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Sealing of cerium oxide coating primers on anodized AA2024-T3 alloy by boiling in Lourier buffers

Stephan Kozhukharov $^{1,2,\boxtimes}$, Christian Girginov³, Stefania Portolesi⁴, Aleksandar Tsanev⁵, Vanya Lilova², Mariya Georgieva⁶, Emil Lilov² and Plamen Petkov²

1 LAMAR Laboratory for Advanced Materials Research, University of Chemical Technology and Metallurgy, 8 Kliment Ohridski blvd., 1756 Sofia, Bulgaria

²Department of Physics, University of Chemical Technology and Metallurgy, 8 Kliment Ohridski blvd., 1756 Sofia, Bulgaria

³Department of Physical Chemistry, University of Chemical Technology and Metallurgy, 8 Kliment Ohridski blvd., 1756 Sofia, Bulgaria

⁴University of Calabria, Via Pietro Bucci, 87036 Arcavacata, Cosenza, Italy

5 Institute of General and Inorganic Chemistry, Bulgarian Academy of Sciences, 1113 Sofia, Bulgaria 6 Institute of Mineralogy and Crystallography, Bulgarian Academy of Sciences, 1113 Sofia, Bulgaria Corresponding authors: [s.kozhukharov@uctm.edu;](mailto:s.kozhukharov@uctm.edu) Tel.: +359-899-837282; Fax: +359 2 68 54 88 Received: June 27, 2023; Accepted: July 11, 2024; Published: August 4, 2024

Abstract

*Although their exceptional re-passivation ability, Al-alloys are susceptible to corrosion due to the amphoteric nature of the alumina passivation films. This issue is exacerbated by the disruption of these films by intermetallics on the surfaces of highly doped ones, like AA2024-T3 aircraft alloy. The combination of anodized aluminium oxide (AAO) and cerium conversion coatings (CeCC) shows promise as a coating primer. However, the defective structures of CeO² and Al2O³ require additional sealing. This research proposes sealing the CeCC/AAO layer by boiling it for 10 minutes in two relatively neutral Lourier buffers, adjusted to pH 7.75, and in a mixture of them. The samples underwent a series of analyses to compare the impact of the sealing procedure on surface topology, properties (*e.g., *colour and wettability on two samples from each set), and corrosion protective ability. It was assessed after 24 hours of exposure to 3.5 % NaCl model corrosive medium on six samples from each set. The assessments included electrochemical impedance spectroscopy (EIS) and potentiodynamic scanning (PDS) techniques. The results indicate that the borate buffer improves the corrosion protection of the coating primers more effectively than the phosphate and mixed ones.*

Keywords

2024-T3 aluminium alloy; cerium conversion coatings; coating sealing; buffer solutions

Introduction

The use of aluminium alloys as structural materials has significantly increased due to their favourable properties, such as high strength-to-weight ratio, ease of fabrication, high workability, considerable ductility, excellent thermal conductivity, high corrosion resistance, and attractive natural finish [1]. However, despite their exceptional ability to form protective oxide layers, these films have amphoteric properties, making them soluble in both alkaline and acidic media. Their susceptibility to decomposition in Cl⁻ ions is another major disadvantage of the native oxide layers on aluminium [2-16]. Additionally, intermetallic inclusions on the surface interrupt the integrity of the oxide layer [17], leading to galvanic corrosion [17-21], which can propagate into other forms of localized corrosion [22-28]. These issues necessitate efficient corrosion protection for aluminiumbased products, equipment and constructions. Cerium compounds have been increasingly used as environmentally compliant inhibitors [29-43] and advanced protective coating components [44-55]. In particular, cerium conversion coatings (CeCC) have recently been recognized as a promising option for primer layers in advanced multilayered coating systems [56-65]. The CeCC deposition process must include preliminary surface treatment and subsequent CeCC sealing procedures [66- 69]. In this sense, anodization is the most appropriate preliminary treatment as it provides an oxide surface suitable for CeCC deposition [70-77]. However, final CeCC layer sealing is essential due to the inherent defectiveness of $Co₂$ structures [78-83]. On the basis of the fundamental concepts and recent trends in the formation of advanced multilayered coating systems [60,61,84-86], it can be inferred that CeCC layers deposited after anodization and appropriate CeCC coating primer sealing will be a primary focus for investigation in the coming years. Figure 1 summarizes the main trends in developing advanced multilayered coating systems for effective and reliable corrosion protection of aluminium alloys.

Figure 1. Trends for elaboration of advanced multilayered coating systems for corrosion protection of aluminium alloys [87,88]

Phosphate sealing is the most desirable approach for CeCC post-treatment, and optimizing its conditions has been the primary focus of research by Heller *et al.* [89,90] and Andreeva *et al.* [91-93]. This method is also proposed for reliable protection of other metals, demonstrating its versatility [94-98].

Summarizing all aspects related to the development of CeCC layers on preliminary anodized AA2024-T3 aircraft alloy, it was concluded that these layers need additional sealing by suitable

phosphate/borate solutions. The aim of the present research is to determine the effects of boiling CeCC/AAO layers in phosphate, borate, or mixed Lourier buffer solutions with pH 7.75.

Experimental

Sample preparation

Four sets of eight AA2024-T3 samples underwent CeCC/AAO primer layer formation under identical conditions. The layer formation procedures included consecutive preliminary surface treatments, anodization, and cathodic CeCC film formation. Preliminary treatments were conducted in alkaline baths (2 minutes in 50 g dm⁻³ NaOH at 50 °C) and acidic baths (2 minutes in diluted HNO₃ 1:1 v/v at 20 °C). Anodization was performed for 15 minutes in 15 wt.% H2SO⁴ at 20 °C. The final cathodic CeCC deposition was carried out for 5 minutes at 5 mA $cm⁻²$ in a pre-activated CeCC deposition solution under conditions similar to those described in [70,99-102]. The solution composition was 0.025 M CeCl₃.7H₂O combined with 0.025 M (NH₄)₂Ce(NO₃)₅.4H₂O, with 5 ml of 30 % H_2O_2 activator added to 250 ml of the solution. Each stage was followed by vigorous washing with tap and distilled water without removing the water film prior to CeCC formation.

For this research, the CeCC/AAO-coated AA2024-T3 alloy specimens were divided into four groups: untreated (Ref), boiled for 10 minutes in borate buffer (BB), phosphate buffer (PB), or in a mixed buffer (MB), following the buffer solution preparation procedures proposed by Lourier [103], which were translated into English in previous work [87]. The surface treatment procedures used in the present study are shown in Figure 2.

Figure 2. Sequence of procedures applied for CeCC/AAO coating primed formation and subsequent phosphatation in buffer solutions

The exact chemical compositions of the buffers used in the present study are shown in Table 1.

Buffer type	Buffer code	Buffer solution compositions		Required volume for 100 ml buffer, cm ³
Borate buffer	BB	First component: Na ₂ B ₄ O ₇ .10H ₂ O	19.069 g dm ⁻³ solution	
		Second component: HCl	0.100 mol dm ⁻³ solution	up to 100
Phosphate buffer	PB	First component: Na2HPO4.2H2O	11.866 g dm ⁻³ solution	92.40
		Second component: KH ₂ PO ₄	9.072 g dm ⁻³ solution	up to 100
Mixed buffer	MB	First component: phosphate buffer	(freshly prepared)	50.00
		Second component: borate buffer	(freshly prepared)	up to 100

Table 1. Compositions of the used buffer solutions

The buffer compositions in Table 1 were prepared to maintain a relatively neutral medium (pH ≈ 7.75), considering the amphoteric nature of the AAO layer.

Sample analyses

The analyses of all 32 investigated specimens were organized to allow a comparative description of surface properties (colour and wettability), surface topology, and corrosion protection ability.

Two samples from each set underwent surface analyses, while the remaining six samples were subjected only to corrosion tests. Surface properties were characterized by data acquisition from four points on each sample surface, and results were statistically analysed as described in previous work [104].

The respective measurements were performed using an RT 100 Lovibond tintometer and a "Theta Lite" sessile drop tester from Biolin Scientific (UK). The tintometer provided colour characteristics in accordance with the *CIE, L*, a*, b** colour space system, while the sessile drop tester measured the contact angle between the sample surfaces and distilled water drops. The results were statistically analysed following procedures described in previous works [70,99,104,105].

The surface topology of each sample couple was observed using optical metallographic microscopy (OMM). Optical micrographs were acquired at 100× magnification using a Boeco optical microscope equipped with a PK-710G camera from A4Tech, with a resolution of 640×480.

Scanning electron microscopy (SEM) images were obtained from the same sample couples used for surface analyses. Both surface topology and coating thickness were observed at 5000x magnification. All SEM images were obtained using a TESCAN SEM/FIB LYRA I XMU in backscattered electron mode.

To perform a comprehensive comparative analysis, the same sample couples were subjected to X-ray photoelectron spectroscopy (XPS) analysis. XPS studies were conducted using a VG Escalab II system with AlKα radiation at an energy of 1486.6 eV. The chamber pressure was 13.3 MPa. The C1s line of adventitious carbon at 284.6 eV was used as an internal standard to calibrate the binding energies. The photoelectron spectra were corrected by subtracting a Shirley-type background and quantified using the peak area and Scofield's photo-ionization cross-section. The accuracy of the binding energy measurements was ± 0.2 eV.

Corrosion tests

These tests were conducted on the remaining six samples from each set after 24 hours of exposure to a 3.5 % NaCl solution. Electrochemical methods used included electrochemical impedance spectroscopy (EIS) and potentiodynamic scanning (PDS). These methods were performed using an Autolab 30 (Metrohm) equipped with an FRA-2 module. Samples were exposed to the corrosive medium in threeelectrode cells, with 0.933 cm² of the sample surface as the working electrode, an Ag/AgCl/3 M KCl reference electrode, and a cylindrical platinum mesh counter electrode.

EIS data were recorded at 50 data points from 10 kHz to 0.01 Hz, using excitation signals ranging from 10 to 25 mV relative to the open circuit potential (OCP).

PDS curves were acquired within a potential range from -50 to +500 mV relative to the reference electrode, with a potential sweep rate of 10 mV s^{-1} .

Results and discussion

Among the examined buffer solutions, only the mixed buffer solution (MB) maintained stability, as pH measurements taken 24 hours after boiling showed that the MB solution retained its initial pH value. In contrast, the borate solution precipitated, whereas the phosphate solution became more alkaline, reaching a pH of 8.12.

Prior to assess the barrier properties, two specimens from each set were subjected to topological observations, as mentioned above.

The observed surfaces did not show significant differences, indicating that boiling in buffer solutions does not cause substantial changes in the CeCC/AAO layer morphologies. According to models developed by Nelson *et al.* [106] and Chen *et al.* [107], the CeO₂ phosphating proceeds via monolayer formation due to the highly defective nature of $Co₂$ crystal surfaces, as described by various authors [108-112]. Interaction with borate anions will likely also result in the formation of rather thin layers. However, all treated specimens appeared paler, possibly due to the formation of phosphate and/or borate monolayers.

Besides the colour characteristics, the wettability of the samples was also measured, and the statistically treated data are shown in Table 2.

Sample set	Value type	L*	a^*	b^*	θ / \degree		
Ref	Maximal value	85.55	-1.28	58.28	93.74		
	Minimal value	83.08	-3.11	54.56	48.13		
	Final result	84.52 ± 1.47	-2.08 ± 1.09	55.84 ± 2.26	63.45 ± 26.47		
BB	Maximal value	86.97	-2.78	62.46	23.27		
	Minimal value	84.13	-5.26	57.56	12.20		
	Final result	85.79 ± 2.05	-3.84 ± 1.67	59.99 ± 3.09	19.13 ± 6.05		
PB	Maximal value	86.72	-3.53	60.46	22.31		
	Minimal value	84.55	-5.35	56.63	7.380		
	Final result	85.40 ± 1.37	-4.62 ± 1.43	58.66 ± 2.12	12.61 ± 9.45		
MB	Maximal value	87.30	-4.12	59.07	21.07		
	Minimal value	84.71	-4.79	54.45	15.04		
	Final result	86.01 ± 2.34	-4.53 ± 0.51	57.24 ± 3.13	18.71 ± 4.69		

Table 2. Statistically treated data regarding the colours and wettability of CeCC/AAO layers subjected to additional sealing by borate and phosphate solutions

The treatments with phosphate/borate solutions did not result in notable changes in colour characteristics. A slight brightening of the colour, from *L** ≈ 85 for the references to *L** ≈ 86 for the samples treated in the mixed solution, was observed. This brightening was accompanied by a minor increase in the greenish hue, from $a^* \approx -2$ to $a^* \approx -4.5$, and a slight saturation of the yellow tonality, from $b^* \approx 55$ to $b^* \approx 60$.

Regarding contact angle, the samples changed from relatively to definitely hydrophilic after boiling in the respective buffer solutions. This effect on contact angle values is likely due to hydration during the final sealing in the buffer solutions. These initial surface characterizations were followed by topological observations of the reference and treated specimen couples using optical metallographic microscopy and scanning electron microscopy. The respective OMM and SEM images of the investigated samples are shown in Figures 3 and 4.

Figure 3. OMM images of reference (Ref) and samples treated in phosphate (PB), borate (BB) and mixed (MB) buffer solutions

The images reveal that specimens treated in borate and phosphate baths appear slightly paler than the reference specimens, although no visible alterations in topology are observable. These effects can be attributed to the inherent white colour of cerium phosphates [113-115] and borates [116,117], as well as the thin borate/phosphate monolayers expected based on described mechanisms [106,107].

SEM images, despite their higher resolution, confirm the OMM observations. All coatings, including the reference ones, appeared cracked. Therefore, the final sealing of the CeCC/AAO coating primers is necessary, as proposed in various studies [72,89-93] and supported by this research.

Figure 4. SEM images of reference (Ref) and samples treated in phosphate (PB), borate (BB) and mixed (MB) buffer solutions

In summary, the surfaces observed using both microscopy methods did not show significant differences. Boiling in the buffer solutions did not result in remarkable changes to the CeCC/AAO layer morphologies. However, all treated specimens appeared paler, likely due to the formation of white phosphate and/or borate monolayers.

The elemental contents calculated from the XPS spectra, acquired from the phosphate/borate sealed CeCC/AAO layers, as well as the reference ones, are summarized in Table 3. This approach has enabled the definition of the impact of the final sealing procedure on the superficial primer coating compositions.

Sample	Content, at.%							
	O	Al	Ce	D	B	Na	К	
REF ₁	75.0	7.1	17.9	0.0	0.0	0.0	0.0	
REF ₂	74.7	8.0	17.3	0.0	0.0	0.0	0.0	
BB ₁	60.0	11.2	2.6	0.0	21.7	4.2	0.0	
BB ₂	58.4	14.4	1.0	0.0	22.9	3.4	0.0	
PB ₁	61.6	2.0	0.8	16.5	0.0	18.1	1.0	
PB ₂	61.4	0.5	1.4	16.4	0.0	19.0	1.3	
MB ₁	61.2	1.1	4.3	16.5	6.5	7.9	2.5	
MB ₂	61.4	1.2	3.5	16.7	7.9	7.2	2.1	

Table 3. Data for the chemical compositions of the investigated specimens acquired through systematic XPS analyses

The data in Table 3 reveal that the reference CeCC/AAO layer covers the alloy substrate completely because the Al-concentration is nearly 7-8 at.%. The subsequent treatments in phosphate (PB) and mixed (MB) buffer resulted in a further decrement of the Al-content, down to approximately 2 at.%. In contrast, the borate buffer treatment results in a rise of the Al content up to about 11 to 14 at.%. The Al 2p core photoelectron spectra of all the sample sets are shown in Figure 5.

Figure 5. Deconvolutions of XPS core photoelectron spectra of Al 2p, O 1s and P 2p. The separate peaks are with the following colours: Al2p: Al-metal - gray; CeAlO³ - blue; Al(OH)³ - red:, AlOOH - magenta and green; O1s: Ce-Al-O - green; Me-O - orange; PO $_4^3$ and/or BO $_3^3$ -groups - pink; OH /H₂O - purple; P2p: PO $_4^3$ - blue; *PO³ 3- - green; PO³ 3- and BO³ 3- - yellow; P2O⁵ - orange*

The XPS spectra of the reference CeCC/AAO layers are split in couples of peaks positioned at 74.6 and 76.1 eV, respectively. The former peak could be attributed to the occurrence of AlPO₄ [118], whereas the latter is related to the presence of AlO(OH) [119]. This energy level slightly exceeds (by 0.9 eV) the characteristic one for AlO(OH), which equals 75.2 eV. This fact can be explained by the higher concentration of hydroxyl groups coordinated around the aluminium cation, which predetermines higher bonding energy, which must be overcome for the expulsion of the 2p-electrons from the electron shell of the Al-ion. The Al 2p spectrum of the Ref 1 sample has an additional lowintensity peak positioned at 72.5 eV due to metallic Al traces [116].

The Al2p spectra of the phosphate-treated samples are composed of a unique, low-intensity peak situated at 75.4 eV, which again is attributed to the presence of the AlO(OH) phase. The respective photoelectron peaks of the borate-treated specimens are more intensive and can be split into lower and higher energy ones. The former one is positioned at 74.6 eV for BB (Sample 1) and slightly shifted to 73.2 eV for BB (Sample 2), respectively. In both cases, the lower energy peak approximates the characteristic one for Al2O3, which, according to Latsunskyi *et al.* [120], should occur at 74.1 eV, possibly as a result of the occurrence of Al(OH)₃ since its typical value is 75.4 eV [121]. Thus, the shifting towards 74.6 eV could be due to $AIBO₃$ traces, whereas the opposite displacement to 73.2 eV corresponds to the characteristic one for $CeAlO₃$ [122].

The higher energy peaks are also slightly shifted to 76.1 eV for BB (Sample 1) and to 74.6 eV for BB (Sample 2), corresponding to Al(OH)3 and AlO(OH), respectively.

Both mixed buffer-treated samples (MB) have only one low-intensity Al 2p peak, positioned at 75.1 eV, related to the occurrence of the AlO(OH) phase. The Al 2p spectra are particularly interesting for the CeCC/AAO layers since they show interruptions of the conversion coating. In other words, these spectra are a response of the AAO layer surface under the cracks of the CeCC layer. Thus, it could be noticed that the phosphate-containing solutions (*i.e.,* PB and MB) result in the occurrence of only the AlO(OH) phase. The samples in these groups possess rather similar corrosion protective performance. On the other hand, the borate solution treatment leads to the simultaneous occurrence of CeAlO₃, Al(OH)₃, and AlO(OH). Especially in the case of BB (Sample 2), the CeAlO₃ phase composes about 1/3 of the registered Al content.

The O1s photoelectron spectra are rather intensive due to the oxide composition of the entire CeCC/AAO conjunction and the presence of hydroxides and crystal hydrate water. All the investigated samples, regardless of which group they belong to, have four peaks at 528.5, 529.6, 530.9 and 532.1 eV. These peaks reveal oxygen included in both Ce-Al-O and Me-O bonds [123], as well as (PO₄)³⁻, (BO₃)³⁻ [124], and OH^{$-$} moieties [125]. The O1s spectra analysis reveals higher OH $$ moieties content in the case of BB treatment.

The XPS analysis continued with the occurrence of phosphor as a result of the final CeCC/AAO layer sealing. The respective P2p core photoelectron spectra of the phosphate and mixed buffertreated samples are shown in Figure 5. The photoelectron patterns of the phosphate buffer (PB) treated specimens have peaks positioned at 134.0 and 134.9 eV, corresponding to the occurrence of (PO₄)³⁻ ions [126] and P₂O₅ [127], respectively. The presence of these phosphorus species is evidence for the participation of the phosphate ions in oxidation/reduction processes.

Couples of peaks are also observable for the mixed buffer-treated (MB) samples. The former peak, staying at 133.4 eV, can be attributed to the occurrence of orthophosphate (PO₄)³⁻ ions. The latter one, situated at 134.4 eV, reveals (PO₃)³⁻ moieties as was already commented. Shifting of the bonding energies by 0.4 eV towards higher values is probably a result of coordinated borate ligands surrounding the metallic ions.

The Ce3d photoelectron patterns were also analysed. These patterns are composed of two distinguishable peaks combined with four satellites, labelled "S" in Figure 6. The first peak is positioned at 882.3 eV, corresponding to Ce3d_{5/2}, while the second one is at 900.7 eV, corresponding to Ce3d_{3/2}. The spin-orbital splitting between the peaks, characteristic of CeO₂, is 18.4 eV. The satellite at 916.7 eV is characteristic of Ce⁴⁺ ions, allowing for the calculation of its percentage of the total Ce content. The calculations reveal that the entire cerium content consists of Ce(IV) species.

Figure 6. Ce3d, Na1s and B1s photoelectron patterns acquired from the respective specimens

The B1s spectra overlap with the P 2s ones. Therefore, only the B1s spectra of the borate buffertreated samples, which lack phosphorus, were analysed. For both samples, the peaks are at a bonding energy of 192.6 eV, typical for borate ions.

The XPS spectra of the investigated coating primers exhibit well-defined repeatability, indicating the consistent application of surface treatment procedures. Additionally, the treated coating primers exhibit distinct occurrences of (PO₃)³⁻, (PO₄)³⁻, and P₂O₅ moieties, which are known for their low solubility [93,128]. The borate-treated coating primers have elevated aluminium contents, suggesting a partially detrimental effect of the borate solution due to the presence of HCl, which is necessary for its preparation. However, the $(B_2O_7)^2$ ions form a barrier layer composed of Al₅BO₉ and/or Al₁₈B₄O₃₃ phases, as proposed by Luhrs *et al.* [129].

In summary, the presence of $CeAlO₃$, as indicated by the XPS spectra analysis, is the reason for the reliable and efficient corrosion protection provided by the proposed coating primers, as observed during the corrosion test results, commented below.

After 24 hours of exposure to a 3.5 % NaCl solution, 6 specimens from each set underwent electrochemical measurements. The averaged EIS data are summarized in Figure 7. The Nyquist plots reveal that despite its lower stability, BB results in notable sealing. The log|*Z*| - *f* curve is positioned above all others. At *f* = 0.01 Hz, the total impedance of the borate buffer-treated samples reaches $|Z|^{BB}$ _{av} = 1.8762 M Ω cm². This value is almost an entire order of magnitude superior to that of the reference samples $|Z|^{Ref}$ _{av} = 0.4983 MΩ cm². The log $|Z|$ - *f* curves of the remaining specimens, treated in phosphate buffer and the mixed system, occupy intermediate positions, still above those of the reference samples. Consequently, notable sealing occurs as a result of the applied final CeCC layer treatment procedures. In brief, the ordering of the log|*Z*|*-f* curves at 0.01 Hz in the respective Bode plots confirms the inferences drawn from the analysis of the Nyquist plots. The curves of the BBtreated coatings occupy the highest position. Finally, the curves of the MB-treated coatings are below those of the BB and PB-treated ones, owing to their lower barrier ability.

The *Z"*-*Z'* arcs of the BB-treated samples are the largest among the investigated sample sets. These are followed by the PB-treated ones. The mixed system shows a weaker effect on the barrier ability of the CeCC/AAO combined coating primer, as the *Z*"*- Z*'-arcs of the MB-treated specimens are closer to those of the references compared to the rest of the curves.

Figure 7. Nyquist (a) and Bode (b, c) plots acquired after 24 hours of exposure to MCM from CeCC/AAO layers subjected to sealing in boiling buffer solutions

The EIS spectra shown in Figure 7 were further analysed *via* data fitting to suitable model equivalent circuits (MECs), as depicted in Figure 8. Following the concepts outlined in [105], these MECs represent the charge transfer across homogeneous media through ohmic resistance elements, while the charge transfer across interfaces is represented by either pure capacitance or by constant phase elements due to the related charge accumulation phenomena.

The MECs shown in Figure 8 are composed considering the main concepts of González-Castaño *et al.* [130] and González-Rovira *et al.* [131]. In both schemes, *C*layer is attributed to the intact, nonconductive layer areas. This element is connected in parallel with time constants, resulting from the penetration of the electrolyte (*R*layer) across the layer defects and the electrochemical corrosion reactions (R_{ct}) after overcoming the double layer (CPE_{dl}). Hence, C_{layer} appeared suitable during the EIS data fitting, although the phase shift-frequency diagrams did not reach 90°. The reason is that the capacitance of the intact areas of the CeCC/AAO layers is connected in parallel to the elements (*i.e., R*layer, CPE_{dl}, R_{ct} and CPE_{diff}) attributed to the defects of the layers, resulting in phase shift values below 90 °.

Figure 8. MECs used for data fitting of the spectra of the reference (a) and sealed CeCC/AAO layer (b). RMCM - resistance of the model corrosive medium; Rlayer - resistance of the electrolyte penetrated across the CeCC/AAO layer defects; Rct - charge transfer resistance; Rdiff - resistance arising from the diffusion of the corrosive species across the coating primer defects; Clayer - capacitance originated from the insulating properties of the CeCC/AAO layer; CPEdl and CPEdiff - incomplete capacitances of the electric double layer and the diffusion processes inside the layer defects

Indeed, a multitude of cracks is notable in the SEM images of Figure 4. When these defects are narrow enough, re-passivation occurs, as outlined in [132,133]. Otherwise, these defects continue as crevices. Their occurrence results in the appearance of CPE_{dl} and R_{diff} time constants. They demonstrate stagnant corrosion, hindered by capillary diffusion across the corrosion products with "Keggin"-type (Al₁₃O₄(OH)₂₄(H2O)₁₂)⁷⁺ and (AlO₄Al₁₂(OH)₂₄(H2O)₁₂)⁷⁺ aluminum polyhydroxychloride structures, like established in [134,135]. The fitting of the EIS spectra in Figure 7 was performed using the MEC illustrated in Figure 8. All the acquired data are summarized in Table 4.

	Sample sets					
MEC component (24 h)	Ref samples	BB samples	PB samples	MB samples		
$R_{\rm MCM}$ / Ω cm ²	225.4 ± 24.5	206.6 ± 20.8	194.0 ± 18.8	163.5 ± 28.1		
C_{layer} / nF cm ⁻²	51.3 ± 4.5	8.5 ± 0.2	10.5 ± 0.3	13.0 ± 0.5		
R_{layer} / kQ cm ²	2.2 ± 0.2	7.1 ± 0.6	2.2 ± 0.3	2.2 ± 0.1		
CPE _{edl} / 10^{-7} s ⁿ Ω^{-1} cm ⁻²	36.5 ± 1.8	5.7 ± 0.4	10.1 ± 2.3	17.9 ± 0.6		
n	0.66 ± 0.01	0.59 ± 0.01	0.53 ± 0.01	0.57 ± 0.01		
$R_{\rm ct}$ / kQ cm ²	988.0 ± 18.6	133.2 ± 4.5	81.6 ± 1.3	85.5 ± 3.0		
CPEdiff / 10^{-7} s ⁿ Ω^{-1} cm ⁻²	$\overline{}$	14.9 ± 0.5	16.7 ± 1.27	16.5 ± 0.7		
n	$\overline{}$	0.89 ± 0.01	0.91 ± 0.01	0.99 ± 0.01		
R_{diff} / M Ω cm ²		65.1 ± 3.1	8.4 ± 0.4	3.4 ± 0.2		

Table 4. Results of the data fitting of the MEC in Figure 8 to EIS spectra shown in Figure 7

The data in the table indicate that the reference samples did not require the addition of diffusion elements (CPE_{diff} and R_{diff}) to fit their spectra, unlike the additionally treated ones. This fact suggests that the CeCC/AAO layer maintained its integrity after 24 hours of exposure. However, all the buffertreated specimens required the addition of diffusion elements, indicating corrosion occurrence combined with diffusion limitations.

Consequently, boiling resulted in coating cracking due to thermal tensions caused by the relatively high temperature. Nevertheless, the R_{diff} element exhibited rather high values, falling within the range of R_{ct} of the reference samples. Moreover, in the case of borate buffer (BB) treated samples, the sum of R_{ct} and R_{diff} was two orders of magnitude higher than the R_{ct} value for the reference specimens. This fact accounts for the superior barrier ability of these samples, as mentioned earlier. The R_{ct} and R_{diff} values in the cases of PB and MB are inferior to those of the borate-treated samples, further emphasizing the superior barrier ability of the BB-treated ones.

This inference is supported by the highest R_{ct} and R_{layer} values obtained from the borate buffertreated samples compared to those of the other specimen sets. Additionally, the highest corrosion

protection of the borate buffer-treated samples is evident from the EIS spectra shapes in Figure 7, as indicated by the highest position of the log|*Z*|*-f* curve at 0.01 Hz in the Nyquist plot (Figure 7a), corresponding to 4.198 MΩ cm², and the largest arc of these specimens in the respective Bode plots (Figure 7b and c).

Following every EIS spectrum recording, potentiodynamic curve acquisition was performed. The averaged PDS curves are summarized in Figure 9. The PDS curves of the sealed specimens are positioned below the reference ones, indicating lower current densities. Consequently, the respective sealings should have a beneficial effect on the barrier properties of the treated CeCC/AAO layers. Furthermore, the PDS curve with the lowest current densities is registered for the BB-treated specimens.

All the curves shown in Figure 9 exhibit passivation regions combined with local corrosion features. In the cases of buffer solution treatments, these regions possess weak slopes, suggesting incomplete passivation. The occurrence of local corrosion is noticeable by the sharp bending of the anodic branches, indicating pitting corrosion. However, the difference among the sample types is evident from the curve positions. The curves of reference samples are positioned at the highest current densities and the most negative potential values, whereas the BB ones are situated at the lowest current densities and the noblest potentials. The final treatments in phosphate and mixed solutions result in intermediate positions of the PDS curves, relatively closer to the curves of the reference samples.

Figure 9. PDS curves obtained from the reference and sealed CeCC/AAO layers

In summary, an ambiguous effect is to be expected from the final sealing procedure. On one hand, the buffer solutions should result in the formation of thin sealing layers, leading to an increment of the resistance values. On the other hand, boiling should cause CeCC/AAO layer cracking. In the present case, the borate buffer (BB) reveals the most notable beneficial effect compared to the rest (PB and MB). The borate and phosphate buffers (MB) mixture reveals the weakest impact on the coating primer performance. The inferior behaviour of the phosphate-treated samples compared to the borate-treated ones can be explained by considering the observed alkalization (mentioned at the beginning of the "Results and Discussion" section) of its solution, which affects the amphoteric AAO layer under the cracks of the CeCC film, as noticeable in the SEM images (Figure 4).

The curves shown in Figure 9 were further subjected to Tafel slope analysis and the results are shown in Table 5.

Sample type	OCP. mV	E_{corr}/mV	i_{corr} / nA cm ⁻²	$R_{\rm D}$ / MQ cm ²	E_{pit} / mV	i_{pit} / nA cm ⁻²
Ref	-767 ± 13	$-761 + 15$	$(2.1 \pm 0.1) 10^3$	0.17 ± 0.02	-488 ± 19	$(8.4 \pm 0.7) 10^3$
BB	-479 ± 17	-475 ± 16	2.75 ± 0.16	1.63 ± 0.28	-114 ± 10	71.12 ± 6.72
PB	-687 ± 12	$-676 + 18$	7.41 ± 0.18	$1.41 + 0.27$	-466 ± 18	74.01 ± 2.04
MВ	-611 ± 18	$-615 + 16$	3.74 ± 0.15	1.40 ± 0.22	-473 ± 27	52.83 ± 6.94

Table 5. Results of the Tafel slope analysis of the PDS curves acquired after 24 hours of exposure to 3.5 % NaCl model corrosive medium

The results in Table 5 confirm the inferences drawn from the analysis of Figure 9, substantiating them with numerical values of the electrochemical measures. Notably, a significant difference is observed among the values of the open circuit potential (OCP) and the corrosion potential (*E*corr), as a result of the final treatments of the CeCC/AAO layers. The data in the table show a rather large range of nearly 300 mV between the reference samples (Ref) and the borate solution-treated (BB) ones. The former tends to about -770 mV, whereas the latter approaches -480 mV. The intermediate OCP and *E*corr values are possessed by the PB and MB buffer solution-treated specimens, with about -670 and -620 mV, respectively.

Further analysis of the data in Table 5 reveals that the most negative *E*corr values of the reference samples are associated with three orders of magnitude higher corrosion current density (*icorr*). Consequently, the final CeCC/AAO layers treatment results in an obvious enhancement of the barrier ability of the CeCC/AAO layers, suppressing the *i*_{corr} down to about 5 nA cm⁻². Once again, the lowest corrosion current density belongs to the BB-treated specimens. A similar difference is also noticeable in the polarization resistance (R_p) values. Hence, the R_p of the additionally treated specimens is more than an order of magnitude higher than those of the references. Besides, the BBtreated CeCC/AAO layers possess the highest R_p values.

Additional analysis was performed on the anodic branches of the PDS curves, providing supplemental data regarding the pitting corrosion parameters, namely pitting potential (E_{pit}) and current density (*i*p**i**t). The former measure provides the determination of the strength against pitting nucleation (SAPN = E_{corr} - E_{pit} [105]), whereas the latter enables the calculation of the pitting nucleation resistance (R_{pit}) . The SAPN values for the references and the borate buffer-treated ones are approximately SAPN_{Ref} \approx 273 mV and SAPN_{BB} \approx 361 mV, respectively. Since a higher SAPN value indicates a larger passivation region, the BB-treated samples also excel in the reference ones in this parameter. The SAPN of the PB and MB samples exhibit even lower values.

The ratio between *i*_{pit} and *E*_{pit} allows the estimation of the *R*_{pit} values for the respective sample sets. Thus, the BB specimens have R_{pit} ^{BB} ≈ 1.603 MΩ cm², whereas R_{pit} ^{Ref} ≈ 0.581 MΩ cm². The rest of the treated sample sets have shown R_{pit}^{PB} ≈ 6.296 MΩ cm² and R_{pit}^{MB} ≈ 8.953 MΩ cm², respectively. These values are slightly higher than those of the borate buffer-treated specimens. Nevertheless, the final treatment in all cases results in rather higher R_{pit} values compared to those of the references.

In summary, comparing the results of both electroanalytical methods (Tables 4 and 5) enables the ordering of the effects of each buffer on the barrier properties of the CeCC/AAO layers. Thus, in accordance with both electroanalytical methods, the highest protective ability belongs to the BB set, followed by PB and finally MB. The same ordering from the best to the worst barrier properties can be seen in Figures 7 and 9, summarized as BB > PB > MB > Ref. The analyses above reveal that this order is confirmed by both electrochemical methods employed. Consequently, in simple words, the treatment by borate buffer possesses a better sealing effect on the CeCC/AAO layer than that by phosphate solution and no synergistic effect between these solutions appeared, although the expectations.

Conclusions

This study is focused on the sealing of CeCC/AAO layers using Lourier buffers with relatively neutral pH. It presents the results of systematic research conducted on four sample sets, each comprising eight CeCC/AAO combined coating primers deposited on AA2024-T3 alloy. Three sets underwent additional sealing in borate (BB), phosphate (PB), and mixed (MB) buffers. Two samples from each set underwent comparative analyses of their properties, including colour and wettability, as well as topological analyses using optical and electron microscopy methods and detailed chemical composition determination via X-ray photoelectron spectroscopy (XPS).

The remaining six samples from each set were subjected to electrochemical tests using electrochemical impedance spectroscopy (EIS) and potentiodynamic scanning (PDS) after 24 hours of exposure to 3.5 % NaCl model corrosive medium.

The results indicate that the applied sealing procedures do not affect the topology of the CeCC/AAO layer but alter the surface chemical composition and resulting surface properties. While the micrographs show identical topologies, the sealed specimens exhibit increased hydrophilicity and paler coloration. These alterations in surface properties are attributed to changes in chemical composition detected by XPS analysis, which reveals the presence of $(PO₄)³⁻, (BO₃)³⁻, (B₂O₇)²⁻$ and ОН¯ moieties, likely forming thin monolayers based on literature analysis.

Corrosion tests demonstrate that among the tested buffer solutions, the borate solutions have a superior effect despite their Cl⁻ ion content. The beneficial effect of the phosphate solutions was comparatively weaker. Additionally, mixing borate and phosphate buffers did not result in synergistic effects as initially expected.

Sealing with phosphate and/or borate buffers enables the formation of sealing layers, but it is important to perform these procedures at lower temperatures. Otherwise, the solutions may change their composition due to precipitation, resulting in pH alterations that affect the amphoteric AAO layer beneath the cracks of the CeCC film.

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