298 K. Prior to the experiments, samples were stored inside a desiccator for 48 h at low RH condition (<15% RH). In the beginning of experiments, climatic chamber was set at 30% RH for about 30 min, while measurement equipment recorded data to make sure that the particulate contaminants were in equilibrium with the air conditions in the chamber.

The SIR patterns contaminated with atmospheric particles were exposed to three different RH ramp rates. The three ramp rates (0.49 % RH min$^{-1}$, 0.10 % RH min$^{-1}$ and 0.05 % RH min$^{-1}$) were chosen on the basis of different considerations. The 0.49 % RH min$^{-1}$ ramp rate is the same as used in previous experiments carried out in the Milano-Bicocca Aerosol Exposure Chamber and detailed in Ferrero et al. (2015) [22], Casati et al. (2015) [23] and D’Angelo et al. (2016) [16]. Thus, this ramp rate was firstly chosen in order to compare the obtained results with that reported in the aforementioned works. However, due to the importance of real exposure conditions the response for other two ramps (0.10 % RH min$^{-1}$ and 0.05 % RH min$^{-1}$) was investigated. Particularly, 0.05 % RH min$^{-1}$ represents the median value of the wintertime RH variation in the Po Valley (where samples were collected), while 0.10 % RH min$^{-1}$ is for a typical summertime humidity conditions. They were determined as the median value of the RH derivative with time observed in the Po Valley (Milan).

Thus, the chosen ramps are representative of real ambient conditions in which the ambient particulate matter samples were collected under field or outdoor use conditions. Most important is representative of real outdoor use conditions.

Further, three SIR patterns were also exposed to three RH cycles (rate 0.49 RH min$^{-1}$) setting surrounding air temperature at 281 K, 298 K, and 308 K (Sect. III.A.3).

As reported in the literature [3] [24]-[26] the aerosol hydration level depends on the relative humidity (RH) to which an aerosol particle is exposed in the atmosphere compared to the values of DRH and CRH in function of the RH history and in function of the given atmospheric temperature. Moreover, both DRH and CRH depend on particle chemistry (section II.D). It is important to note that several models present in literature (i.e. E-AIM, ISORROPIA) [27]-[28] allow us to calculate the DRH as a function of chemistry and temperature with a good degree of accuracy; however, present models [25] are unable to predict CRH in terms of whole particle complexity and temperature [22].

Thus in this present work, the whole dynamic complexity of the atmospheric contaminant particle behaviors with seasonal variations in the field use conditions is captured and results are presented for PCBA based tin corrosion.

C. Impedance and leakage current measurements

The formation of thin water-layer was investigated with “BioLogic VSP” potentiostat by measuring the impedance at 1 kHz frequency with 25 mV sinus amplitude. The impedance measurement at 1 kHz mainly corresponds to the resistance between the conductor lines, while the contribution from the capacitive component is minor and only for the dry SIR pattern. The use of 1 kHz frequency for monitoring conductive water film formation benefits from both higher frequency AC measurement minimizes the polarization effect at the electrodes, while the impedance corresponding to the resistance and not the capacitance results in more pronounced changes in impedance i.e. change of resistance due to formation of conductive water layer is more significant compared to the capacitance change related to the build-up of water layer due to deliquescence.

Data shown for impedance measurement are the mean values within a RH range of RH-0.5% ≤ RH < RH+0.5% (error bar represents the standard deviation of the signal variation within
the same RH range of RH-0.5% ≤ RH < RH+0.5%). Although Ferrero et al. (2015)[22] showed the effectiveness of the DC voltage in order to identify both DRH and CRH on PTFE filters, it is well known[17] that the AC impedance measurements with low perturbation voltage limits corrosion and avoids dendrites formations due to electrochemical migration (ECM) during measurements allowing the DRH and CRH to be accurately determined. In this respect, the DRH and CRH were determined using a gradient method as detailed in Ferrero et al. (2015) [22]. Particularly, DRH and CRH corresponded to the RHs at which the absolute value of the impedance gradient reached the maximum value (i.e. a sharp change in the electric signal); the corresponding DRH and CRH regions (DRHstart/end and CRHstart/end) were determined within the ranges of RH in which the gradient values deviated significantly from zero and contained its maximum.

This definition was validated in previous works [22-23] [16]. It is based on the observation that in impedance graphs (Fig. 3), along the humidification curve, a dry region is followed first by a deliquescent region (sharp decrease in impedance), and then by a hygroscopic growth region (due to continuous condensation of water); the reverse happens during dehumidification when evaporation took place, followed by crystallization (sharp increase in impedance), which in turn led once again to a dry state. The ECM is typically studied by applying constant voltage and measuring the LC between interdigitated electrodes [29]. Under these conditions, the “BioLogic VSP” potentiostat was used to measure the low current passing through a conductive layer at 5 V DC and as a function of RH (30.0-90.0% RH).

In addition to the response induced by particles deposited on SIR, the dendrite formation was investigated (under condensing conditions) on the ceramic chip capacitors (size 0805, Yageo Phycomp, Taiwan Type 2238 580 15649) contaminating them with WS compounds extracted from PM2.5 samples collected on PTFE filters (see Sect. II.D). They were diluted four times (from 1:2 up to 1:16), and 2.5 µL micro-droplets were placed on the ceramic chip capacitors (size 0805, Yageo Phycomp, Taiwan Type 2238 580 15649). Thus, five solutions for three PM2.5 samples were tested and LC was measured at 5 V DC and as a function of RH (30.0-90.0% RH).

D. Chemical characterization

Morphology of the corrosion products and tin dendrites after the LC measurements was characterized by means of scanning electron microscope (SEM JEOL 5900 instrument) and analyzed for elemental composition with energy dispersive X-ray spectroscopy technique (EDS Oxford Link ISIS).

In order to chemically characterize the WS fraction of the collected contaminants, atmospheric PM2.5 collected on PTFE filters were extracted in ultrasonic bath for 20 min (SONICA®, SOLTEC, Italy) in 3 mL of ultrapure water and the solutions were then recovered with a syringe and filtered (0.45 µm PTFE Syringe Filters, Phenomenex). The extraction efficiency was already investigated elsewhere [30]. The PM2.5 WS compounds were then analyzed with ion chromatography technique by means of Dionex ICS-90 (analytical column: Ion Pac CS12A-5 µm Analytical 3x150 mm, Dionex) for cations (Na+, K+, Ca2+, Mg2+, NH4+) and ICS-2000 (Ion Pac AS14A-5 µm Guard and Analytical 3x150 columns, Dionex) for anions (F-, acetate, formiate, Cl–, succinate, NO2-, SO42-, oxalate and PO43-).

The WS compound extraction was also carried out in order to evaluate the corrosion behavior of this fraction on pure tin. Similar to the volume extraction used for chemical characterization, the sampled compounds on the filters were extracted in 10 mL of ultrapure water in order to carry out at least three repetitions of the same measurements.

E. Micro-Electrochemical measurements

The corrosiveness of the WS particles was analyzed by performing potentiodynamic polarization measurements using electrolyte solution from collected contaminants. The overall inorganic ionic concentration of the tested solution accounted for 101.3 ppm. Prior to the start of the electrochemical measurement, the ionic strength of solution was measured by current measurement under applied potential of 5 V using a pair of closely spaced platinum electrodes immersed in 100 µL droplets of the solution. The results of the tests allowed identifying the strength of solution in terms of NaCl equivalent. The measurements were performed on a sample of pure tin, which is most commonly used metal for the surface finish on the PCBAs. The sample was cleaned with 2-propanol and ultrapure water prior to the testing. However, no polishing of the sample was done, in order to maintain the oxide layer naturally existing on tin finish surfaces on the PCBAs.

The potentiodynamic polarization measurements were performed using micro-electrochemical cell, which allows measurements using low volume of electrolyte. The detailed description of the cell can be found elsewhere [31]. Micro-electrochemical cell is designed for the measurements with high lateral resolution which is defined by the diameter of pipette tip, which makes the contact with a local region of the working electrode (1 mm in this current study).

The potentiodynamic polarization measurements (cathodic and anodic) were performed at a scan rate of 1 mV/s relative to Ag/AgCl reference micro electrode. Prior to the potentiodynamic polarization, the open circuit potential (OCP) measurements were performed. The potential stabilization was observed within first 5 min of the OCP measurement.

III. RESULTS

The degree of contamination of the SIR patterns was determined by means of data provided by the optical particles counter. Assuming a spherical shape, the particles volume concentration was calculated and log-normally distributed data were fitted in order to estimate the volume of particles with a diameter down to 30 nm. Once a complete volume size distribution was obtained in the range 30 nm – 32 µm, the cumulated volume of particles was used in order to estimate the PM2.5 density as the ratio between the mass of particle collected...
on PTFE filters (PM$_{2.5}$) and the calculated volume. The results indicate that the density of PM$_{2.5}$ fraction sampled in Milan during the campaign period was 2.1±0.1 g cm$^{-3}$.

Literature data [32–35] had shown that the particle density can vary from 0.6 g cm$^{-3}$ for freshly emitted diesel exhaust particles to 3.2 g cm$^{-3}$ for coal combustion particles. McMurry et al. (2002) [36] measured 0.1-0.3 µm sized particles density finding a mean range between 1.54 – 1.77 g cm$^{-3}$ for urban spherical hygroscopic particles in dry conditions, but density within the range 1.7 – 2.2 g cm$^{-3}$ were also observed. In this context, the obtained results agreed with the previous findings.

In order to estimate the mass of particles deposited on SIR patterns, density was assumed constant within the whole size distribution range of interest. Hence, the average mass of atmospheric particles collected was estimated to be 0.345±0.264 mg, which corresponds to 106.1±81.1 µg cm$^{-2}$ on SIR patterns. On the other hand, the contamination degree depends on the particles concentration in the atmosphere.

Impedance measurements carried out on SIR patterns contaminated with NaCl and atmospheric particles are shown in Sect. III.A in connection with the DRH and CRH of the contaminants.

A. The impedance responses of contaminated SIR patterns

In this section, the results regarding the effects of different exposure conditions on impedance response are shown for NaCl and atmospheric particles contaminated SIR patterns. In Sect. III.A.1, the conditions used by D’Angelo and co-authors [16] are reproduced, highlighting the role of the RH in impedance response. In Sect. III.A.2, the effect of the RH ramp rates is shown followed by the effect of exposure temperature in Sect. III.A.3.

A.1. The effects of RH ramps

The SIR patterns were contaminated with 0.345±0.264 mg of airborne particles, corresponding to 106.1±81.1 µg cm$^{-2}$. The contaminated SIR patterns were firstly exposed to increasing and decreasing RH ramps with the same setting conditions previously used in D’Angelo et al. (2016) [16] and described in Sect. II.B.

In Fig. 3 the example of three impedance responses are shown for a clean patterns (a), NaCl contaminated one (b) and atmospheric particles one (c). The presence of a contaminating agent substantially decreases the impedance with the increase in RH whereas for the clean SIR, the decrease is gradual and very weak.

For the SIR contaminated with NaCl (Fig. 3b), a steep decrease in impedance was observed within 74.0–77.0% RH corresponding to the DRH of NaCl. Above the DRH, water uptake still occurred and impedance decreased until 99% RH. On the opposite, with decreasing RH, the electrolytic solution formed on the SIR surface tends to decrease the conductive properties and a strong increase in impedance occurred between 59.0 – 52.0% RH, interpreted as CRH.

The atmospheric contaminants (Fig. 3c) caused a weak decrease in impedance even at low RH conditions due to an early water uptake as reported in D’Angelo et al. (2016) [16]. Further increase in RH caused a drop in impedance within the range of 55.0–63.0% RH. Above this range, impedance continued to decrease with the RH increasing, reaching an impedance values below 20 kΩ at 90.0% RH. During the decreasing RH period, firstly, a slow increase in impedance was observed followed by faster increase suggesting a recrystallization of water-soluble compounds within the range of 50.0–44.0% RH and then a complete water evaporation. An important conclusion is that ambient particles presented a hysteresis loop as pure salts making the presence of an electrolytic solution (and thus the hazard for PCBA) a function of the RH history.
A.2 The kinetic effects on hydration

Fig. 4 a shows the impedance profile of a SIR pattern contaminated with NaCl during RH cycles at different variation rates (Sect. II.B). Deliquescence was observed for the three RH cycles at 75.0% RH until 79.0% RH was reached. The impedance responses showed similar behavior, however, with lower impedance values for the slowest rate (0.05% RH min⁻¹), suggesting higher water uptake. The same hypothesis seems supported from the decreasing RH ramps, which clearly showed crystallization within 60.0 – 58.0% RH range. Above CRH range, the slowest cycle kept lower impedance values until 70.0% RH and then reached similar values as for 0.10% RH min⁻¹ rate.

![Fig. 4](image)  
**Fig. 4** Kinetic effects on impedance on SIR patterns contaminated by NaCl (a) and atmospheric particles (b). Full markers are for increasing RH ramp whereas open ones are for decreasing ramp. All measurements were conducted at 298 K.

Impedance response of SIR pattern contaminated with atmospheric particles is shown in Fig. 4b. The curves highlighted decrease in determined DRH and CRH with decreasing rate of RH ramping. The results showed that there was no difference in minimum impedance reached due to the effect of water uptake. Nevertheless, when comparing the impedance values for the fixed RH after deliquescence, lower impedance values were observed with decrease of RH variation rate.

A.3 The effect of the surrounding air temperature

In order to observe the effects of the surrounding air temperature on atmospheric particles deposited on SIR patterns, three RH cycles were carried out at different temperatures (Sect. II.B). Fig. 5 shows an impedance response during both the humidification (Fig. 5a) and dehumidification processes (Fig. 5b) for different temperatures. The results clearly highlighted that both DRH and CRH decreases with the increasing air temperature as expected [32]. The data analyses carried out on three SIR patterns contaminated with ambient particles suggested that there was a decrease in DRH of 0.4±0.1% RH K⁻¹ and in CRH of 0.5±0.2% RH K⁻¹. Thus, a second important conclusion is that a PCBA, contaminated by the same particles, should experience different degree of corrosion on different components with respect to exposure temperatures.

![Fig. 5](image)  
**Fig. 5** Impedance responses during increasing (a) and decreasing (b) RH ramp obtained at different temperatures.

B. The effect of atmospheric contaminants and RH on leakage current on SIR patterns

Leakage current was measured by applying 5 V DC voltage during a RH cycle from 30.0% to 90.0% RH and reverse with a single ramp duration of 2 h 20 min (rate 0.49 RH min⁻¹) at 298 K. In Fig. 6 the impedance response (Fig. 6 a) and the leakage current measurements (Fig. 6 b) of the same sample are shown. The SIR pattern was contaminated with an estimated particles mass surficial density of 270.6 μg cm⁻². The impedance response revealed that atmospheric contaminants enhanced water adsorption even at low RH conditions, which caused an increase in LC (Fig. 6 b). At the beginning of the experiment, measured LC was lower than 10⁻³ mA and it increased up to 10⁻² mA at 57.6% RH. Even though the starting RH of the deliquescence process was not clear, the impedance profile states that the end of this processes occurred at 64.0% RH.
Below this threshold, the increase in LC was already observed, but starting from 64.0% RH, the LC showed a further increasing trend. This suggests that water-soluble compounds dissolved and the electrolytic layer formed on the SIR surface allowing the conduction of charges and the electrolytic cell was active (corrosion of SIR pattern). The leakage current spikes seen in the graph can be explained by the formation of conductive bridges between the electrodes. The conductive bridges can be formed by aligning the atmospheric particles, and the tin corrosion products in a thin water film under the influence of electric field. These conductive bridges are forming and breaking off throughout the duration of the water film presence between the conductor lines on the SIR pattern.

At 71.8% RH a sharp increase in LC – from 5.3×10⁻² mA to 1.7×10⁻¹ mA at 72.1% RH – was detected. The maximum measured LC of 3.3×10⁻¹ mA was reached at 73.0% RH during the increasing RH ramp. Further increase in RH was not followed by an increase in LC which irregularly decreased instead and then exponentially diminished when RH was reduced inside the climatic chamber.

In order to compare the effect of atmospheric particle contamination with behavior of an uncontaminated SIR pattern, a cleaned SIR pattern was exposed to a longer RH cycle, i.e. 24 h per RH ramp. As Fig. 6 c shows, LC for cleaned SIR pattern was affected by changing in RH exposure conditions as well but with an exponential growth for the increasing RH ramp and with a reversing trend in the case decreasing RH ramp. Beyond the sharp increases in LC observed on the contaminated samples, the drop in current before the end of the increasing RH ramp suggested a different behavior in contrast to the clean SIR pattern.

C. The ECM probability in condensing conditions due to atmospheric contamination

The condensing conditions were simulated by placing a 2.5 µL droplet with WS compounds of atmospheric particles on ceramic chip capacitors (Sect. II.C).

The current on four chip capacitors in simulated condensing condition was firstly recorded in order to evaluate the effect of a pure water droplet. This showed that pure water on clean chip do not promote ECM and LC was observed to be less than 10⁻² mA. The contamination due to atmospheric WS compounds showed that LC easily reached values higher than 1 mA due to ECM processes. The dendrite bridge formation of the terminal of the chips was observed by means of a light optical camera during the experiment.

The contaminants content resulted to affect the probability of ECM occurrence (Fig. 7). In particular, the three tested solutions showed that the probability is not linearly correlated with the WS compounds content, but shows a bell-shape relationship.

D. Potentiodynamic polarization in aqueous electrolyte containing WS dust

Fig. 8 shows potentiodynamic polarization curves obtained in solution containing extracted compounds from WS atmospheric particles and a reference electrolyte containing NaCl at concentrations equivalent to 15.6 µg cm⁻² (based on relation 1 µl / 1 mm²) of NaCl. Polarization curves in electrolyte from WS particles showed a passive region followed by breakdown of the passive film. For the NaCl electrolyte, in contrast, the anodic polarization did not exhibit a clear breakdown of oxide film as characterized by significant...
increase in anodic current density. Current density oscillations during the anodic polarization were observed for both electrolytes, although it was more for the electrolyte from atmospheric particles. The corrosion potential was higher in electrolyte solution from atmospheric compounds, (-126 ± 11) mV vs (-198 ± 3) mV than in NaCl solution. The corrosion current density was also slightly higher in the electrolyte from atmospheric WS compounds (1.21 ± 0.31) × 10⁻³ mA cm⁻²) compared to NaCl solution (8.71 ± 3.87) × 10⁻⁴ mA cm⁻²). A pronounced increase in current density was observed for samples tested in electrolyte from atmospheric compounds above the breakdown potential indicating severe pitting of the surface.

Fig. 8 Potentiodynamic polarization of tin in electrolytes of NaCl and WS compounds of atmospheric particles.

IV. DISCUSSION

The results showed in sect. III.A highlighted the effect of RH exposure conditions on the impedance due to the hygroscopicity of NaCl salt and atmospheric particles deposited on the SIR patterns. Electrical methods were used by many authors in order to observe the phase-transitions of inorganic salts [7], [17], [19], weak organic acid [8], coarse dust [18] or atmospheric particles [23] deposited on different surfaces. The authors highlighted the effectiveness of this method to characterize the DRH of the contaminants. The resistance drop becomes dramatic when RH reaches the DRH since the contaminants form an electrolytic solution on the substrate surface. In fact, at DRH the Gibbs free energy of the solution state becomes equal to the energy of the salt solid state, thus a phase-transition from solid to aqueous solution take place for NaCl deliquescence occurs at 75% RH [26], perfectly in agreement with the observation presented in this paper. Under decreasing RH conditions, the opposite behavior was observed and the impedance showed an increase. According to D’Angelo et al. (2016) [16], Schindelholz et al. (2014) [17] and Ferrero et al. (2015) [22], when an abrupt increase happened, crystallization of NaCl salt occurred. In this regards, two observations were allowed by means of the data showed in Fig. 3. Primarily, evidence of the hysteresis behavior was seen due to the difference in the DRH and CRH humidity levels. This is because before crystallization, all the water needs to be evaporated from the solution and a high supersaturation is needed in order to crystallize. Nevertheless, the observed CRH was higher than the one commonly found in literature for homogeneous crystallization of NaCl particles, i.e. 45% RH [37]. This suggests the presence of crystallization nuclei, which increased the CRH of NaCl. Many authors, such as Han et al. (1999) [38] and Oatis et al. (1998) [39], pointed out the role of solid impurities, insoluble materials, which can act as nucleation seeds, in particular the metal oxides present in these seeds at significant levels. Thus, the obtained results point toward the fact that the AC signal and high NaCl contamination level caused little corrosion of tin, which will provide seeds for premature heterogeneous nucleation.

The analyses carried out on 9 SIR patterns contaminated with atmospheric particles indicated that the DRH occurred within 51.7±2.9% RH and 59.0±4.5% RH range at 298 K. Evaporation, which occurred while drying the air inside the climatic chamber cause reduction in the thickness of the water layer and increases the electrolytes concentration of the solution. Data showed that the evaporation increased the impedance until a sharp reduction was observed starting from 49.0±5.3% RH. This period, which ends at 41.5±4.8% RH, was identified as the RH range in which crystallization of the solid material occurs.

The results from ion chromatography (Sect. II.D) stated that the PM2.5 samples had a high content in nitrate (25.7±4.6% w/w) and a lower content in sulfate ions (12.1±0.6% w/w), with an average ratio NO₃⁻/SO₄²⁻ of 2.1±0.5. Inorganic ionic fraction accounted for 50.1±4.8% w/w of the total mass of particles collected on filters. In addition to nitrates and sulfates, ammonium ions (10.5±1.0% w/w) represented the main species found. Other cations (Na⁺, K⁺ and Mg²⁺) and inorganic anions (F⁻, Cl⁻, PO₄³⁻) accounted for less than 1% each. The organic acids were found representing about 1.1±0.4% of the mass. These findings allow to explain the agreement of DRH and CRH identified in this work with those provided in D’Angelo et al. (2016) [16]. In fact, the authors highlighted that PM2.5 samples with high NO₃⁻ contents showed DRH range between 51.6±0.7% and 58.5±0.7% RH while CRH was observed in average within 48.1±0.5 – 44.3±0.6 % RH range. Moreover, it has to be noticed that the three main components of the water-soluble inorganic fraction (the responsible for DRH and CRH behavior) were NO₃⁻, SO₄²⁻ and NH₄⁺ (explain the 94.3±1.5% of water soluble mass). Their concentration during the field tests was quite constant as demonstrated by the low standard deviation values reported here above, making the sample very close to each other as often happens in the Po Valley. This ensures the absence of significant deviations in the impedance and ECM measurement responses from sample to sample.

Note that the samples used for the kinetic and temperature tests (Sect. III.A.2 and 3) were not considered in the previous discussion since the data were collected on April 20th – 24th, 2015 and the chemical characterization showed a different composition. In fact, ion chromatography indicated that WS fraction was mainly constituted by sulfates (12.4±7.0% w/w) with significantly lower content in nitrates (2.8±0.5% w/w), resulting in average ratio NO₃⁻/SO₄²⁻ of 0.3±0.2. Such a high
content in sulfate ions leads to higher values of DRH and CRH [16]; comparing with the previous dataset, DRH and CRH (at 298 K and 0.49% RH min$^{-1}$) occurred at higher RH conditions, i.e., 78 – 81% RH and 64 – 58% RH, respectively.

A. The role of exposure conditions in impedance response

The formation of thin electrolytic layer and its conductive properties is mainly affected by the RH variation rate and the temperature of the surrounding ambient conditions (Sect. III.A).

This gives the reasons in defining a RH variation rate in order to simulate the most similar conditions of the circuits operating ambient. Thus, since lots of electrical equipment are nowadays exposed to outdoor conditions, a thermo-hygrometric sensor (DMA 572.1, LSI-Lastem S.r.l.) have been recorded RH and temperature of the air in Milan for seven months from March 2015 to September 2015 with a time resolution of 5 min. Relative humidity variation rate was then calculated and an average rate of 0.14±0.17% RH min$^{-1}$ was obtained. This means that in order to simulate a real RH cycle, a single RH ramp should last about 8 h 12 min.

On the other hand, many electronic systems are protected within electronic enclosures, which can make difficult for the heat dissipation and also slow down the variation rates of humidity in respect to the external conditions. In this regard, nighttime low temperatures can promote very high RH values or condensation within the electrical panel boxes and when the electrical circuits are switching off, there is no source of heat to keep the humidity below the condensation conditions. As a consequence, when circuits are switched on and current flows, humidity can affect electrical properties of the electrolytic layer depending on the air temperature.

In Sect. III.A.3, it was shown that there is a negative dependence between phase transitions RH of atmospheric particulate contaminants and air temperature. Many authors [40]–[42] highlighted the T-dependence of DRH for pure inorganic salts or mixtures. Although many evidences are available regarding the DRH-temperature dependence, few works were carried out to understand the effect of the temperature on the crystallization process. Zeng and co-authors (2014) [43] used an ATR-FTIR flow system in order to study the temperature dependence of methanesulfonate sodium salt (CH$_3$SO$_2$Na), one among the most abundant atmospheric methanesulfonate salt due to dimethyl-sulfide reactivity with OH radical [44]. They found a negative trend of DRH and CRH with temperature. The same trend was obtained for (NH$_4$)$_2$SO$_4$ but not for NaCl, which showed a negative trend for DRH and a weak positive trend for CRH.

Beside a lower DRH, a reduction in impedance values with the increase in temperature was recorded in agreement with findings in Song et al. (2013) [18]. Song and co-authors linked this behavior with the lower DRHs, which permit more moisture absorption and an increase in water layers at fixed RH, and with higher solubility of WS compounds forming atmospheric particles and corrosion products.

B. Evidences of corrosion below condensing conditions

Sect. III.B highlighted that high values of LC can be reached above the DRH of atmospheric contaminants. This suggested that ECM processes can occur even far from condensing conditions as confirmed by SEM-EDS investigations (Fig. 9 and Table I). Although no dendrite-like structures were observed, EDS revealed the presence of oxides/hydroxides and tin deposits, thus the distance between the conductive lines was shortened causing increase in LC.

![Figure 9 SEM image of the corrosion promoted by atmospheric particles and 5 V DC biased](image)

**Table I** EDS RESULTS OF CORROSION PRODUCTS REFERRING TO FIG. 9. ALL DATA ARE GIVEN IN WEIGHT PERCENTAGE.

<table>
<thead>
<tr>
<th>Spectrum</th>
<th>O</th>
<th>Br</th>
<th>Si</th>
<th>S</th>
<th>Cu</th>
<th>Sn</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>49</td>
<td>&lt;0.1</td>
<td>4</td>
<td>1</td>
<td>46</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>42</td>
<td>&lt;0.1</td>
<td>5</td>
<td>4.6</td>
<td>47</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>40</td>
<td></td>
<td>5</td>
<td>1</td>
<td>54</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>44</td>
<td>3</td>
<td>0.6</td>
<td>4</td>
<td>48</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>33</td>
<td></td>
<td>2</td>
<td>-</td>
<td>-</td>
<td>64</td>
</tr>
</tbody>
</table>

In Fig. 6 a temporary bridge seemed to be formed at 73% RH because of the steep increase in LC. On the other hand, at higher RH, LC decreased likely due to dendrites breakdown. Although high RH conditions were expected to further promote an increase in LC, with the beginning of the decreasing RH ramp, the electrical signal decreased with a similar behavior of the clean SIR pattern (Fig. 6 c). Such a similar effect could be ascribed to tin passivation. On the other hand, since atmospheric contaminants are characterized by a very complex chemical composition, a clear understanding of corrosion inhibitors formation was not carried out.

In this regard, EDS analyses (Table I) highlighted the presence of sulfur and oxygen forming the corrosion products. Moreover, ion chromatography carried out by the water-soluble compounds extracted from the PTFE filter – sampled simultaneously with the SIR pattern – showed many other chemical species both organic and inorganic compounds which can affect the corrosion process. Inorganic ionic fraction represents the 50.1% (mass percentage) of the collected particles mass and 1.1% is represented by mono- and dicarboxylic acids. The unresolved fraction (48.8%) was mainly
represented by elemental carbon, other not-analyzed water-soluble organic and hydrophobic organic compounds. In this regards, many authors investigated the chemical composition of organic matter within atmospheric particle fine fraction [45]– [47] in urban sites. Sannigrahi and co-authors (2006) [46] found that the main representative functional groups in aliphatic fraction (about 95% of water-soluble organic compounds) were alkyl and oxygenated alkyls (about 80%), carboxylic acids (about 10%) and aromatic functional groups (about 4%) were also found. This chemical composition makes difficult to the understanding of the processes involved in corrosion behavior induced by atmospheric particles and humidity.

C. Corrosion in condensing conditions

The WS contaminants of atmospheric particles resulted to promote ECM processes in condensing conditions, with dendrites formations in several cases (Sect. III.C). In this regards, Fig. 9 shows an image provided by means of SEM-EDS analysis highlighting a major contribution in tin (about 60% w/w), oxygen (<30% w/w) and titanium (<10% w/w, from the ceramic of the chip capacitor). The burn-off of the dendrites was likely due to the high current transported through them [48]. Both black and white corrosion products were observed on the ceramic chip capacitor surface. The same findings were described in Verdingovas et al. (2013) [49] due to the presence of tin hydroxides. EDS analysis also revealed the presence of other inorganic species, such as Na and Cl. On the other hand, further investigations are needed in order to identify the crystalline phases presented in the corrosion products. Ion chromatography technique was used to chemically characterize the WS composition of the solutions and the results are summarized in Table II.

D. Polarization curve and corrosion rate

Potentiodynamic polarization of tin in the WS atmospheric particles containing electrolytes indicated an influence of particles composition on the anodic dissolution and passivation of electrode. In electrolyte containing atmospheric particles, the initial anodic current density was lower, however, the breakdown of oxide layer was also more pronounced (potential when significant increase in current density is measured), as compared to NaCl electrolyte. Above the breakdown potential, a significant increase in anodic current by an order of magnitude was observed, and the resulting overall current density in anodic polarization was higher compared to NaCl. Pronounced fluctuations observed in the anodic polarization above breakdown potential indicated influence of compounds in the atmospheric contaminants on the re-passivation and anodic dissolution of tin. The latter observations also may play a role in electrochemical migration and influence probability for formation of tin dendrites. The electrochemical polarization and passivation of tin in electrolytes containing chlorides and other halides studied elsewhere [51]-[53], suggests importance of atmospheric particles composition on the corrosion behavior and passivation. The composition of atmospheric particles in an electrolyte solution will have an effect on both the anodic dissolution of tin, and the stability of tin corrosion products in electrolyte; thereby it will have an effect on probability and rate of dendrite formation.

The corrosion behavior of Sn-based solder alloys in NaCl and

![Fig. 10 SEM image of dendrite formed on chip capacitor contaminated with atmospheric WS compounds.](image-url)
acid electrolytes was also reported [53], [54]. Overall comparison of corrosion potential and corrosion current density between above literature studies and our current study indicates similar corrosion rates, although different influence on passivation and pitting of tin was observed. The corrosion potential of tin in electrolyte containing atmospheric particles was approximately 75 mV higher as compared to NaCl electrolyte. The atmospheric particles were characterized as more corrosive, due to low breakdown potential and resulting significant increase in anodic current density.

V. CONCLUSIONS

The effects of urban atmospheric particle contamination on PCBA were investigated at different ambient RH and temperature conditions. The deliquescence and the crystallization of water-soluble compounds were observed by monitoring the impedance responses while varying the RH. The existence of hysteresis of phase transitions between increasing and decreasing RH ramp was present in relation with the chemical composition of atmospheric particles.

The corrosive behavior of atmospheric contaminants on tin conductors was also investigated at 5 V DC. A strong increase in the LC was observed at RH values higher than deliquescence. Under condensation conditions, as simulated by placing a micro-droplet of electrolyte solution of dissolved atmospheric contaminants, promoted tin corrosion over chip capacitors through ECM process and their probability was affected by electrolyte concentration. Potentiodynamic polarization of tin showed that corrosion potential and the initial current density during anodic polarization in electrolyte containing ambient particles was significantly higher when compared to similar studies with pure NaCl. Also, the breakdown of oxide layer was more pronounced and an order of magnitude increase in anodic current was observed.

A practical implication of this study relates to the free-cooled data center operations under lowest carbon footprint levels as present in the literature studies and our current study indicates significant increase in anodic current density.

REFERENCES


