Protective Film Formation on AA2024-T3 Aluminum Alloy by Leaching of Lithium Carbonate from an Organic Coating

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An investigation into corrosion inhibition properties of a primer coating containing lithium carbonate as corrosion inhibitive pigment for AA2024 aluminum alloy was conducted. It was found that, during neutral salt spray exposure, a protective film of about 0.2 to 1.5 μm thickness formed within the area where an artificial defect was introduced by scratching through the coating to the base alloy. This film showed a multilayered structure consisting of a relatively compact layer near the alloy substrate, a porous middle layer and a columnar outer layer. The thicknesses of the layers varied, as a consequence of the difference in the local concentration of lithium species leaching from the primer coating. The presence of the film in the scribed area significantly reduced corrosion of the alloy, with little consumption of the metal substrate within the scribed area during neutral salt spray exposure. Furthermore, the presence of the protective film resulted in a significant increase of impedance modulus, measured after the salt spray exposure.

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The corrosion protection of aluminum alloys is critical for their successful application in the aerospace industry. Typically, several steps are employed to protect the alloy: surface cleaning and deoxidizing,1 conversion coating2,3 or anodizing,4 organic primer coating and decorative top coating.5–7 Cleaning and deoxidizing remove debris, lubricants, and near-surface deformed layers. The subsequent conversion coating or anodizing forms an extended oxide layer. The organic coatings control the access of the external environment to the underlying alloy and contain relevant functional components usually in the form of pigments. Where the coating is locally damaged, corrosion inhibitors, which are added into the organic primer as pigments, leach into the aqueous phase and react with the exposed alloy to form a protective film. Traditionally, strontium chromate is widely used as the corrosion inhibitive pigment in the organic primer for aerospace applications and has demonstrated to be very effective.5–8 However, chromium (VI) species have toxic effects on kidneys, liver and blood in addition to carcinogenic effects.9,10 Their use for corrosion protection applications must be limited or avoided, by finding suitable alternatives.

In order to develop suitable “green” active corrosion inhibitors, numerous research activities have been carried out since 1980s. Some of the most promising results were obtained with metal salts containing oxoanionic, e.g. molybdates,11–13 silicates,14 metavanadate,15–17 permanganate,18,19 phosphates,19–24 borate25,26 and tungstate.16 Rare earth27–35 and other cation compounds11,13,27–35 were also found to be effective. Organic coatings containing ion exchange inhibitors,36,37 metals,36,38,41 and organic inhibitive compounds42–46 were also found to increase corrosion resistance of metals.

In 1987, lithium containing aluminum alloys were developed for use in aerospace applications; this triggered the need to study the role of the lithium cations when present in a salt solution.37 It was found that a passive region becomes evident during anodic polarization of aluminum alloy 6061-T6 in 0.05 M Li2CO3 solution. Such passive region was not present when Li2CO3 was replaced by Na2CO3 or K2CO3.47 Lithium-based conversion coatings were developed in early 1990s.48–51 Using various lithium salts in alkaline solutions. Lithium cations were found to be intercalated into aluminum hydroxide to form insoluble layered double hydroxides.47–51 The protective properties provided by these lithium conversion layers are thought to originate from the formation of a lithium-aluminum hydroxide-carbonate-hydrate (Li3[Al2(OH)6]2 · CO3 · nH2O) layer,48,49 which greatly increases the corrosion resistance of various substrates, by increasing the pitting potential and reducing the corrosion current density.47–51

Although many approaches have been attempted to achieve acceptable alternatives, few systems have achieved the effectiveness of chromate-containing systems.39,46 A potential replacement for the chromate-containing pigments in organic coatings using lithium salts has been developed recently.32,33 Neutral salt spray testing (ASTM B-117) revealed that when such pigments are added into organic coatings, the corrosion resistance increases, resulting in comparable behavior to chromate pigments. In a previous study,30 the general filming behavior of the film formed on damaged area of AA2024 aluminum alloy coated with a lithium carbonate and lithium oxalate pigmented coating was investigated. It was found that lithium ions were able to leach from the organic coating, forming a film in the damaged area of the coated alloy. This film provided effective corrosion protection and comprised multiple layers, exhibiting barrier, porous and columnar morphologies.

In the present study, the evolution of such a film in the damaged area of AA2024 aluminum alloy coated with a lithium carbonate pigmented coating was investigated. It was found that the films formed in the damaged area are able to effectively protect AA2024 aluminum alloy from corrosion during neutral salt spray exposure and have varied morphologies in different locations. The mechanisms of film formation were investigated using advanced microscopic techniques and electrochemical impedance spectroscopy.

Experimental

Analytic grade purity lithium carbonate, from Sigma Aldrich was used as the corrosion inhibitive pigment, which was formulated into a model organic coating. The weights and sources of individual organic and inorganic compounds used in the coating are listed in Table I. The pigment volume concentration (PVC) of lithium carbonate was 15 v.%. The pigmented coatings were prepared according to the following procedure:

The individual components, including polyurethane polyol building block (Desmophen MPA), N-Butylacetate solvent, lithium carbonate, magnesium oxide, titanium dioxide and barium sulfate, used for Component A were added sequentially into a 370 ml glass jar

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Manuscript submitted September 8, 2015; revised manuscript received November 25, 2015. Published December 9, 2015.
with stirring, before 400 grams of Zirconox pearls (1.7–2.4 mm) were added to the mixture to aid dispersion. The mixture was then shaken for 20 minutes on a Skandex paint shaker to achieve a dispersed pigment particle size smaller than 25 μm. The Zirconox pearls were filtered after shaking. Component B that consists of polyisocyanates cross-linking agents (Tolonate HDB75MX) and an organosilane (Dynasilan Glymo) was mixed separately and then added to Component A. Finally, the mixture was stirred to a homogeneous state.

AA2024 aluminum alloy was anodized in tartaric-sulfuric acid according to aerospace requirements (AIPI 02-01-003) at Premium AEROTEC, Bremen, Germany. The primer coatings were applied to the anodized AA2024-T3 alloy with a HVLP (high-volume-low-pressure) spray gun. The coated alloy panels were then cured at 80° C for 16 hours. The thicknesses of the coating after curing were between 20 and 25 μm.

A mechanical milling device was used to artificially damage the organic coating on AA2024 aluminum alloy with a U-shaped scribe. The coated panels, with dimensions of 7 × 7 cm, were scribed from corner to corner, leaving a 1 mm wide scribe that penetrated to a depth of 100–150 μm into the metal. After scribing, the samples were exposed to the neutral salt spray test (ASTM-B117) chamber, for varying periods of time (4, 8, 24, and 168 h). After exposure, the samples were rinsed for 2 min with flowing deionized water to remove any residual chlorides and, subsequently, air dried.

After salt spray exposure for various times, the surface and cross sectional observation of the scribed area was carried out by scanning electron microscopy (SEM) using a ZEISS Ultra 55 instrument with an acceleration voltage of 0.5 kV. The cross sections of the scribed area were also examined by transmission electron microscopy (TEM) using a JEOL FX 2000 II instrument operated at 120 kV. The electron transparent TEM foils, of nominal thickness of 15–50 nm, were prepared using a Leica EM UC6 ultramicrotome with a diamond knife or by a focused ion beam technique with a FEI Quanta 3D SEM/FIB.

High resolution TEM and electron energy loss spectroscopy (EELS) was conducted on a TECNAI F30 TEM fitted with a Gif 2001 detector, operated at 300 kV.

Apart from the electron microscopy, electrochemical impedance spectroscopy (EIS) was also employed to evaluate the protective behavior of the lithium carbonate pigmented organic coatings using a three-electrode cell with a working electrode of exposed area 12.56 cm², a platinum counter electrode and a saturated calomel reference electrode (SCE). The testing cells were made by a cylindrical tube of 4 cm diameter, clamped to the testing samples with a rubber ring. Coated alloy samples with scribing before and after neutral salt spray exposure for 168 h were immersed in 0.05 M NaCl solution for 2 h. Impedance measurements were taken at the free corrosion potential using an Autolab-PGSTAT30 computer-controlled potentiostat, with 10 mV sine amplitude and a frequency range from 0.01 Hz to 3 × 10⁴ Hz.

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<th>Table I.</th>
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<td>Component A</td>
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<td>Bayer Materials science</td>
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<td>Lithium carbonate</td>
<td>Sigma Aldrich</td>
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<tr>
<td>Magnesium oxide</td>
<td>Sigma Aldrich</td>
</tr>
<tr>
<td>TiOxide TR 92</td>
<td>Huntsman</td>
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<tr>
<td>Blanc Fixe N (Ba(SO₄))</td>
<td>Sachtleben</td>
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<td>Component B</td>
<td>Supplier</td>
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<td>Tolonate HDB 75 MX</td>
<td>Vencorex</td>
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<tr>
<td>Dynasilan Glymo</td>
<td>Evonik</td>
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Figure 1. Scanning electron micrographs of the scribed area on AA2024-T3 before and after 168 h neutral salt spray exposure: (a) before; (b) after, with primer coating without lithium carbonate; and (c) after, with primer coating pigmented with lithium carbonate.

Results

Appearance of the scribed area before and after salt spray exposure.— Scanning electron micrographs of the scribed areas of the coated AA2024 aluminum alloy before and after salt spray exposure for 168 h are displayed in Fig. 1. The scribed areas of the alloy substrate, where the coatings were removed over a width of 1 mm, are in the center with the coated alloy substrate evident on the left and right sides. After the exposure to neutral salt spray conditions, a significant amount of corrosion products was formed on the alloy with a non-inhibitive coating (Fig. 1b), suggesting severe corrosion within the scribed area. However, when lithium carbonate was added into the coating, the scribed area appeared much cleaner (Fig. 1c), indicating effective protection within the scribed area by the coating.

Fig. 2 shows the surface and cross sectional views of the scribed area before salt spray exposure, revealing a mixture of shingling, cracking, and parallel scribing lines, as well as metal debris, resulting from the high levels of shear strain imposed during the scribing process (Fig. 2a). Cross sectional examination of the shingling within the scribed area showed partial detachment of metal from the surface, suggesting delamination of surface/near-surface regions as a result of the scribing (Fig. 2b). Away from the shingled region, the surface was relatively flat, as shown by Fig. 2c, which displays the cross section of the framed region in Figure 2a. Fig. 3 shows the surface and cross sectional views of the scribed area after salt spray exposure for 168 h. After 168 h salt spray exposure, for the primer coating without lithium carbonate, the initial scribe lines and shingles had disappeared. Instead, corrosion products were deposited on the surface, suggesting delamination of surface/near-surface regions as a result of the scribing. The corrosion products showed fine flake-shaped morphologies on the surface (Fig. 3b). Cross sectional examination (Fig. 3c–3e) of the area showed a thick layer of corrosion products, of about 25 μm thickness, in the middle of the scribed area (Fig. 3c). The majority of the corroded area showed a smooth alloy/corrosion product interface, indicating that active alkaline corrosion was the predominant process.
Pitting corrosion (Fig. 3d) and intergranular corrosion (Fig. 3e), associated with acidified environment at the corrosion front, were also observed locally in the cross sectional view. After 168 h salt spray exposure, for the primer that was pigmented with lithium carbonate, the scribed area showed a different morphology to that of the non-inhibited coating. Although the original surface features were still visible, they appeared less prominent (Fig. 4a). The scribed area showed bright and dark regions when studied with SEM (Figs. 4a and 4b). Dark regions of about 140 µm width were observed on the two sides close to the primer coating (Fig. 4a). In the center and near center region of the scribed area, the dark region was mixed with the bright region, with circular and irregular appearances (Fig. 4b). Figs. 4c and 4d show the regions A and B indicated in Fig. 4b, respectively, at increased magnification, revealing that the dark regions observed in Fig. 4b are associated with the presence of columnar/petal morphologies, while the bright regions exhibit a relatively flat appearance. Isolated oval or irregular features (as indicated by the dash-lined circles) observed in Fig. 4c are associated with the partial delamination of the alloy resulting from the mechanical scribing (Fig. 2b), or the presence of second phase particles beneath, as shown in Figs. 4f–4g.

Early stage of corrosion and filming behavior during salt spray exposure.—The early stage of the corrosion and filming behavior within the scribed area on the alloy with coatings in the absence and presence of lithium carbonate during salt spray exposure for 4, 8, and 24 h were also examined.

Coating without lithium carbonate.—After salt spray exposure for 4 h, localized corrosion was observed in the edge region of the scribed area, associated with a cluster of second phase particles. Away from the second phase particles, intergranular corrosion also developed deep into the metal substrate (Fig. 5a). In the center of the scribed area, corrosion products were evident across the surface. The thickness of the corrosion products was about 1–2 µm over the alloy matrix. Pores and cracks were also observed in the corrosion products (Fig. 5b). The corrosion product with porosity provides little corrosion protection. After salt spray exposure for 8 h (Figs. 5c–5e), corrosion products with similar features were observed (Fig. 5c). A dense barrier layer was formed closer to the alloy/corrosion products interface, with a layer

Figure 2. Scanning electron micrographs of the scribed area on AA2024-T3 aluminum alloy before neutral salt spray testing: (a) surface view; (b) and (c) cross sectional views of the near-surface region.

Figure 3. Scanning electron micrographs of surface (Figs. 3a and 3b) and cross sectional (Figs. 3c–3e) views of the scribed area on AA2024-T3 aluminum alloy after neutral salt spray exposure for 168 h in the absence of lithium carbonate in the primer coating: (a) surface view of the scribed area, showing significant corrosion products; (b) The corrosion products with fine flake-shaped morphologies; (c) cross sectional view, revealing the thickness of corrosion products being about 25 µm in the middle of scribed area; (d) cross sectional view showing pitting corrosion at local region; and (e) cross sectional view showing intergranular corrosion at local region.
Coating with lithium carbonate.—Cross sectional examination of the scribed area on the alloy with the lithium carbonate pigmented coating after salt spray exposure for 4, 8 and 24 h is displayed in Figs. 6a–6i, showing that a protective film formed within a few hours of exposure. Interestingly, there is little evidence of corrosion after exposure for 24 h. After 4 h exposure (Figs. 6a–6c), thin columnar layer has developed at the edge region of the scribed area (Fig. 6a). Furthermore, a porous outer layer and a dense inner barrier layer are evident near central region (Fig. 6b). The pores are larger at the outer region, showing a transition from porous to columnar/petal morphology. Barrier inner-layer with a thin porous outer layer was observed in the central region (Fig. 6c). After 8 h salt spray exposure, the film morphology is similar to that developed after 4 h, but the film became thicker, as shown in Figs. 6d, 6e, and 6f. Significant columnar and porous layer were developed close to the edge region of the scribed area (Fig. 6d) while porous/barrier film was developed near the center of the scribed area (Fig. 6e). In the central region, mainly barrier-layered film was developed (Fig. 6f). After 24 h exposure, a film with a columnar/petal morphology and almost no barrier inner layer was revealed at edge region of the scribed area (Fig. 6g). Moving away from the edge region, a three-layered morphology with significant columnar layer was observed (Fig. 6h). Near center of the scribed area, film developed a barrier inner layer, with a thin porous and columnar outer layer (Fig. 6i).

The composition and structure of the protective film.—The ultramicrotomed cross sections of the scribed area of the AA2024 alloy coated with the primer without lithium carbonate pigment before and after salt spray exposure were examined by transmission electron microscopy (TEM). Before the salt spray exposure, it is evident that a near-surface deformed layer, approximately 0.8–1 μm thick, with a sharp transition to the underlying, large grained, bulk alloy microstructure, was generated by the scribing (Fig. 7a). The deformed layer was observed previously when thermomechanical processing or mechanical polishing was carried out in aluminum alloys.56–60 The deformed layer microstructure is characterized by very fine grains, approximately 20–100 nm diameter (Fig. 7a). For the organic coating without lithium carbonate, after the salt spray exposure, the TEM image taken from the scribed area on the alloy shows a thin film at the alloy surface, with the majority of the corrosion products detached from the surface (Fig. 7b). The near-surface deformed layer is now absent from the alloy surface, indicating that corrosion completely consumed the near-surface deformed layer.

Fig. 8 displays TEM micrographs of the cross sections taken from the scribed area of the AA2024 alloy with lithium carbonate pigmented coating after salt spray exposure for 168 h, revealing thin films formed on the alloy surface. The cross sections were prepared by ultramicrotomy (Figs. 8a–8b) and focused ion beam (Fig. 8c) in various locations of the scribed area. The thicknesses of the films formed during salt spray exposure ranged from 0.2 to 1.5 μm. Fig. 8a reveals a barrier film in the center of the scribed areas. A film with significant columnar/petal outer layer, porous middle layer and barrier inner layer was observed near the edge of the scribed area (Fig. 8b). Fig. 8c shows a TEM micrograph obtained from the area between the edge region and the center of the scribed area. Interestingly, beneath the thin film formed during neutral salt spray exposure, the near-surface deformed layer introduced by the scribing was still present, indicating that corrosion of the alloy substrate was very limited and a fast, effective inhibition mechanism had operated. At increased magnification, area A indicated in Fig. 8c exhibits the three layer morphology (Fig. 8d), with lithium distributed in these layers, as shown by the corresponding EELS map (Fig. 8e).

Fig. 9a captured the film formed on the partially delaminated alloy (as indicated by the frame in Figure 4f), which shows a barrier inner layer, surrounded by the porous and columnar/petal layers. The EELS elemental maps of the film are displayed in Fig. 9b–9d, revealing the presence of lithium, oxygen and aluminum.

Fig. 10a shows a high-resolution TEM image of the film formed in the scribed area during salt spray exposure, revealing a mixture of
Figure 5. Cross sectional view of corrosion development in the scribed area on the AA2024 alloy coated with the primer without lithium carbonate after neutral salt spray exposure for 4, 8 and 24 h: (a)-(b) 4 h, revealing intergranular corrosion in the alloy matrix (a), and morphology of corrosion products (b); (c)-(e) 8 h, revealing the similar featured corrosion products and pitting corrosion (e); (f)-(g) 24 h, showing alkaline corrosion, pitting and intergranular corrosion.

Figure 6. Cross sectional view of protective film formed in the scribed area on the alloy coated with the lithium carbonate pigmented primer after salt spray exposure: (a)-(c) 4 h; (d)-(f) 8 h; (g)-(i) 24 h.
amorphous and nano-sized crystalline structures, as indicated by the dash-lined circles. Figs. 10b and 10c display two of the nano-crystals at increased magnification, exhibiting the lattice fringe images. The interplanar spacings and the angles between the planes measured from Figs. 10b and 10c are presented in Tables II and III. Comparing the measured values from a number of the lattice fringe images obtained from the nano-crystalline regions with the lattice parameters of the possible crystal structures presented in Tables II and III, it was evident that the characterization of the crystal structure is not conclusive since the measured lattice parameters in this study could not distinguish between the lithium-aluminum-carbonate-hydroxide hydrate version of hydrotalcite,49,63 \( \alpha \)-LiAlO\(_2\)61,62 and Al(OH)\(_3\).64,65

Electrochemical impedance spectroscopy.— The corrosion resistance of the film formed in the scribed area during salt spray exposure of the alloy coated with the lithium carbonate pigmented coating was evaluated by electrochemical impedance spectroscopy (EIS). The samples were immersed in 0.05 M NaCl before and after salt spray exposure; this represents the two conditions of the scribed area, i.e. with and without the thin film formed during the salt spray exposure. Further, a reference sample coated with the organic coating without lithium carbonate pigment was also evaluated by EIS after neutral salt spray exposure. From the bode plot of the impedance modulus (Figure 11a), it is evident that the lithium carbonate containing sample displays low-frequency impedance values that are about one order of magnitude higher than those measured from other two samples at low frequency, i.e. the lithium carbonate containing sample before salt spray exposure, and the reference sample coated with the coating without lithium carbonate pigment after salt spray exposure. The impedance modulus spectrum acquired for the specimen with lithium containing pigments after salt spray exposure also displayed higher values of impedance in the medium frequency range, which is due to the formation of a protective film. The Bode phase diagram (Figure 11b) of the sample with lithium carbonate pigmented coating before salt spray exposure shows clearly the time constant of the natural oxide film in the scribed area at the

Figure 7. Transmission electron micrograph of the ultramicrotomed cross sections of the scribed area of the AA2024 alloy coated with organic coating without lithium carbonate pigment examined by transmission electron microscopy: (a) before exposure to the salt spray, the near surface region revealed a deformed layer; and (b) after exposure to salt spray for 168 h, the deformed layer is absent. The samples were prepared by ultramicrotomy.

Figure 8. TEM cross sectional examination of the film formed in the scribed area of AA2024 alloy coated with lithium carbonate pigmented coating after 168 h salt spray exposure: (a) the film with barrier layer; (b) the film with the columnar/petal layer, porous layer and barrier layer; (c) the three layered film and the near surface deformed layer introduced by scribing; (d) area A indicated in (c) at increased magnification; (e) corresponding EELS map of (d), showing the presence of lithium in the film.

Figure 9. (a) Transmission electron micrograph of the film formed on the partial delaminated alloy, as indicated by the frame in Figure 4f; corresponding EELS elemental maps of the film: (b) lithium map; (c) aluminum map; and (d) oxygen map.
Figure 10. (a) TEM image of the film formed in the scribed area during salt spray exposure, revealing a mixture of amorphous and nano-sized crystalline structures; (b)-(c) lattice fringe images of the nano-crystals.

frequency at or above \( \sim 10 \text{ Hz} \). After salt spray exposure the sample shows an increase and broadening of the phase, again indicating that a thicker and stable film was formed in the scribed area during the exposure. On the other hand, for the sample coated with primer without lithium carbonate, a broad but lower phase range with lower impedance modulus was observed. These results confirm that the formation of the lithium containing film on the scribed area during the salt spray exposure resulted in an improvement of the corrosion resistance, with a significant increase in the overall value of impedance compared to the reference samples.

**Discussion**

Direct observation and comparison of the scribed area on the coated AA2024 aluminum alloy before and after salt spray exposure for 168 h in Fig. 1 have revealed the significant role of lithium carbonate

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<th>Table II. Interplaner d-spacings measured from the lattice image in Figure 10b and the lattice parameters of the possible crystal structures.</th>
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<th>Table III. Interplaner d-spacings measured from the lattice image in Figure 10c and the lattice parameters of the possible crystal structures.</th>
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as leachable inhibitor in the organic coating. This has led to a detailed investigation of the protection mechanism of the alloy by lithium carbonate as corrosion inhibitor, as discussed below.

Comparing Figs. 2a–2c with Figs. 3a–3e, it is evident that extensive corrosion had developed in the scribed area when lithium carbonate was absent, which changed the surface appearance in the scribed area and developed relatively thick corrosion products. Such products are mainly aluminum hydroxide. Such hydroxide showed low protective properties, as suggested by electrochemical impedance results. On the contrary, for the alloy coated with the lithium carbonate loaded coating, the scribed area showed little evidence of corrosion after the neutral salt spray exposure. Examination of this region revealed that a thin film, containing lithium, aluminum and oxygen, was formed on the alloy surface during the salt spray exposure. The presence of lithium species in the film indicates that lithium species are able to leach out from the primer coating to the scribed alloy surface and form a stable protective film that protects the alloy from subsequent corrosion. Furthermore, such films provide corrosion protection, as evident from EIS measurement after neutral salt spray exposure.

From the observation of the filming behavior after exposure to salt spray for 4, 8, 24, and 168 h, it is evident that the film formation commenced in the early stage of neutral salt spray exposure. With increasing exposure time, the film thickens with significant development of a columnar/petal layer on the outer region, which may be due to the increased lithium species available for the film formation. On the other hand, the film thickness was influenced by the local cathodic/anodic activities, especially the presence of cathodic CuAl₂ particles.

The morphology of the protective film showed three distinct layers, containing a barrier layer close to the alloy interface, a porous central layer, and a columnar/petal outer layer. However, the relative thickness of the three layers varied. Some regions developed a significant barrier layer while the others developed columnar/petal layer with little barrier and/or porous layer. Such varied morphology may be related to local lithium concentration and local pH. High lithium concentration and high pH may result in the formation of mainly columnar/petal morphology. Adjacent to the primer coating, where relatively high concentration of lithium species is expected, a significant development of the columnar/petal morphology was observed.

In the center of the scribed area, the pathway for leaching is increased and the concentration of lithium species may be relatively low, consequently, barrier-type morphology was formed. On the other hand, the film morphology at local regions may be altered due to the redistribution of the lithium species by the stream of the salt sprayed to the sample surface, or by the variation of local pH due to local anodic and cathodic activities associated with the different phases in the alloy or the individual grains. The grains with relatively high stored energy might display more anodic activities than that of grains associated with relatively low stored energy. A schematic diagram of the various film morphologies is illustrated in Fig. 12, categorizing five main types of the morphologies observed at different locations in the scribed area. These include high columnar/petal layer with little barrier and porous layers, significant columnar/petal layer with the presence of porous and barrier layers, significant presence of three layered morphology, porous and barrier layers, and barrier layer. Protrusion of the film can be correlated to the scribing-introduced partial delamination of the alloy, the presence of a second phase particle beneath the film, or the presence of the metal debris. The TEM image of the cross section of the alloy substrate in the scribed area revealed the presence of a near-surface deformed layer of about 0.8 to 1 μm thickness, as shown in Fig. 7a. The presence or absence of the near surface deformed layer provides a handy tracer to measure the consumption of the alloy during corrosion and filming in the scribed area. In Fig. 8c, a substantial deformed layer of 0.9 μm thickness was still present, suggesting that the consumption of the alloy due to the initial corrosion or filming was very limited.

EIS measurements after salt spray exposure confirmed that the presence of lithium carbonate in the primer coating results in the formation of a protective film in the scribed area. Such film formed a physical barrier, resulting in a considerable increase in the values of low-frequency impedance.

Conclusions

(1) During salt spray exposure of the AA2024 aluminum alloy coated with a lithium carbonate pigmented primer, lithium species were able to leach out from the primer and to migrate to the damaged area. The leached lithium species reacted with aluminum hydroxide
and formed a film of about 0.2 to 1.5 μm thickness on the alloy surface within the damaged area. The morphology of the film varied, with the regions close to the primer coating showing a significant outer columnar/petal layer, and the central region of the scribed area exhibiting mainly a barrier film. The variation of the morphology of the film might be associated with the local concentration of the leached lithium species.

(2) The film is a mixture of amorphous and nano-sized crystalline materials. The lattice fringe images of the nano-crystals suggested that Li2Al2CO3(OH)2−3·3H2O and/or α-LiAlO2 and/or Al(OH)3 are present in the film.

(3) The film formed in the damaged area was able to effectively protect the AA2024 aluminum alloy from corrosion during neutral salt spray exposure. Beneath the film, the presence of the near-surface deformed layer of approximately 0.8–1 μm thickness, introduced by scribing, indicates a fast and effective inhibition by the leaching process.

(4) Electrochemical impedance spectroscopy showed that such film can form a physical barrier between the bare metal and the aggressive environment, as evident from the presence of an additional time constant and the increase in the impedance at low and medium frequencies.

Acknowledgment

The authors thank AkzoNobel for sponsoring this work.

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