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Electrochemical Behavior and Self-Sealing Ability of Zirconium Conversion Coating Applied on Aluminum Alloy 3005 in 0.5 M NaCl Solution

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Chemical conversion coatings are formed by immersion, spraying, rinse or no-rinse process in which metal reacts with ions/substance from conversion bath forming an insoluble oxide-based coating at the surface of a metal. They are usually inorganic in nature. The main purpose of these coatings is to improve the adhesion of organic paints and lacquer to the substrate; to facilitate adhesion, organic polymers and additives are often added. Additionally, conversion coatings should improve the corrosion resistance of the substrate and the whole painted system.

For the last 100 years, state of the art conversion coatings were chromium conversion coatings (CCC) for aluminum/light metals and phosphate conversion coatings for ferrous metals. As CCCs are nowadays being replaced due to carcinogenicity and phosphate conversion coatings are reconsidered due to environmental and economic issues, there is a great need to find a comparable, economical and green conversion processes. The most promising and already commercialized processes are the zirconium conversion coatings (ZCCs), so-called Zirconization, and the trivalent chromium process (TCP). Zirconization has developed very quickly since the 1st generation ZrCCs in 2002 and 2nd generation in 2010. Zirconization coating the corrosion resistance in NaCl solution is much thicker at/in the proximity of IMPs. Zinc, steel, and many other metals can also produce thicker conversion coatings. Addition of a small amount of Cu speeds up the conversion process and produces thicker coating. In addition, while a small amount of Si can also produce thicker conversion coatings. Surface pre-conditioning has an important role in the microstructure and properties of ZrCCs. Additionally, surface pre-treatment influences acid-base properties and the amount of OH⁻ groups in the final ZrCC. Coating thickness increases with the agitation of the bath by factor 2–4 depending on the substrate. ZrCCs are usually 30–80 nm and were reported to protect a bi- or even tri-layer structure. ZrCCs improve adhesion of top organic coats, reduce the rate of cathodic delamination and improve corrosion resistance on different substrates. Addition of corrosion inhibitors such as Ce, V, Mo into ZrCC can provide a certain degree of active corrosion protection to these coatings.

Aluminum-manganese alloy AA3005 has good workability, medium strength, good electrical and thermal conductivity and excellent corrosion resistance. It is used as a heat exchanger in automotive and other industries, in architectural industries, for brazing and cladding, etc. There have been several studies on the nature and stability of Al₂O₃ formed on 3xxx alloys, but only a few studies about conversion pre-treatment of this alloy. Smit et al. studied no-rinse pre-treatment of AA3003 in conversion bath containing 0.1 M solution of H₂TiF₆. When coated with titanium conversion coating the corrosion resistance in NaCl was initially not so good but it improved with immersion time. TEM images showed that the coating was 30 nm thick. Thermal treatment had a beneficial effect on the corrosion properties of H₂TiF₆-pre-treated...
AA3003. Banczek et al. studied Ce conversion coatings on AA3003 and compared it with a commercial TCP conversion coating (SurTec 650 Chromital).

Therefore, there is a lack of studies about zirconium conversion coatings applied to AA3xxx alloys. Moreover, the extraordinary behavior of titanium conversion coating on AA3003 which improved with time, as noticed by Smit et al.56 deserves further mechanistic understanding. In our previous study, the commercial TCP coating SurTec*650 containing Cr(III) and Zr deposited on AA3003 was studied.72 This commercial coating exhibited good corrosion resistance in simulated acid rain and sodium chloride solution but only after longer immersion in solution, i.e., 24 h. This interesting behavior is studied in more detail in this study devoted to Zr-based coatings only deposited over AA3005. The electrochemical behavior of ZrCCs prepared by immersion of AA3005 in the conversion bath of hexafluorozirconic acid (H₂ZrF₆) was investigated.

Microstructural characterization of the coated substrate was carried out using scanning electron microscopy equipped with energy X-ray dispersive spectrometer (SEM/EDS), cross section focus ion beam microscopy (FIB), transmission electron microscopy (TEM), X-ray photoelectron spectroscopy (XPS) and time-of-flight secondary ion mass spectrometry (ToF-SIMS).

Electrochemical characterization of samples was carried out in a 0.5 M NaCl solution by measuring open circuit potential (OCP) vs time curves, potentiodynamic polarization curves and electrochemical impedance spectra. Special attention was devoted to the long-term corrosion resistance in NaCl, up to 40 days. An interesting behavior of improvement of corrosion characteristics of coated samples with time was confirmed. Based on the results obtained we were able to address this phenomenon and postulate the mechanism of self-sealing or active protection of Zr-conversion coatings on Al-Mn alloy AA3005.

**Experimental**

*Materials, sample preparation and calculations.*—Aluminum alloy AA3005 / UNS-A93005 (Mn 1–1.5 wt%, Mg 0.2–0.6 wt%, Fe 0–0.7 wt%, Si 0–0.6 wt%, Cu 0–0.3, Zn 0–0.25 wt%, Cr 0–0.1 wt%, Ti 0–0.1 wt%, residuals 0.15 wt%) in form of foil (thickness 70 μm) produced by Impol 2000 d.d. (Slovenska Bistrica, Slovenia) was used as a substrate. The composition given was provided by the supplier. Samples used in experiments were cut in a rectangular shape with dimensions of 3 cm × 3 cm.

The following chemicals were used for experiments: NaCl (for analysis, Fischer Scientific, Leicestershire), H₂ZrF₆ (50 wt% in water, Sigma-Aldrich, Saint Louis, USA), NH₄HCO₃ (reagent grade, Sigma-Aldrich, Steinheim, Germany) and Metaclean® (alkaline cleaner, Amity international, Barnsly, United Kingdom). Milli-Q Direct water with the resistivity of 18.2 MΩ·cm at 25 °C (Merck, Darmstadt, Germany) was used for rinsing and for solution preparation.

pH was measured using pH meter 827 pH-lab equipped with Unitrode/Pt1000 (Metrohm AG, Herisau, Switzerland). Teflon beakers (V = 1 L) were used for conversion baths, while glass beakers (V = 600 ml) were used for alkaline cleaner and rinsing baths. Sample holders were made of teflon and conversion coating solutions were stored in polyethylene bottles.

**Preparation of conversion coatings.**—Conversion bath contained 1 · 10⁻³ mol L⁻¹ (200 ppm) of H₂ZrF₆ and pH was set to 4.8 using 15 wt.% NH₄HCO₃. The conversion was carried out at room temperature.

Samples were chemically cleaned by immersion in 5 wt.% Metaclean solution (T = 57 °C, stirring rate 150 rpm) for 3 min. Immediately afterward, samples were washed off with Milli-Q Direct water and then rinsed by dipping in Milli-Q Direct water bath for 45 s. Immediately afterward, the sample was dipped in a conversion coating bath (T = 25 °C, stirring rate 450 rpm) for 10 min. Immediately afterwards the sample was rinsed with Milli-Q Direct water and washed off by dipping it in Milli-Q Direct water bath for 1 min. The sample was then dried with N₂ stream and left to dry in air for 24 h.

As-received (bare) samples refer to AA3005 used in as-received conditions without any chemical cleaning.

**Electrochemical characterization.**—Electrochemical experiments were carried out with a Multi Autolab/M204 (Metrohm Autolab, Utrecht, Netherlands) potentiostat/galvanostat controlled by Nova 2.1 software. Electrochemical measurements were repeated for each condition for at least three different samples. Representative measurements are presented in plots and average values together with standard deviation in tables. Measurements were conducted in a standard three-electrode flat cell with a volume of 250 mL. The sample was the working electrode, a carbon rod was the counter electrode, and a saturated Ag/AgClₑₛₜₑₐₑₜ electrode was the reference electrode (E_Reference = 0.198 V). All potentials in the text refer to the Ag/AgClₑₛₜₑₐₑₜ scale. The electrolyte was freshly prepared 0.5 M NaCl.

Prior to electrochemical measurement samples were allowed to rest at open circuit potential (OCP) for 5 min, 1 h, 2 h, 1 day, 2 days, 3 days, 4 days, 5 days and 10 days.

Electrochemical measurements were conducted in the following sequence from least destructive to more destructive: (i) measurement of the OCP as a function of time during for 5 min, (ii) recording of linear polarization curves, and (iii) recording of potentiodynamic polarization curves. The system was allowed to rest for 10 s before switching to the next electrochemical technique.

Linear polarization curves (LPR) were run in the range from —10 to —1 mV vs OCP using a 0.1 mV s⁻¹ scan rate. Polarization resistance (R_p, LPR) was determined as the slope of the fitted potential (E) vs current density (j) curve using Nova 2.1 software.

Potentiodynamic polarization curves measurements were made in the potential region starting from —250 mV vs OCP in the anodic direction until current reached 0.1 mA. The scan rate was 1 mV s⁻¹.

Corrosion current density and corrosion potentials were determined using Nova 2.1 software based on the Tafel extrapolation method.73 Polarization resistance was calculated also from parameters obtained by Tafel extrapolation (R_p, Tafel) method using Nova 2.1 software based on Eq. 1 in accordance with standard ASTM G59–97.

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R_p,Tafel = \frac{B}{j_{corr}}
\]  

where B is the Stern-Geary constant.

Electrochemical impedance spectroscopy (EIS) spectra were recorded from 100 kHz to 3 mHz frequency using an AC potential amplitude of 10 mV. Spectra were recorded at the OCP, periodically every day up to 40 days of immersion or until the samples were pitted.

**Microstructural characterization.**—Surface characterization was performed using scanning electron microscopy combined with energy dispersive X-ray spectrometry (SEM/EDS), focused ion beam microscopy (FIB) equipped with EDS, transmission electron microscopy (TEM) and X-ray photoelectron spectroscopy (XPS).

SEM images were taken in secondary electron (SE) and back-scattered electron (BSE) modes using a FEI Helios Nanolab 650 microscope. EDS spectra were taken using an Oxford Instruments Aztec system with X-max SDD (50 mm²) detector. The analysed area was 100 nm² with an analysis depth of 100 nm (near surface condition) at 3 kV beam energy, to few micrometers in diameter and the analysed depth around 1 μm for higher beam energies. EDS mapping was recorded with 3 kV beam energy and resolution of 1024 lines. Prior to analysis, samples were coated with a thin carbon layer to reduce the charging effect.

FIB was used to analyse the cross-section of conversion coatings. A FEI Helios Nanolab 650, equipped with EDS, was first used to...
The TEM specimen was the disc center (Dimple grinder, Gatan Inc., Warrendale PA, USA). Chemical analysis was performed using a conventional 200 kV transmission electron microscope (JEM-2010F, Jeol Ltd., Tokyo, Japan). Spectra were calibrated relative to the position of C 1s peak at 284.8 eV.

ToF-SIMS ion depth profiles were made using a ToF-SIMS V spectrometer (ION TOF GmbH—Munster, Germany). The analysis chamber was operated at a pressure below ~10⁻⁸ mbar. Ions depth profiles for positive ions were recorded by interlacing a pulsed 30 keV Bi⁺ primary ion source delivering 1 pA of target current over a 100 × 100 μm² area with sputtering using a 2 keV O₂ source beam delivering 544 nA of target current over a 400 × 400 μm² area. Depth profiles for negative ions were recorded by interlacing a pulsed 30 keV Bi⁺ primary ion source delivering 1.2 pA of target current over a 100 × 100 μm² area with sputtering using a 2 keV Cs⁺ source beam delivering 523 nA of target current over a 400 × 400 μm² area.

### Results and Discussion

**OCP measurements during the formation of conversion coating**—OCP during coating formation was measured in order to follow the kinetics of conversion coating formation and to determine when the formation of the coating was completed. OCP curves measured during immersion in the H₂ZrF₆ conversion bath are presented in Fig. 1. The formation of the coating from the conversion bath can be described with the following steps.¹,³²,⁴⁴

Activation of the surface caused by the initial attack of aggressive ZrF₆⁻ ions leads to the dissolution of naturally formed aluminum oxide layer Al₂O₃ (Eq. 2) that corresponds to decay of OCP (Fig. 1, region I). At the same time, oxygen reduction and hydrogen evolution reactions take place at cathodic sites and lead to local alkalization up to pH = 8.5 near cathodic sites (Eqs. 3 and 4).⁴⁵ This establishes the conditions for the initiation of coating formation (Eq. 5). At the OCP minimum the processes of natural oxide dissolution and conversion coating formation are in equilibrium. Region II, denoted by the progressive shift in OCP to more positive values, marks an increased growth of conversion coating over film dissolution. In this region, the formation and growth of conversion coating in the form of zirconium oxide/zirconium oxyfluoride takes place caused by the increased value of pH.

\[
\begin{align*}
\text{Al}^{3+} & (aq) + \text{ZrF}_6^- & \rightarrow \text{AlF}_6^{3-} (aq) + \text{Zr}^{4+} (aq) \quad [2] \\
\text{O}_2(g) + 2 \text{H}_2\text{O} + 4e^- & \rightarrow 4\text{OH}^- (aq) \quad [3] \\
2\text{H}^+ (aq) + 2e^- & \rightarrow \text{H}_2(g) \quad [4] \\
\text{ZrF}_6^- (aq) + 4\text{OH}^- (aq) & \rightarrow \text{ZrO}_2 \cdot 2\text{H}_2\text{O(s)} + 6\text{F}^- (aq) \quad [5]
\end{align*}
\]

At the end of region II, the OCP again makes an inflection point indicating that a steady state deposition process is reached. This point denotes the beginning of region III. At longer conversion times the conversion coating still continues to grow but due to thickness increase the cracking of the coating occurs and it loses its protective ability.⁵⁴,⁷⁵⁻⁷⁷ Based on the OCP measurements during ZrCC formation we conclude that the coating formation was completed after 10 min. This conversion time was then taken for the preparation of ZrCCs for all further experiments.

**Open circuit potential during immersion of conversion coatings in 0.5 M NaCl**—Within the first 1000 s to 1200 s after immersion of conversion coated AA3005 sample in 0.5 M NaCl, the OCP decreases rapidly (Fig. 2). This indicates the initial dissolution of the conversion coating in chloride containing solution and XPS was carried out using a PHI-TFA XPS spectrometer (Physical Electronic Inc.). The vacuum during the XPS analysis was in the range of 10⁻⁹ mbar. The analysed area was 0.4 mm in diameter and the depth about 3–5 nm. X-rays were provided from a monochromatic Al Kα source at a photon energy of 1486.6 eV. XPS spectra were analysed by Multipack software, version 8.0 (Physical Electronics Inc.). Spectra were calibrated relative to the position of C 1s peak at 284.8 eV.

Deposit a 1 μm thick Pt protection layer. Then, a relatively large cross-section was made by a rough removal of material with Ga ions (30 keV/48 nA) followed by a rough polishing (30 keV/22 nA) and a fine polishing (30 keV/9 nA). EDS maps of the cross-section were recorded with a resolution of 1024 lines. The average thickness of oxide layers was evaluated using ImageJ programme.

TEM was used to analyse the cross section of the conversion coatings and assess its thickness, composition and structure. Detailed structural investigations of the sample were performed using a conventional 200 kV transmission electron microscope (JEM-2010F, Jeol Ltd., Tokyo, Japan). Chemical analysis was performed using a Si(Li) EDS detector.

For TEM cross-section analysis, the sample was mounted and fitted into 3 mm brass cylinders using epoxy glue between two silicon blocks on each side to improve strength. The TEM specimen was ground to a thickness of 100 μm and dimpled down to 15 μm at the disc center (Dimple grinder, Gatan Inc., Warrendale PA, USA). The TEM specimen was finally ion-milled (PIPS, Precision Ion Polishing System, Gatan Inc., USA) using 3 kV Ar⁺ ions at an incidence angle of 8° until perforation.

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**Figure 1.** Open circuit potential measurement during the formation of zirconium conversion coating on AA3005 in 1·10⁻³ M H₂ZrF₆ conversion bath. Average value of 3 repetitions on different samples is presented by solid curve and standard deviation as grey area. Numerals I, II and III denote the regions of surface activation, initial coating formation and growth, respectively.

**Figure 2.** Open circuit potential during the first 2 h after immersion of zirconium conversion coated AA3005 samples in 0.5 M NaCl. Average value of 4 repetitions on different samples is presented by solid curve and standard deviation as grey area.
Potentiodynamic polarization curves were recorded for as-received conversion coatings. This is a OCP started to recover and during the next 6000 s shifted towards substrate and intermetallic particles (IMPs). After that period, the penetration of electrolyte through the conversion coating to the substrate. Electrochemical data deduced from polarization curves are given in Table I.

As-received (bare), chemically pre-treated and zirconium conversion coated AA3005. As-received and chemically pre-treated samples were stabilized for different periods of time (from 5 min up to 10 days). Electrochemical data deduced from polarization curves are given in Table I.

Table I. Electrochemical data deduced from potentiodynamic polarization measurements for as-received, chemically pre-treated and zirconium conversion coated AA3005 exposed to 0.5 M NaCl for different periods of time prior to electrochemical measurement. In table, average values and standard deviation (SD) are given obtained on different samples.

| Sample          | $J_{\text{corr}}$ A cm$^{-2}$ | OCP V | $E_{\text{corr}}$ V | Epit V | $\Delta E$ V | $j_0$ mV/dec | $\beta_{\text{corr}}$ mV/dec | $R_{\text{P,LPR}}$ Ω cm$^2$ | $R_{\text{P,Tafel}}$ Ω cm$^2$ |
|-----------------|-------------------------------|-------|---------------------|--------|--------------|---------------|-----------------------------|-----------------------------|-----------------------------|
| As-received-1h  | 3.9E-08                       | −0.83 | −0.82               | −0.67  | 0.15         | 220           | 116                         | 8.6E+05                     | 9.3E+05                     |
| Pre-treated-1h  | 1.1E-08                       | 0.02  | 0.03                | 0.03   | 0.04         | 23            | 19                          | 1.5E+05                     | 3.2E+05                     |
| SD              | 7.5E-08                       | −0.74 | −0.78               | −0.68  | 0.10         | 176           | 102                         | 4.6E+05                     | 3.8E+05                     |
| SD              | 1.2E-08                       | 0.01  | 0.01                | 0.02   | 0.02         | 44            | 4                           | 2.0E+05                     | 4.4E+04                     |
| ZrCC-5 min      | 2.4E-07                       | −0.87 | −0.83               | −0.61  | 0.22         | 367           | 320                         | 1.8E+05                     | 3.0E+05                     |
| SD              | 4.5E-08                       | 0.02  | 0.03                | 0.02   | 0.02         | 72            | 43                          | 1.3E+04                     | 3.0E+04                     |
| ZrCC-1h         | 2.3E-07                       | −0.93 | −0.94               | −0.63  | 0.31         | 417           | 177                         | 2.6E+05                     | 2.4E+05                     |
| SD              | 4.4E-08                       | 0.01  | 0.03                | 0.02   | 0.02         | 56            | 30                          | 3.3E+04                     | 3.7E+04                     |
| ZrCC-2h         | 1.6E-07                       | −0.89 | −0.90               | −0.59  | 0.34         | 285           | 194                         | 3.6E+05                     | 3.1E+05                     |
| SD              | 1.1E-08                       | 0.02  | 0.02                | 0.01   | 0.04         | 28            | 9                           | 6.7E+04                     | 3.5E+04                     |
| ZrCC-1d         | 1.4E-08                       | −0.71 | −0.70               | −0.40  | 0.29         | 150           | 188                         | 2.8E+06                     | 3.2E+06                     |
| SD              | 8.1E-09                       | 0.02  | 0.02                | 0.12   | 0.11         | 50            | 64                          | 1.2E+06                     | 1.2E+06                     |
| ZrCC-2d         | 8.0E-09                       | −0.69 | −0.69               | −0.36  | 0.33         | 176           | 188                         | 3.0E+06                     | 5.4E+06                     |
| SD              | 3.5E-09                       | 0.03  | 0.03                | 0.11   | 0.11         | 26            | 44                          | 1.5E+06                     | 1.4E+06                     |
| ZrCC-3d         | 6.9E-09                       | −0.69 | −0.68               | −0.25  | 0.43         | 209           | 189                         | 5.8E+06                     | 6.7E+06                     |
| SD              | 2.6E-09                       | 0.03  | 0.03                | 0.09   | 0.09         | 46            | 29                          | 5.6E+06                     | 1.5E+06                     |
| ZrCC-4d         | 5.5E-09                       | −0.67 | −0.67               | −0.32  | 0.35         | 203           | 197                         | 4.1E+06                     | 9.1E+06                     |
| SD              | 2.1E-09                       | 0.03  | 0.04                | 0.10   | 0.12         | 23            | 27                          | 2.8E+06                     | 3.5E+06                     |
| ZrCC-5d         | 5.1E-09                       | −0.66 | −0.67               | −0.17  | 0.49         | 220           | 184                         | 4.0E+06                     | 9.6E+06                     |
| SD              | 2.0E-09                       | 0.03  | 0.04                | 0.20   | 0.20         | 55            | 41                          | 3.4E+06                     | 3.2E+06                     |
| ZrCC-10d        | 3.2E-09                       | −0.65 | −0.67               | −0.22  | 0.45         | 235           | 173                         | 5.3E+06                     | 1.6E+07                     |
| SD              | 1.8E-09                       | 0.03  | 0.04                | 0.13   | 0.09         | 29            | 54                          | 5.0E+06                     | 5.1E+06                     |

$\Delta E = |E_{\text{pit}} - E_{\text{corr}}|$. With the prolongation of immersion time above 1 h, the $j_{\text{corr}}$ of ZrCC samples progressively decreased, $E_{\text{corr}}$ shifted more positive and $\Delta E$ increased. After 24 h of immersion, the $j_{\text{corr}}$ was smaller by 1 order of magnitude, while OCP was shifted towards more positive values reaching ~0.71 V. This was a significant improvement in the performance of ZrCC compared to the short immersion.
of 1 h. The trend of decreasing corrosion current densities continued up to 10 days of immersion in 0.5 M NaCl solution reaching OCP of $-0.65$ V, $j_{corr}$ of $3 \cdot 10^{-9}$ A cm$^{-2}$, and $\Delta E$ of 0.45 V. This electrochemical behavior indicates that ZrCC on AA3005 acts as an anodic inhibitor and that some kind of self-sealing/active corrosion protection takes place during prolonged immersion in 0.5 M NaCl.

**Electrochemical impedance spectroscopy.**—To evaluate the performance of conversion coatings the electrochemical impedance spectra (EIS) were measured at OCP for as-received, chemically pre-treated and ZrCC coated AA3005 samples until pitting occurred. Curves of OCP and impedance modulus at 3 mHz ($|Z|_{3\text{mHz}}$) measured during the course of immersion are presented in Fig. 4. Bode and Nyquist diagrams for all repetitions are presented in Supplement material (Figs. S4–S6). Impedance at low frequencies represents total polarization resistance of the coating and it is proportional to the corrosion resistance of a coating.81–83 For ZrCC samples the impedance modulus was $3 \cdot 10^6$ Ω cm$^{-2}$ after 1 h of immersion in 0.5 M NaCl; however, it increased over one order of magnitude during the first 6 to 7 days of immersion reaching the average value of $7.5 \cdot 10^6$ Ω cm$^{-2}$. After 10 days of immersion, $|Z|_{3\text{mHz}}$ impedance modulus formed a plateau at the average value $8.5 \cdot 10^6$ Ω cm$^{-2}$ (Fig. 4b). The same trend was noticed with Bode phase angles diagrams (Figs. S4–S6) where phase angle at low frequencies (ϕ) continuously increased up to 20 days of immersion, or even longer, depending on the sample. The highest increase was noted during the first 10 days of immersion indicating self-sealing/active corrosion protection within ZrCC. Drops in $|Z|_{3\text{mHz}}$ and ϕ occurred after 20 days of immersion (Fig. 4b) but afterwards $|Z|_{3\text{mHz}}$ recovered to a higher average value of $6.5 \cdot 10^6$ Ω cm$^{-2}$; this indicates increased probability that pitting will occur after 20 days of immersion in 0.5 M NaCl solution. Finally, after 30 days of immersion, a decrease in $|Z|_{3\text{mHz}}$ and ϕ indicates that samples underwent pitting corrosion.

According to Buchheit et al., if conversion coating applied on AA3003 exhibits $R_c$ (defined as apparent integrated pitting resistance) of $2.3 \cdot 10^6$ Ω cm$^{-2}$ (range $1.1 \cdot 10^7$ to $4.8 \cdot 10^7$ Ω cm$^{-2}$) after 168 h immersion in 0.5 M NaCl, it has a probability of $>80\%$ to pass 168 h neutral salt spray test (NSST) test.84,85 It was pointed out that EIS is more sensitive than salt spray chamber exposure for detecting the corrosion of a coating when the inspection interval was $>24$ h.86 The ZrCCs on AA3005 showed exceptional performance with impedance modulus of $8.5 \cdot 10^6$ Ω cm$^{-2}$ after 168 h of immersion. This value is smaller than that proposed by Buchheit et al.85 but it has to be taken into account that the low frequency limit was higher than in the present work (10 mHz and 3 mHz, respectively). When taking that into account, the fitted average values of EIS parameters obtained in this work are $R_c$ (96 h) = $1.2 \cdot 10^7$ Ω cm$^{-2}$ and $R_c$ (168 h) = $1.61 \cdot 10^7$ Ω cm$^{-2}$ after 96 h and 168 h of immersion in 0.5 M NaCl. Based on these values we therefore assume that the coatings will pass 168 h NSST mark.84,85 More data are given in supplement material (Table SVII).

The time dependence of EIS spectra and progressive improvement of corrosion resistance is in agreement with results of PD measurements indicating that ZrCC coating on AA3005 has excellent corrosion performance, even better than TCP conversion coatings.71,72 During the course of immersion in NaCl, the time dependence of OCP indicate that ZrCCs act as anodic inhibitors: OCP shifted from approximately $-0.9$ V after 1 h of immersion, to $-0.7$ V after 24 h of immersion, reaching values $-0.6$ V to $-0.65$ V after 10 days of immersion (Fig. 4a). Trends in OCP and impedance modulus were in correlation since the drop in $|Z|_{3\text{mHz}}$ was followed by a drop in OCP, as characteristic for pitting and exposure of AA3005 substrate to the electrolyte.

**Figure 4.** Impedance module at 3 mHz and open circuit potential as a function of immersion time for: (a, b) zirconium conversion coated AA3005; (c, d) as-received AA3005 (black symbols) and chemically pre-treated AA3005 (red symbols). Average values of 3 repetitions on different samples are presented by solid curves and standard deviation as error bars. Lines were drawn to guide the eye.
Compared to ZrCCs coated samples, the EIS behavior was quite different for bare and chemically pre-treated samples. For the latter, the $|Z|$ showed an initial increase with immersion time but the values are approximately 3–4 times smaller compared to ZrCCs coated samples (Fig. 4d) and reached the maximum of $2 \cdot 10^6 \Omega \cdot \text{cm}^2$. However, this value is still comparatively high and indicates that even bare AA3005 has substantial corrosion resistance. At longer immersion times, $|Z|$ decreased, reflecting the initiation of the localized corrosion process. The time dependencies of OCP and Bode plots show a similar trend (Fig. 4c, S5). The same trend was noted with Bode phase angle plots (Fig. S5). Chemically pre-treated AA3005 showed very similar behavior as bare AA3005 but it was less corrosion resistant which is in good agreement with PD curves (Fig. 3, Table I).

**SEM/EDS characterization of the coating surface.**—SEM images of surface morphology and microstructure of chemically pre-treated AA3005, as-prepared ZrCC coated AA3005, and after immersion in NaCl are given in Fig. 5 recorded at lower and higher magnification. As-prepared ZrCC (Fig. 5b) was obviously very thin as no significant change was observed at the majority of the surface using SE at lower magnification mode compared to a chemically pre-treated sample (Fig. 5a, 5b). At higher magnification the coating is visible, especially around IMPs (inset in Fig. 5b). After 2 days of immersion in 0.5 M NaCl, the most significant feature was the appearance of a bloated area around IMPs (denoted as a "hat") (Fig. 5c). After 5 days of immersion, this area was not bloated anymore. There were no visible corrosion or pits in conversion coatings after 5 days of immersion in 0.5 M NaCl.

As-received AA3005 was composed of: Al (96.7 at. %), O (1.8 at. %), Mn (0.6 at. %), Fe (0.3 at. %), Mg (0.5 at. %) and Si (0.1 at. %) (Table II). The microstructural characteristics of AA3005 were typical as reported for this alloy.86,87 The main intermetallic particles were Al$_3$(Mn, Fe) composed of Mn and Fe in an atomic ratio approximately 1, 1.3 and 1.6, respectively, for each spot, (Fig. 6a, spot 3, Table II and Fig. 6b, spots 4, 5, Table III). α-Al(Fe, Mn)Si IMPs containing Mn, Fe and Si were also detected (Fig. 6a, spot 4, Table II and Fig. 6b, spots 3, 6, Table III). The atomic ratios Mn/Fe and Mn/Si were similar, i.e. between approximately 1 and 2 (Table III). A large number of dispersoids (little white dots) sized below 1 μm containing mainly of Mn were present (Fig. 6a spot 2, Table II). There were no visible changes in the microstructure and elemental composition of AA3005 after alkaline cleaning using Metaclean (Fig. 6b, Table III). This confirms that Metaclean pre-treatment removed just grease and organic contamination from the surface.

SEM-BSE images of the as-prepared ZrCC and after immersion in 0.5 M NaCl for 2 and 5 days are presented in Figs. 7b–7d (SE images are presented in Fig. 5b–5d). For the sake of comparison, Fig. 6b for chemically pre-treated AA3005 is repeated as Fig. 7a. These samples were analysed by point EDS using an electron beam voltage of 3 kV or 11 kV corresponding to the analysed depth of approximately 100 nm and 1 μm, respectively (Tables IV–VII and SII–SIV). Results of EDS point analysis are presented in the form of atomic ratios (Table IV) or atomic % (Tables V–VII and SII–SIV). Further, EDS mapping was recorded for all the samples at 3 kV electron beam voltage (Fig. 8).

After conversion for 10 min in the conversion bath, the formation of ZrCC was completed over the whole surface (Fig. 7b and Tables IV, VII, SIII). The coating was cracked and much thicker at the (Mn, Fe) IMPs and in their proximity. These areas are denoted as hats. The atomic ratios of O/Zr, F/Zr and Al/Zr in the matrix of the as-prepared ZrCC were 2.39, 0.61 and 4.86 respectively (Fig. 7b, spot 2, Table VII), while the atomic ratios of O/Zr and F/Zr in coating at the hat above (Mn, Fe) IMPs were 1.68 and 0.31, respectively (Fig. 7b, spot 4, Table VII). The latter values indicate that the hat consists of ZrO$_{1.68}$F$_{0.31}$ and does not contain Al. There were darker areas in the coating with a smaller concentration of Zr, F and O, and a higher concentration of Al indicating that coating was thinner at these areas (Fig. 7b, spot 5, Table IV). EDS mapping
clearly shows that ZrCC was preferentially formed at and around IMPs (Fig. 8).

After 2 days of immersion in a 0.5 M NaCl solution, the most visible change at the coating was a dark circle around (Mn, Fe) IMPs (Fig. 7c and Tables V, VII, SIII). Apparently, the cracks around (Mn, Fe) IMPs were weak spots in the coating where electrolyte penetrated and the corrosion process initiated. This would usually lead to the dissolution of IMPs, degradation of conversion coating and pit growth but in this case, a densification and closing of cracks/pit around (Mn, Fe) IMPs occurred. This would usually lead to the dissolution of IMPs, degradation of conversion coating and pit growth, but in this case, a densification and closing of cracks/pit around (Mn, Fe) IMPs occurred. This is clearly visible from EDS analysis/mapping which shows a very high concentration of O (65.8 at.%) and Al (30.7 at.%) at dark circle area around (Mn, Fe) IMPs indicating the formation of mostly oxide-containing area (Fig. 7c, spot 5, Table V). The atomic ratio of O/Zr at the hat increased to 2.69, while Al and F were not detected (Fig. 7c, spot 4, Table VII); therefore, the composition changed from \( \text{ZrO}_{1.68}\text{F}_{0.31} \) to \( \text{ZrO}_2 \cdot 0.7 \text{H}_2\text{O} \). This indicates that the process of densification and closing of pores occurs in the whole coating and not just around (Mn, Fe) IMPs. The most significant change in ZrCC occurred during immersion in NaCl was a 5-fold reduced concentration of fluorine, i.e., from 6.9 at.% down to 1.3 at.% (Tables IV, V, spot 2). Atomic ratios of Al/Zr, O/Zr and F/Zr changed to 5.63, 2.86, and 0.125 (Fig. 7c, spot 2, Table VII). A decrease in fluorine concentration in the ZrO2 hats and around them is clearly visible from EDS mapping (Fig. 8).

After 5 days of immersion in 0.5 M NaCl solution, the most visible change was a change of the dark circle around the Zr-oxide hat from dark to light white color with further spreading to a surrounding area (Fig. 7d and Tables VI, VII, SIV). It was difficult to select the right area for the comparison due to different samples. The area selected to be similar as in Fig. 7c seems to contain a significantly reduced O concentration (down to 31.0 at.%) and increased concentrations of Zr and Al (up to 10.2 at% and 57.7 at%, respectively) (Fig. 7d, spot 5, Table VII).

| Spot | at.% Al | at.% O | at.% Mn | at.% Fe | at.% Mg | at.% Si |
|------|--------|--------|---------|---------|---------|---------|
| 1    | 89.2   | 10.4   | —       | —       | 0.4     | —       |
| 2    | 87.5   | 12.1   | —       | —       | 0.4     | —       |
| 3    | 79.2   | 20.8   | —       | —       | —       | —       |
| 4    | 81.5   | 18.5   | —       | —       | —       | —       |
| 1*   | 96.7   | 1.8    | 0.6     | 0.3     | 0.5     | 0.1     |
| 2*   | 98     | 1.5    | —       | —       | 0.5     | —       |
| 3*   | 79.3   | 2.8    | 8.8     | 9.0     | —       | —       |
| 4*   | 83.6   | 4.1    | 5.0     | 3.2     | —       | 4.1     |

Table II. EDS analysis of as-received (bare) AA3005 (Fig. 6a); electron beam voltage of analysed spots 1–3 was 3 kV (denoted 1-4) or 11 kV (denoted 1*-4*) differing in the depth on the analysed region (approximately 100 nm and 1 μm, respectively).

| Spot | at.% Al | at.% O | at.% Mn | at.% Fe | at.% Mg | at.% Si |
|------|--------|--------|---------|---------|---------|---------|
| 1    | 88.8   | 10.6   | —       | —       | 0.6     | —       |
| 2    | 88.8   | 10.6   | —       | —       | 0.6     | —       |
| 3    | 82.3   | 17.7   | —       | —       | —       | —       |
| 4    | 83.1   | 16.9   | —       | —       | —       | —       |
| 5    | 82.4   | 17.6   | —       | —       | —       | —       |
| 6    | 87.7   | 12.3   | —       | —       | —       | —       |
| 1*   | 96.6   | 1.9    | 0.6     | 0.3     | 0.5     | 0.1     |
| 2*   | 97     | 1.8    | 0.5     | 0.1     | 0.5     | —       |
| 3*   | 78.4   | 2.1    | 7.2     | 7.8     | 0.2     | 4.3     |
| 4*   | 87.3   | 3.7    | 5.0     | 3.9     | —       | —       |
| 5*   | 88.2   | 3.7    | 4.7     | 3.0     | 0.4     | —       |
| 6*   | 87.9   | 2.0    | 5.1     | 2.5     | —       | 2.6     |
5, Table VI) compared to that after 2 days immersion. The fluorine concentration was only 1.3 at.%. Atomic ratios Al/Zr, O/Zr and F/Zr were 5.14, 3.15 and 0.142 (Fig.7d, spot 2, TableVII) indicating densification of ZrCC. In the zirconium oxide hat the atomic ratio O/Zr was 3.16 (Fig.7d, spot 4, TableVII).

Summarizing, during the course of immersion in NaCl, the Al/Zr and O/Zr ratios at the coating matrix increased and that of F/Zr decreased indicating densification with Al- and Zr-oxides and depletion with fluoride or oxyfluoride. At the hat above (Mn,Fe) IMPs, Al was not detected and F was detected only prior to immersion. A change of O/Zr ratio reflects a progressive transformation from ZrO$_{1.68}$F$_{0.31}$ to ZrO$_2$ $\cdot$ 0.7 H$_2$O and finally ZrO$_2$ $\cdot$ 1.2 H$_2$O.

Table IV. EDS analysis of as-prepared zirconium conversion coatings on AA3005 (Fig. 7b); electron beam voltage of analysed spots 1–5 was 3 kV corresponding to the depth on the analysed region of approximately 100 nm. The results of the analysis carried out at 11 kV are presented in Table SII.

| Spot | at.% Al | at.% Zr | at.% O | at.% F |
|------|--------|--------|--------|--------|
| 1    | 55.4   | 12.7   | 25.5   | 6.4    |
| 2    | 54.9   | 11.3   | 27.0   | 6.9    |
| 3    | 22.3   | 23.9   | 45.1   | 8.7    |
| 4    | —      | 33.5   | 56.2   | 10.3   |
| 5    | 74.9   | 7.6    | 13.9   | 3.6    |

Table V. EDS analysis of zirconium conversion coatings on AA3005 immersed in 0.5 M NaCl solution for 2 days (Fig. 7c); electron beam voltage of analysed spots 1–6 was 3 kV corresponding to the depth on the analysed region of approximately 100 nm. The results of the analysis carried out at 11 kV are presented in Table SIII.

| Spot | at.% Al | at.% Zr | at.% O | at.% F |
|------|--------|--------|--------|--------|
| 1    | 53.1   | 10.9   | 34.6   | 1.4    |
| 2    | 58.6   | 10.4   | 29.7   | 1.3    |
| 3    | 33.0   | 17.8   | 47.8   | 1.5    |
| 4    | —      | 27.1   | 72.9   | —      |
| 5    | 30.7   | 0.7    | 65.8   | 2.8    |
| 6    | 30.3   | 5.0    | 62.1   | 2.6    |

Figure 7. SEM-BSE images of AA3005: (a) pre-treated with Metaclean; (b) zirconium conversion coating as prepared; (c) zirconium conversion coating after 2 days of immersion in 0.5 M NaCl and (d) zirconium conversion coating after 5 days of immersion in 0.5 M NaCl. EDS mapping of areas denoted by square 1 is presented in Fig. 8. Squares 1, 2, 3, 4, 5, 6, present spots in Tables IV–VI and SII–SIV. Figure 7a is the same as 6b and is given for comparison's sake.

SEM/EDS characterization at the coating cross-section.—SEM images of cross section and related EDS mapping of as-prepared ZrCC and ZrCC after 5 days immersion in 0.5 M NaCl were taken in the region near (Mn, Fe) IMPs of the AA3005 sample (Fig. 9). In this area the coating was thicker and its structure is fully developed. The thickness of the coating before and after immersion in NaCl solution was 200 nm. Based on SEM images and EDS mapping it appears that coating had a tri-layer structure. In the as-prepared ZrCC, Zr and O were concentrated in the upper layer of the coating, while F, Mn, Mg were equally distributed across whole coating (Fig. 9a). Distribution of elements in ZrCC changed after 5 days of immersion in 0.5 M NaCl solution: an upper layer of the coating consisted mainly of Zr, Mg and Mn; the middle layer consisted of O, F and Al, while the bottom layer contained mainly Al, O, F, Mn and Mg (Fig. 9b).

Atomic ratios of elements within ZrCC better describe the structure of coatings (Table VIII). (Please note that the atomic concentrations at spots 1–4 were reconstructed from EDS maps where the analysis spot is larger than the analyzed spot. Therefore, relative error is always similar and allows for relative comparison of concentrations). Atomic ratios O/Zr, Al/Zr, Mg/Zr increased from the upper to the inner part of the ZrCC coating indicating the enrichment of the inner part with Al, O and Mn. Ratios F/Zr and Mn/Zr were lower, or F and Mn were not detected at all, in the middle layer of ZrCC. In the bottom layer of the ZrCC, atomic ratios F/Zr and Mn/Zr were higher in comparison with the top layer of coating.
The same trend was found after 5 days of immersion in 0.5 M NaCl (Table VIII). A substantial increase in O/Zr ratio up to 17.75 was noticed at the coating/substrate interface, and increased Mn/Zr and Fe/Zr ratios are the most significant changes. These indicate that Mn, Fe and O were incorporated in the bottom layer. The ratio of F/Zr was larger as well compared to the coating top. These findings are in agreement with SEM/EDX analysis of coating (Figs. 5–8) and confirm the densification and self-sealing of ZrCC during immersion in a 0.5 M NaCl solution.

SEM images and EDS mapping of the cross-sections of as-prepared ZrCC and Zr-oxide hat above (Mn, Fe) IMP after 5 days of immersion in 0.5 M NaCl are presented in Fig. 11. As shown by EDS mapping, the hat was composed mainly of Zr, O and Al with atomic ratios for O/Zr, Al/Zr of 1.41 and 0.25, respectively (Figs. 11c, 11e and Table X, spot 1). The main change in the composition of Zr-oxide hat was the complete disappearance of F. IMPs had an Mn/Fe atomic ratio of 0.75 (Fig. 11c, spot 2) and 1.42 (Fig. 11c, spot 3). In the area between two IMPs there was a dense layer of Al, O and Zr with atomic ratios Al/Zr, O/Zr of 5.33 and 5.63, respectively (Fig. 11c, spot 4). It seems that the corrosion process was initiated around (Mn, Fe) IMPs but the channels around them then filled with a dense layer of aluminum oxide and passivated after 5 days of immersion in a 0.5 M NaCl solution. A small amount of Mn, Fe, Zr and Al was present in aluminum oxide as well. These elements may be therefore incorporated in the growing layer of aluminum oxide formed around and above (Mn, Fe) IMPs (Fig. 11e). Figure 11d presents the cross-section taken in the proximity of IMP (detailed in Fig. 11b and imaged at higher magnification in Fig. 9b). The coating was uniform and undamaged in the proximity of (Mn, Fe) IMPs even after 5 days of immersion in 0.5 M NaCl. It had a tri-layer structure with a thickness of 200 nm.

**TEM characterization of the coating cross section.—** Transmission electron images of cross-sectional ZrCC taken at the coating matrix are presented in Fig. 12. As-prepared ZrCC exhibited a tri-layer structure with a total thickness of 29 nm. The top layer had a thickness of 21.5 nm, middle layer 5 nm and the inner layer 2.5 nm. The coating was homogenous and without cracks. It is noteworthy that there is a difference in thickness obtained using TEM (Figs. 9–11) and TEM (Fig. 12) analyses, the latter being much smaller. This difference can be ascribed to the uneven thickness of the coating over the AA3005 surface, i.e., the coating is much thicker at and around the IMPs (Figs. 7, 9, 10, 11) than at the matrix (Fig. 12). The ratio between thickness near (Mn, Fe) IMPs and that of coating matrix far from IMPs (dZrCC-IMPs/dZrCC) was 6.9 as determined based on SEM and TEM analyses, respectively. Based on the thickness of individual layers within the coating determined by TEM analysis (Fig. 12), the contribution of individual layers before and after immersion in NaCl was calculated (Table XI). Based on such normalized layer thickness it can be stated that the contribution of the top layer decreased from 74.2% to 49.2%; the middle layer increased from 17.2 to 21.0%, and that of bottom layer increased from 8.6% to 29.8%. These findings are in good correlation with SEM/EDS data that confirmed an increased concentration of O, Mn and Fe in bottom layers of ZrCC after 5 days of immersion in NaCl (Fig. 9).

SEMs and EDS mapping of the cross-sections of ZrCC and Zr-oxide hat above (Mn, Fe) IMP after 5 days of immersion in 0.5 M NaCl are presented in Fig. 11. As shown by EDS mapping, the hat was composed mainly of Zr, O and Al with atomic ratios for O/Zr, Al/Zr of 1.41 and 0.25, respectively (Figs. 11c, 11e and Table X, spot 1). The main change in the composition of Zr-oxide hat was the complete disappearance of F. IMPs had an Mn/Fe atomic ratio of 0.75 (Fig. 11c, spot 2) and 1.42 (Fig. 11c, spot 3). In the area between two IMPs there was a dense layer of Al, O and Zr with atomic ratios Al/Zr, O/Zr of 5.33 and 5.63, respectively (Fig. 11c, spot 4). It seems that the corrosion process was initiated around (Mn, Fe) IMPs but the channels around them then filled with a dense layer of aluminum oxide and passivated after 5 days of immersion in a 0.5 M NaCl solution. A small amount of Mn, Fe, Zr and Al was present in aluminum oxide as well. These elements may be therefore incorporated in the growing layer of aluminum oxide formed around and above (Mn, Fe) IMPs (Fig. 11e). Figure 11d presents the cross-section taken in the proximity of IMP (detailed in Fig. 11b and imaged at higher magnification in Fig. 9b). The coating was uniform and undamaged in the proximity of (Mn, Fe) IMPs even after 5 days of immersion in 0.5 M NaCl. It had a tri-layer structure with a thickness of 200 nm.

**Table VI.** EDS analysis of zirconium conversion coatings on AA3005 immersed in 0.5 M NaCl solution for 5 days (Fig. 7d); electron beam voltage of analysed spots 1–6 was 3 kV corresponding to the depth on the analysed region of approximately 100 nm. The results of the analysis carried out at 11 kV are presented in Table SIV.

| Spot | at.% Al | at.% Zr | at.% O | at.% F |
|------|---------|---------|-------|-------|
| 1    | 53.9    | 11.0    | 33.7  | 1.4   |
| 2    | 54.5    | 10.6    | 33.4  | 1.5   |
| 3    | 9.4     | 23.5    | 67.2  | —     |
| 4    | —       | 24.0    | 76.0  | —     |
| 5    | 57.7    | 10.2    | 31.0  | 1.1   |
| 6    | 35.2    | 18.7    | 45.0  | 1.1   |

**Table VII.** Atomic ratios in the matrix area of zirconium conversion coatings and at the hat for as-prepared ZrCC and ZrCC immersed in 0.5 M NaCl solution for 2 and 5 days. Atomic ratios were calculated from spots 2 and spots 4 (Figs. 7b–7d and Tables IV–VI) for matrix area and hat respectively.

| Sample | Al/Zr | O/Zr | F/Zr |
|--------|-------|------|------|
| ZrCC-0d, spot 2 | 4.86 | 2.39 | 0.61 |
| ZrCC-2d, spot 2 | 5.63 | 2.86 | 0.125 |
| ZrCC-5d, spot 2 | 5.14 | 3.15 | 0.142 |
| ZrCC- hat–0d, spot 4 | — | 1.68 | 0.31 |
| ZrCC- hat–2d, spot 4 | — | 2.69 | — |
| ZrCC- hat–5d, spot 4 | — | 3.16 | — |
Figure 9. SEM images and EDS mapping of a cross-section of zirconium conversion coatings on AA3005: (a) zirconium conversion coating as-prepared; (b) zirconium conversion coating after 5 days of immersion in 0.5 M NaCl. The area where the mapping was recorded is denoted by vertical rectangles, circles and horizontal rectangles denote the spots where EDS analysis was carried out (Table VIII).

Table VIII. EDS analysis given in atomic ratios of elements at the cross-section of as-prepared zirconium conversion coating on AA3005 (Fig. 9a) and after 5 days immersion in 0.5 M NaCl (Fig. 9b).

| Spot       | Al/Zr | O/Zr | F/Zr | Mn/Zr | Mg/Zr | Fe/Zr |
|------------|-------|------|------|-------|-------|-------|
| as-prepared|       |      |      |       |       |       |
| 1          | 7.37  | 1.53 | 0.233| 0.023 | 0.035 | —     |
| 2          | 15.60 | 1.49 | 0.189| —     |       | 0.076 |
| 3          | 34.92 | 2.15 | —    | —     | 0.192 | —     |
| 4          | 39.17 | 2.43 | 0.435| 0.087 | 0.217 | —     |
| Mapping average | 14.30 | 1.87 | 0.226| 0.038 | 0.076 | —     |

| mapping average | 18.12 | 3   | 0.220| 0.123 | 0.098 | 0.073 |

After 5 days immersion

| Spot | Al/Zr | O/Zr | F/Zr | Mn/Zr | Mg/Zr | Fe/Zr |
|------|-------|------|------|-------|-------|-------|
| 1    | 6.91  | 1.88 | 0.174| 0.070 | 0.070 | 0.023 |
| 2    | 6.74  | 1.84 | 0.202| —     | 0.061 | —     |
| 3    | 15.16 | 3.04 | 0.255| 0.020 | 0.020 | 0.020 |
| 4    | 226.5 | 17.75| 0.5  | 1.25  | 1     | 1     |
| Mapping average | 18.12 | 3   | 0.220| 0.123 | 0.098 | 0.073 |

Figure 10. SEM images with EDS mapping of cross section of as prepared ZrCC. (a) top view, (b,c) side view, (d) the cross-section of coating in the proximity of (Mn, Fe) IMPs, as indicated by an arrow in (b), and (e) EDS mapping of the cross-section. The results of EDS analysis are given in Table IX and S5.
XPS characterization of the coating surface.—XPS analysis was carried out on as-received and chemically pre-treated AA3005, as prepared ZrCC and ZrCC immersed in 0.5 M NaCl solution for 2 and 5 days. XPS technique has a probing depth up to 10 nm thus giving information from the upper-most layer of the ZrCC. The composition of samples deduced from XPS survey spectra are presented in Table XII. The surface of an as-received sample contained Al, O, Mg, and N, while chemically pre-treated sample contained Al, O, Mg, P, Si and Na. Na and P originate from Metaclean alkaline cleaner that contains NaSiO3 and probably some P based corrosion inhibition/surfactant. Chemical cleaning does not cause an enrichment in Mn (detected concentration < 0.2 at.%).

Aluminum and oxygen are related to the formation of aluminum oxide/hydroxide. After the conversion coating process with hexafluorozirconic acid, aluminum almost diminished due to coverage with ZrCC, as evidenced by the appearance of Zr and F originating from the conversion coating. After 2 days of immersion in NaCl, however, F completely disappeared from the top layer of ZrCC and, at the same time, Al appeared in the top layer. The concentration of Al increased with immersion time, while that of Zr decreased. Si appeared after 2 days of immersion in a 0.5 M NaCl solution and originates from AA3005.

High resolution XPS spectra recorded for major elements detected in survey XPS spectra are presented in Fig. 13. High resolution Al 2p spectra are presented in Fig. 13a. The Al 2p peak is usually presented as a single peak rather than two split component peaks 2p3/2 and 2p1/2. Two 2p peaks are noted in the Al 2p spectrum for as-received and chemically pre-treated Al centered at binding energy (E_b) 72.4 and 74.8 and corresponding to Al metal and Al(OH)3, respectively.88,89 Considering the intensity ratio between the two peaks, it can be stated that the surface is covered by a thin, few nanometers thick, layer of Al(OH)3 which allows the detection of Al metal. After the conversion coating process, however, peaks for Al diminished due to coverage of AA3005 with ZrCC. After 2 and 5 days of immersion, a small peak at 74.5 eV re-appeared suggesting the formation of Al(OH)3 in the top layer of coating.

High resolution O 1 s spectra are presented in Fig. 13b. For as-received and chemically pre-treated samples, the peak was centered at 531.9 eV suggesting the prevalence of hydroxide over oxide.
Considering the presence of hydrated Zr oxide (see below), this peak could be deconvoluted into peaks for bound H2O, Al(OH)3 and ZrO2·2H2O at binding energies at 533 eV, 532 (531.9) eV and 531.5 eV, respectively. After the conversion process, the O 1 s peak shifted to 531.2 eV and is now related to predominantly hydrated Zr oxide and/or oxyfluoride (ZrO2·2H2O/ ZrFxOy). After 2 and 5 days of immersion ZrCC in a 0.5 M NaCl solution, the center of the O1 s peak was shifted back to slightly higher E_b of 531.6 eV corresponding to the presence of ZrO2·2H2O and Al(OH)3.

The F 1s peak was located at E_b of 684.8 eV and was noted only for the as-prepared ZrCC indicating the presence of ZrF4/ZrFxOy in the top layer of as-prepared coating (Fig. 13d). After 2 and 5 days of immersion, the F 1s peak disappeared, indicating dissolution of fluorine and transformation of ZrF4/ZrFxOy into ZrO2·2H2O.

High resolution spectra Zr 3d spectra recorded after the conversion coating process are presented in Fig. 13c. The Zr 3d peaks for the as-prepared coating are centered at 185.5 eV and 183.2 eV, respectively. Considering a relatively high fluorine and oxygen contents (Fig. S7), this spectrum reflects the presence of a mixture of ZrF4/ZrFxOy and ZrO2·2H2O. After 2 and 5 days of immersion in 0.5 M NaCl solution, however, the peaks shifted to lower E_b of 184.9 eV and 182.5 eV, respectively, indicating the progressive transformation of Zr-fluoride/oxyfluoride to ZrO2·2H2O. These findings are in correlation with the disappearance of the fluorine peak in the F 1s spectrum.

Table XI. Percentage of individual layer thickness to total coating thickness for as prepared zirconium conversion coating and zirconium conversion coating immersed in 0.5 M NaCl solution for 5 days (calculated from layer thickness in Fig. 12).

| Layer        | ZrCC-0d | ZrCC-5d |
|--------------|---------|---------|
| Top layer, % | 74.2    | 49.2    |
| Middle layer, % | 17.2    | 21.0    |
| Bottom layer, % | 8.6     | 29.8    |

Considering the presence of hydrated Zr oxide (see below), this peak could be deconvoluted into peaks for bound H2O, Al(OH)3 and ZrO2·2H2O at binding energies at 533 eV, 532 (531.9) eV and 531.5 eV, respectively. After the conversion process, the O 1 s peak shifted to 531.2 eV and is now related to predominantly hydrated Zr oxide and/or oxyfluoride (ZrO2·2H2O/ ZrFxOy). After 2 and 5 days of immersion ZrCC in a 0.5 M NaCl solution, the center of the O1 s peak was shifted back to slightly higher E_b of 531.6 eV corresponding to the presence of ZrO2·2H2O and Al(OH)3.

Table XII. Concentration of elements obtained by XPS analysis of as-received, chemically pre-treated and zirconium conversion coated AA3005 samples.

| Sample      | at.% Al | at.% Zr | at.% O | at.% F | at.% Mg | at.% Si | at.% P | at.% Na | at.% N | at.% Cl |
|-------------|---------|---------|--------|--------|---------|--------|-------|--------|-------|--------|
| as received | 19.6    | —       | 69.1   | —      | 9.9     | —      | —     | —      | 1.4   | —      |
| pre-treated | 18.9    | —       | 66.6   | —      | 5.9     | 0.5    | 3.7   | 4.3    | —     | —      |
| ZrCC-0d     | 1.0     | 23.4    | 67.2   | 8.4    | —       | —      | 3.0   | —      | —     | —      |
| ZrCC-2d     | 4.9     | 17.4    | 74.7   | —      | —       | —      | —     | —      | 8.1   | —      |
| ZrCC-5d     | 8.1     | 14.6    | 69.2   | —      | —       | —      | —     | —      | —     | —      |

Figure 12. Transmission electron microscopy images of (a, c) as-prepared zirconium conversion coating and (b, d) after 5 days immersion in 0.5 M NaCl solution.
rate was 0.30 nm s$^{-1}$ ($\pm$ 0.02 nm s$^{-1}$). As-prepared ZrCC was approximately 30 to 40 nm thick. It consisted of an outer layer that had the highest concentration of ZrO$^-$ and a lower concentration of AlOF$^-$ (Fig. 14, region 1); therefore, its structure can be correlated with ZrO$_2$ \cdot 2H$_2$O/ZrO$_x$F$_y$ species. A significant amount of CuO$^-$ ions was detected in the outer layer. The highest amount of AlOF$^-$ ions was detected in the middle layer, accompanied with peaks for AlO-, MgO- and Al$_2$O$_4^-$ ions (Fig. 14, region 2). ZrO$^-$ ions were present in the middle layer but in a smaller amount than in the outer layer. The same ions were found in the middle layer and at the interface (Fig. 13, region 3) so it was hard to distinguish the border between these two regions.

After 5 days of immersion in 0.5 M NaCl, the profile of ZrO$^-$ ions remained approximately the same with a maximum in the outer layer (Fig. 14, region 1). The most important change was a decrease in intensity (concentration) of F$^-$ and AlOF$^-$ ions. At the same time, the concentration of Cl$^-$ ions increased significantly following the same profile shape as that of F$^-$ and AlOF$^-$ ions. This indicates that Cl$^-$ ions were exchanged with F$^-$ ions and incorporated within the ZrCC structure. The concentration of AlO$^-$ was larger throughout the ZrCC showing two maxima in the outer and, especially, the middle part, respectively. This increased concentration extended over the interface with the substrate for approximately 10–15 nm.

Mechanism of active corrosion protection/self-sealing. — Electrochemical results obtained for zirconium conversion coatings on AA3005 suggest that these coatings enable self-sealing and active corrosion protection when immersed in a 0.5 M NaCl solution. These following results substantiate this statement: (i) rapid decay and recovery of OCP during the first 2 h immersion, (ii) a 50-fold increase in impedance modulus at 3 mHz and the ability to sustain it at values in the megaohm-cm$^2$ range (7.5 \cdot 10^6 to 9.5 \cdot 10^6 Ω-cm$^2$) and to resist pitting up to at least 21 days, and (iii) increased passive window (from 0.22 to 0.45 V) and decreased corrosion current density of two orders of magnitude (from 2.32 \cdot 10^{-7} A-cm$^{-2}$ down to 3 \cdot 10^{-9} A-cm$^{-2}$) after 10 days of immersion.

The electrochemical study was complemented with microstructural and compositional studies. The microstructural investigation by XPS showed the disappearance of fluorine in the top layer of ZrCC after 2 days in 0.5 M NaCl and the transformation of ZrF$_4$/ZrF$_x$O$_y$ into Zr(OH)$_2$ \cdot 1.2H$_2$O and Al(OH)$_3$. SEM/EDS investigation confirmed that the concentration of F decreased and that of Al and O increased during immersion. FIB/EDS investigation showed that coating was 200 nm thick in the proximity of IMPs, homogenous with tri-layer structure, while (Mn, Fe) IMPs were covered with a dense hat of zirconium oxide. During immersion in NaCl, channels around (Mn, Fe) IMPs became progressively filled with a dense layer of aluminum oxide containing a small content of Zr and F. This process results in the passivation of the IMPs. At the same time, (Mn, Fe) IMPs were dissolving, leading to the incorporation of Mn and Fe in ZrCC. We believe that their incorporation has a role in the stabilization of aluminum oxide around IMPs and contributes to the passivation. Especially important seems to be Mn which was integrated in the bottom layer of ZrCC accompanied with the
highest increase in O. TEM investigation showed that the tri-layer structure consisting of top, middle and bottom layer changed from 74.2%, 17.2%, 8.6% of layer in the as-prepared ZrCC to 49.2%, 21.0% and 29.8% after 5 days immersion in 0.5 M NaCl solution. Based on these facts we propose the following mechanism of self-sealing/active corrosion protection in ZrCC applied on AA3005 (Fig. 15).

First, H2O diffuses inside ZrCC through cracks and pores down to IMPs and Mn dispersoids where hydrogen evolution and oxygen reduction reactions take place. This leads to an increased concentration of OH\(^-\) and release of hydrogen gas (Eqs. 6 and 7). Released H\(_2\)(g), especially around (Mn, Fe) IMPs, causes a bloating around IMPs after 2 days of immersion in NaCl (Fig. 7c). At the same time, OH\(^-\) ions cause a dissolution of aluminum oxide and its precipitation in the form of dense Al(OH)\(_3\) around (Mn, Fe) IMPs thus closing cracks and channels (Eq. 8). Excess in OH\(^-\) ions together with H\(_2\)O is transported towards outside part of ZrCC expelling F\(^-\) ions from the coating interior and thus transforming ZrF\(_4\)/ZrO\(_x\)F\(_{4-x}\) to ZrO\(_2\) · 2H\(_2\)O\(_x\) (Eq. 9). Excess of F\(^-\) ions is transported to a solution where it is balanced with Na\(^+\) ions. It can be also postulated based on obtained results that, along with the H\(_2\) evolution reaction, manganese is dissolved from IMPs and dispersoids (Eq. 11). Dissolved Mn\(^{2+}\) incorporates in the bottom layer of ZrCC forming spinel MnAl\(_2\)O\(_4\)\(_x\) (Eq. 12). It is known that spinels have great chemical and thermal stability. 92,93

\[
\begin{align*}
\text{H}_2\text{O} & \rightarrow 2\text{H}^+\,(\text{aq}) + \text{OH}^-\,(\text{aq}) \\
2\text{H}^+\,(\text{aq}) + 2\text{e}^- & \rightarrow \text{H}_2(g) \\
\text{Al}^{3+}\,(\text{aq}) + 3\text{OH}^-\,(\text{aq}) & \rightarrow \text{Al(OH)}_3(\text{s}) \\
\text{ZrF}_4/\text{ZrO}_x\text{F}_{4-x} & \rightarrow \text{ZrO}_2 \times 2\text{H}_2\text{O}_x + 4\text{F}^-\,(\text{aq}) \\
\text{ZrF}_4/\text{ZrO}_x\text{F}_{y(\text{aq})} + 4\text{OH}^-\,(\text{aq}) & \rightarrow \text{ZrO}_2 \times 2\text{H}_2\text{O}_x + 4\text{F}^-\,(\text{aq}) \\
\text{Mn} & \rightarrow \text{Mn}^{2+} \,(\text{aq}) + 2\text{e}^- \\
\text{Al}_2\text{O}_3 \cdot \text{H}_2\text{O}_x + \text{Mn}^{2+} \,(\text{aq}) & \rightarrow \text{MnAl}_2\text{O}_4\,(\text{s}) + 2\text{H}^+\,(\text{aq}) 
\end{align*}
\]

Conclusions

1. The formation of zirconium conversion coating on AA3005 was completed after 10 min of immersion in 200 ppm hexafluorozirconic acid bath at pH = 4.8 at room temperature.
2. Zirconium conversion coating applied to AA3005 showed excellent corrosion performance. Bode modulus at 3 mHz increased 50-fold during immersion in 0.5 M NaCl solution.
and sustained at 7.5 - 105 Ω·cm2 to 9.5-106 Ω·cm2 for at least 21 days with no pitting observed. The shift of the open circuit potential in nobler direction indicates anodic inhibition behavior of these coatings.

3. The as-prepared coating was uniform with thickness that varied from 30 nm at the coating matrix up to 200 nm at around IMPs. A tri-layer structure was noted, with the upper layer being enriched in Zr and O and bottom layer enriched in Al, F, Mn and Mg.

4. After immersion of 5 days in 0.5 M NaCl solution, the overall thickness of the coating did not change, but the proportion of individual layer changed, as well as the structure and composition. The thickness of the top layer decreased and that of the middle, and especially, the bottom layer increased. A transformation from ZrF4/ZrOxFy to ZrO2 · 1.2H2O(s) occurred of the middle, and especially, the bottom layer. The formation of a small amount of Al formed above and around (Mn, Fe) IMPs.

5. Upon immersion in NaCl, a hat consisting mainly of ZrO2 with a small amount of AI formed above and around (Mn, Fe) IMPs. However, at prolonged immersion channels around IMPs became filled with a dense layer of aluminum oxide leading to passivation. At the same time, the concentration of fluorine decreased, while that of O increased, especially in the middle and bottom layer of ZrCC.

6. Based on the above facts it can be stated that ZrCC on AA3005 alloy exhibits a self-sealing/active corrosion protection behavior within a 0.5 M NaCl solution. Its corrosion protection seems to be equal or better compared to that usually observed for chromium conversion coatings.

7. It seems that Mn from AA3005 together with Zr from ZrCC are able to trigger a self-sealing and active corrosion protection within the coating by incorporation in the bottom part of the conversion coating. This is very promising results and we suggest that future formulas of ZrCC should include Mn.

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