Article

LDH Post-Treatment of Flash PEO Coatings

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Abstract: This work investigates environmentally friendly alternatives to toxic and carcinogenic Cr (VI)-based surface treatments for aluminium alloys. It is focused on multifunctional thin or flash plasma electrolytic oxidation (PEO)-layered double hydroxides (LDH) coatings. Three PEO coatings developed under a current-controlled mode based on aluminate, silicate and phosphate were selected from 31 processes (with different combinations of electrolytes, electrical conditions and time) according to corrosive behavior and energy consumption. In situ Zn-Al LDH was optimized in terms of chemical composition and exposure time on the bulk material, then applied to the selected PEO coatings. The structure, morphology and composition of PEO coatings with and without Zn-Al-LDH were characterized using XRD, SEM and EDS. Thicker and more porous PEO coatings revealed higher amounts of LDH flakes on their surfaces. The corrosive behavior of the coatings was studied by electrochemical impedance spectroscopy (EIS). The corrosion resistance was enhanced considerably after the PEO coatings formation in comparison with bulk material. Corrosion resistance was not affected after the LDH treatment, which can be considered as a first step in achieving active protection systems by posterior incorporation of green corrosion inhibitors.

Keywords: PEO; LDH; active protection; corrosion; aluminium

1. Introduction

Plasma electrolytic oxidation (PEO) is a plasma-assisted electrochemical surface treatment characterized by the utilization of high voltages (100–600 V) in alkaline electrolytes to produce ceramic-like coatings on light alloys such as aluminium [1], magnesium [2] and titanium [3]. This technology may become a potential alternative to conventional toxic and highly carcinogenic chromic acid anodizing (CAA) for niche applications [4–6].

The PEO process can be conducted under direct current (DC) [7], alternate current (AC) [8], unipolar [9] or bipolar pulsed regimes [10] involving polarization of the alloy to voltages above the dielectric breakdown of the oxide. This results in the formation of multiple short-lived microdischarges on the metallic surface that trigger the formation of highly stable ceramic phases [1,11,12]. The resulting PEO coatings present high hardness, thermal stability and adherence to the substrate, which lead to enhanced corrosion and tribological properties. Moreover, the coatings’ microstructures and compositions can be tailored as required by controlling electrical parameters during the coating synthesis by electrolyte selection and by the application of pre- and post-treatments [13,14].

However, the costs associated with PEO technology are relatively high, which is mainly due to high energy consumption. General current densities and voltages for the PEO process are in the range of 1.5–15 A·dm⁻² and 400–500 V for Mg [15], 6–20 A·dm⁻² and 200–500 V for Ti [16] and 10–60 A·dm⁻² and 200–400 V for Al [17]. Typical PEO treatment times are within a 15–60 min range.
In the particular case of Al and its alloys, the efficiency of conversion of the anodic charge into the final coating material may be as low as 20%, due to gas generation and dissolution processes [18–20]. Therefore, energy consumption of the process must be reduced in order to make this technology commercially viable for high-volume applications [10,14,20,21]. Strategies to achieve this goal involve waveform design and cell geometry [22], electrolyte design [23–27]) and a pre-anodizing approach [18]. An additional strategy consists of “flash” PEO—processes of a short duration (1–3 min) that produce thin coatings (1–5 µm) suitable and sufficient for many corrosion resistance-demanding applications.

PEO coating properties can be improved even further by incorporating active functionalities. This can be achieved by introducing smart agents into the coatings matrix, which are released in situ, triggered by an external stimulus [23–25]. Coatings based on LDH have increasing potential not only for developing smart functionalities on ceramic materials, but in many other fields, such as biomaterials, due to their low toxicity [23].

LDHs can be expressed by the general formula $[M^{2+}]_{1-x}M^{3+}X_2(OH)_2X^+(A^{m-})_{X/m}nH_2O$, where $M^{2+}$ and $M^{3+}$ represent divalent metallic cations (e.g., Mg$^{2+}$, Zn$^{2+}$, etc.) and trivalent metallic cations (e.g., Al$^{3+}$, Co$^{3+}$, etc.), respectively, and $A^{m-}$ is the interlayer anion (e.g., NO$_3^-$, Cl$^-$, etc.) [26]. It consists of uniform flakes that grow roughly vertically on the substrate and act as intelligent nanocontainers that release, in a controlled way, the previously loaded corrosion inhibitors. The active corrosion protection mechanism is based on anion-exchange reactions induced by particular triggers, such as increases in the aggressiveness of the surrounding environment or initiation of the corrosion process [26,27].

The most common approach to synthesizing LDH is the in situ growth method, due to its low cost and ease of synthesizing at laboratory and industrial scales [28]. In this method, the source of Al$^{3+}$ ions required to grow the LDH structure is the metallic substrate, and it results in coatings with better adherence compared to LDH coatings synthesized by other routes such as hydrothermal treatment [29–31], which is characterized by the use of autoclave to get high temperatures, or the urea hydrolysis method, which is used to form well-crystallized large LDH [32–34].

The knowledge of multifunctional surfaces with active protection based on ceramic-like coatings (PEO) is at an embryonic stage [35]. Y. Zhang [36] investigated the growth behavior of LDH layers on PEO-coated aluminum alloy. It was found that LDH grains are preferentially formed on the micro-pores of PEO coatings to provide effective film repairs. The most dominant factor which determines the PEO/LDH system behavior is the interaction between the electrolyte anions and Al(OH)$_2^+$ cations from PEO coating surface. This phenomenon has a major impact on the kinetic mechanism of the formation of LDH coating [37,38]. F. Chen [39] demonstrated that flakes were preferentially formed in the PEO pores at the initial stage of LDHs growth; it was also found that long-term corrosion resistance was significantly enhanced due to the limited penetration of corrosive ions [38]. The corrosive degradation of the substrate coated with PEO is accelerated by chloride ions in the environment. From a corrosion-relevance perspective, the entrapment of chloride anions via the ion-exchange mechanism is the most critical factor for designing efficient corrosion protection strategies. In fact, partial entrapment is possible when the PEO layer is sealed with LDH [40].

Although the concept of LDH-based post-treatment for sealing the PEO layers has been recently proven, many variables remain unexplored. In particular, this work is focused on the formation of in situ Zn-Al LDH on low energy consumption PEO coatings. For that, three PEO coatings were selected from 31 processes (with different combination of electrolytes, electrical conditions and time) based on corrosive behavior and energy consumption. The selected materials were thoroughly characterized and evaluated in terms of corrosion resistance.

2. Materials and Methods

2.1. Material

The composition of the 1050-H18 aluminum alloy (Famimetal S.L., Madrid, Spain) in wt.% is: 0.07 Zn, 0.05 Mn, 0.25 Si, 0.05 Cu, 0.40 Fe, 0.05 Mg, 0.05 Ti, 0.03 V and Al balance.
2.2. Specimens Preparation

The samples were cut from sheets into specimens of $30 \times 20 \times 1.5$ mm$^3$ dimensions, ground to P1200 silicon carbide abrasive paper, rinsed in distilled water and methanol and dried in warm air. Prior to the PEO processing, specimens were etched in 15 wt.% sodium hydroxide solution for 20 s, rinsed in deionized water and dried in warm air. The working area was then limited to $\sim 3$ cm$^2$ using a commercial resin (Lacquer 45, MacDermid plc., Birmingham, UK).

2.3. Surface Treatment Based on PEO

PEO coatings were obtained using different electrolytes and conditions in order to grow thin PEO coatings with low energy consumption (Table 1) under vigorous agitation. The experimental system was carried out with a DC voltage/current-controlled power supply (SM400-AR-8 Systems electronic) equipped with a thermostatic jacket (25 ± 1 °C) under continuous electrolyte agitation. An AISI 316 steel plate of 7.5 × 15 cm$^2$ size was used as a counter electrode. After the PEO process, all samples were rinsed in distilled water and isopropanol and dried in warm air.

Table 1. PEO conditions of AA1050 alloy.

| Coating | Electrolyte (g/L) | Coating | Electrolyte (g/L) | Coating | Electrolyte (g/L) |
|---------|------------------|---------|------------------|---------|------------------|
| A1.1    | 4                | KOH     | (Na$_3$PO$_4$)$_3$ | KOH     | Na$_2$SiO$_3$   |
| A1.2    | 4                | 3.3     | P1.1             | 30      | 1                | S1.1             | 10.5             | 2.8             |
| A1.3    | 4                | 5.6     | P1.3             | 30      | 5                | S1.3             | 10.5             | 4.6             |
| A2.1    | 7                | 1.0     | P2.1             | 40      | 1                | S2.1             | 15.0             | 2.8             |
| A2.2    | 7                | 3.3     | P2.2             | 40      | 3                | S2.2             | 15.0             | 3.5             |
| A2.3    | 7                | 5.6     | P2.3             | 40      | 5                | S2.3             | 15.0             | 4.6             |
| A3.1    | 10               | 1.0     | P3.1             | 50      | 1                | S2.4             | 15.0             | 6.0             |
| A3.2    | 10               | 3.3     | P3.2             | 50      | 3                | S2.5             | 15.0             | 8.0             |
| A3.3    | 10               | 5.6     | P3.3             | 50      | 5                | S2.6             | 15.0             | 10.0            |
| –       | –                | –       | –                | –       | –                | –                | –                | –               |

PEO treatment conditions: $V$ (V): 400; $j$ (A·cm$^{-2}$): 0.1; $t$ (s): 180; PEO treatment conditions *: $V$ (V): 350; $j$ (A·cm$^{-2}$): 0.05; $t$ (s): 200; ** Na$_2$SiO$_3$ alludes to the use of water glass with specific gravity of 1.3 g/L.

2.4. Synthesis of Zn-Al-LDH Growth

Zn-Al LDH-nitrate (LDH-NO$_3$) was synthesized on AA1050-H18 aluminium alloy. The specimens were immersed in the solution for different times under continuous stirring in order to form LDH (Table 2), then rinsed in distilled water and isopropanol and dried in warm air.

Table 2. LDH synthesis conditions of AA1050-H18.

| LDH Treatment | Chemical Composition (M) | Exposure Time (min) |
|---------------|--------------------------|---------------------|
| 1             | Zn(NO$_3$)$_2$·6H$_2$O: 0.01 NH$_4$NO$_3$: 0.06 | 30                  |
| 2             | Zn(NO$_3$)$_2$·6H$_2$O: 0.01 NH$_4$NO$_3$: 0.06 | 60                  |
| 3             | Zn(NO$_3$)$_2$·6H$_2$O: 0.01 NaNO$_3$: 0.06 | 30                  |
| 4             | Zn(NO$_3$)$_2$·6H$_2$O: 0.01 NaNO$_3$: 0.06 | 60                  |

All treatments were developed in 100 mL of aqueous solution at 95 °C. pH values were adjusted to 6.5 using 1 vol.% ammonia.

2.5. Characterization

Planar and cross-sectional views of the specimens were examined using a JEOL JSM 6335F (Tokyo, Japan) field emission scanning electron microscope (FESEM) working at 20 kV and equipped with an energy dispersive (EDS) spectrometer (OXFORD X-MAX, Oxford, UK). Coating cross-sections were ground through successive grades of SiC paper and polished to a 1 µm diamond finish.
Phase composition was examined by X-ray diffraction (XRD), with a Philips X’Pert MRD (Amsterdam, The Netherlands, Cu Kα = 1.54056 Å). The XRD patterns were taken using grazing incidence with a step size of (0.01°–1°) and a dwell time of 6 s per step at room temperature.

The specific energy consumption was calculated by integration of the voltage-time and current-time transients acquired by the power supply (SM 400AR-8 Systems electronic) during the PEO treatment (Equation (1)). With the obtained results, the energy consumption was calculated in terms of kW·h·m⁻² according to Equation (1), and the specific energy consumption in kW·h·m⁻²·µm⁻¹ was obtained dividing \( P_{tot} \) by the coating thickness.

\[
P_{tot} = \int_{t_0}^{t} [V \cdot j] \left[ \frac{W \cdot s}{m^2} \right]
\]

2.6. Electrochemical Behavior

Electrochemical impedance spectroscopy (EIS) was used to evaluate the corrosion resistance of the different coatings in an aqueous saline solution (NaCl 3.5 wt.%) at 25 °C. For that, a GillAC (ACM Instruments, Cumbria, UK) computer-controlled potentiostat and a three-electrode cell were used. The specimen was connected as a working electrode with an exposed area of 1 cm². A graphite electrode and a silver–silver chloride (Ag/AgCl) electrode used as the counter and the reference electrode, respectively. The solution inside the reference electrode was KCl 3 M, which provided a potential of 0.210 V with respect to the standard hydrogen electrode. The tests applying a sinusoidal perturbation of 10 mV RMS amplitude in the frequency range of 30 kHz–0.01 Hz were carried out after 1 h of immersion. All measurements were duplicated to ensure reproducibility.

3. Results and Discussion

3.1. PEO Coating Screening

The first screening process to select one PEO coating per electrolyte type was conducted in accordance with three factors: (i) presence of microdischarges during coating formation, (ii) visually uniform coating morphology and (iii) coating thickness \( \geq 1 \) µm.

Among the different alternatives based on different electrolyte compositions, PEO coatings developed in phosphate and silicate electrolytes showed more promising results (with lower breakdown voltage values, beneficial for low energy consumption) (Table 3) and therefore additional process conditions were also tried (\( V \) (V): 350; \( j \) (A·cm⁻²): 0.05; \( t \) (s): 200) (Table 1).

| Coating | Electrolyte Composition (g/L) | \( \sigma \) (mS/cm) | pH | \( U_{bd} \) (V) | Thickness (µm) | Growth Rate (µm·min⁻¹) | Energy Consumption (KWh·m⁻²·µm⁻¹) |
|---------|-------------------------------|----------------------|----|-----------------|----------------|------------------------|-----------------------------------|
| A3.1    | NaAlO₂: 10 KOH: 1             | 13.9                 | 12.70 | 320            | 1.1 ± 0.3      | 0.37                | 4.98                             |
| P2.1    | (Na₃P₂O₇)₃·40 KOH: 1          | 11.1                 | 12.5  | 257            | 2.4 ± 0.4      | 1.31                | 4.70                             |
| S4      | Na₂SiO₃·5H₂O·25 KOH: 2.8      | 32.9                 | 12.7 | 108             | 1.3 ± 0.2      | 0.39                | 2.20                             |

Finally, only the A3.1, A3.2, P2.1, S1.3, S2.6 and S4 PEO coatings (Table 1) fulfilled the previous requirements.

The last step of the screening process consisted of a corrosion evaluation based on the value of the modulus of the impedance obtained by electrochemical impedance spectroscopy (EIS). Figure 1 shows the \( |Z| \times 10^{-2} \) Hz values of the studied materials, providing an estimation of the corrosion resistance, where higher values of \( |Z| \) indicate a lower corrosion rate [41].
With the aim of studying the influence of electrolytes on LDH growth, three PEO coatings were selected (one per electrolyte composition: aluminate A3.1, phosphate 2.1 and silicate S4). The specific energy consumption of selected coatings was calculated (Table 3) by integration of the voltage-time and current-time transients (Figure 2) recorded during the PEO process to verify that the developed coatings were energy efficient. The obtained values reveal the effect of electrolyte composition on energy consumption, coating growth rate and breakdown voltage values (Table 3).

![Figure 1. Scatter diagram of impedance modulus at 10^{-2} Hz; selected PEO coatings on AA1050 alloy, citing two measurements per condition.](image)

![Figure 2. Voltage-current time curves for (a) A3.1, (b) P2.1 and (c) S4 PEO coatings on AA1050 alloy.](image)

As can be seen in Figure 2, the current drop was observed only in aluminate and silicate electrolyte cases (Figure 2a,c) when 400 and 350 V limitations (Table 1) were achieved after 40 and 75 s of treatment, respectively, and the power supply switched to a constant voltage-control mode. Typically, when the current density was below 20 mA·cm^{-2}, the microdischarges extinguished; the anodizing, however, was carried on till set time in order to repair microdefects in the oxide material [42].
The fact that the limiting voltage was reached (hence the current drop) could be explained by high coating density and, consequently, higher resistance of the oxide to charge and mass transfer that were achieved at early stages [18]. High value of energy consumption in the case of aluminate is mainly due to the fact that aluminate species in the electrolyte gave rise to the formation of coating composed of nearly pure alumina, which has very low electron conductivity (i.e., the current flows mainly by ion and not by electron transfer) [12]. Further, the high breakdown voltage (320 V) in the aluminate electrolyte compelled the use of a higher voltage limit (400 V) in order to ensure a long enough period of sparking in order to achieve a uniform coating of a significant thickness; this yielded a higher specific energy consumption value.

As a result, the dielectric breakdown voltage was high, the limiting set voltage was achieved quickly and, as a consequence of the current drop, the sparking period was short, hence the low coating growth rate [22,43]. Similarly, high value of energy consumption in the case of phosphate electrolyte, where sparking was observed until the end of the treatment and coating growth rate was relatively high, was due to the absence of current drop, because the limiting 350 V were never reached. In this case, higher resistance of the oxide to charge and mass transfer were achieved at 60 s, giving rise to intense microdischarges and voltage variations, and therefore the treatment was stopped at 115 s in order to maintain coating uniformity.

The lowest energy consumption of 2.2 kW·h·m⁻²·µm⁻¹ was achieved in case of silicate electrolyte. This was mainly the result of its high electrical conductivity and, therefore, low $U_{bd}$. The onset of microdischarges early in the treatment and the relatively long sparking period before current decay resulted in the intermediate coating growth rate value (Table 3).

The specific energy consumption values obtained under DC conditions in the present work are similar to those reported in studies carried out under AC conditions. For example, E. Matykina et al. developed a PEO coating on pure aluminium using silicate electrolyte, and obtained a growth rate of 1.3 µm·min⁻¹ and energy consumption values of 4.77 KW·h·m⁻²·µm⁻¹ [18]. Y.L. Cheng et al. reported a growth rate of 11.3 µm·min⁻¹ and energy consumption values of 5 KW·h·m⁻²·µm⁻¹ for Al-Cu alloy in concentrated aluminate electrolyte [44]. It is well known that DC conditions promote low growth rates in comparison with AC conditions [18,22]; however, in this study the values obtained were considerably lower compared with the available data for different PEO treatments on commercial Al alloys, which can be as high as 26.7 kW·h·m⁻²·µm⁻¹ [18]. The present findings demonstrate that in order to reduce specific energy consumption under DC conditions it is necessary to (i) limit the final forming voltage that ensures a current drop, and (ii) use electrolytes with conductivity, which ensures low microdischarges onset voltages and extended sparking periods, as in the case of the S4 electrolyte.

3.2. PEO Coatings Characterization

Figure 3 shows the planar view and cross-section scanning electron micrographs of AA1050 coated by selected PEO coatings. All selected treatments show a thin oxide layer of 1–2.5 µm (Table 3). This is particularly evident in aluminate electrolyte-based PEO coating (A3.1), where the Al-Fe intermetallic compounds from the substrate are still visible in the coating (Figure 3a, inset) due to its low thickness. This coating is also more heterogeneous (Figure 3b) than the rest, which is attributable to its high breakdown voltage values (Figure 2) [37,45,46] and, as mentioned before, its low coating growth rate. Phosphate electrolyte-based PEO coating (P2.1) showed a homogeneous (Figure 3c) surface appearance (Figure 3c), and the highest thickness value (Figure 3d). This was mainly due to the presence of polyphosphate species that participated in PEO coating formation and favored its high coating growth rate [47]. Silicate electrolyte-based PEO coating (S4) (Figure 3e), with the lowest breakdown voltage, showed a homogeneous surface morphology with very sparse submicrometric pores. The latter may be attributable to the formation of a thin superficial glassy layer of SiO₂, which can be surmised from the EDS analysis where the presence of 1.5 at.% Si in the coatings and a greater content of oxygen than in the other two coatings was confirmed.
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| PEO Coating | EDS (at.%) |
|-------------|------------|
| A3.1        | O 45.7     |
|             | Al 52.8    |
|             | Fe 1.1     |
|             | P  –       |
|             | Si  –      |
| P2.1        | O 53.3     |
|             | Al 46.6    |
|             | Fe  –      |
|             | P 1.1      |
|             | Si  –      |
| S4          | O 64.3     |
|             | Al 33.6    |
|             | Fe  –      |
|             | P  –       |
|             | Si 1.5     |

Figure 3. Planar view (a,c,e) and cross-section view (b,d,f) of secondary electron images of the PEO coatings (A3.1, P2.1, S4), respectively. EDS analysis was performed on the areas corresponding to the planar views of the coatings.

3.3. LDH Screening

The effect of reactant composition and treatment time during the growth of LDH coatings were investigated. Figure 4 depicts the XRD patterns of the different LDH treatments (Table 2) grown on the bulk material. The presence of peaks at 9.6° and 19.9° corresponding to the characteristic (003) and (006) reflections of LDHs intercalated with NO₃⁻ [32,48,49] indicates the formation of LDH under the different conditions.
Figure 4. XRD patterns of Zn-Al-LDH-coated AA1050 alloy at different conditions.

It was revealed that treatments containing NaNO₃ in the solution led to more defined and intense peaks, probably due to the presence of sodium ions in the LDH gallery [50]. On the contrary, the presence of NH₄NO₃ drove the formation of broadened peaks [51]. In fact, just in the case of LDH grown in NH₄NO₃, a small peak was revealed at 9.9° that could be associated with an LDH phase intercalated with carbonate, due to the formation of LDH layers under atmospheric conditions [52].

LDHs formed under long treatment times (LDH 2 and 4) showed very strong peaks in comparison with LDHs formed under short treatment times (LDH 1 and 3), mainly because an increment in the LDH degree of crystallinity took place [51].

The correlation between XRD patterns of studied LDH coatings and planar view scanning electron micrographs were investigated. Figure 5 shows secondary electron images of the LDH treatments (Table 2) grown on the pure aluminium.

The typical flake-like LDH structure could be clearly observed for LDH carried out in NH₄NO₃ at short treatment times, whereas LDH developed in NaNO₃ formed this structure at long exposure times. According to Figure 5a, the LDH structure carried out in the presence of NH₄NO₃ is in good agreement with the XRD patterns that showed broadened peaks and, consequently, a highly open LDH structure. Additionally, Figure 5b clearly shows (also consistent with the XRD pattern) a non-defined LDH structure, which is usually attributed to the incorporation of carbonate ions into the LDH gallery [48]. However, LDH developed in NaNO₃ showed typical curved plate-like LDH microcrystals at long exposure times, and a non-defined structure at short exposure times (Figure 5c,d). This was mainly due to an increment in the degree of LDH crystallinity that was observed in XRD patterns (Figure 4).

In order to evaluate the correlation between the corrosion protection and the structure of studied LDHs, a screening process based on corrosion performance (EIS) was carried out. Figure 6 depicts the Bode and Nyquist plots for AA1050 alloy with studied LDH coatings.
Figure 5. Secondary electron images of the LDH 1 (a,b) LDH 2 (c,d), LDH 3 (e,f) and LDH 4 (g,h) coatings.

Figure 6. Bode plots for AA1050 alloy with studied LDH coatings.
From the point of view of coating structure, the presence of spheroidal particles in flake-like LDH 1 and LDH 4 (Figure 5) [14] was associated with the presence of secondary phases [35–37] that favored aluminium cation dissolution due to their highly cathodic behavior [48]. In addition, the porous structure of these spheroidal particles also favored Cl\(^-\) anion penetration into the LDH gallery, and for this reason these coatings showed the lowest corrosion protection among the studied LDH coatings (Figure 6) [53–55]. From the point of view of corrosion protection of non-defined LDH coatings, it should be noted that LDH 2 showed similar corrosion behavior in comparison with LDH 4, which can be attributed to its intermediate non-porous LDH structure (Figure 5). On the contrary, LDH 3 provided a beneficial effect to corrosion protection in comparison with all the studied LDH coatings. This may be due to the presence of sodium ions in the LDH gallery, which facilitate the formation of non-porous corrosion-protective LDH coating (Figure 6). For this reason, LDH 3 was the selected treatment to use for the selected PEO coatings and study their anti-corrosion properties (Figure 6).

3.4. PEO-LDH Coating Characterization

Figure 7 depicts the XRD patterns of selected PEO coatings (A3.1, P2.1 and S4) with LDH 3 and discloses the effects of the PEO coating compositions on LDH growth.

![XRD patterns of Zn-Al-LDH-coated A3.1, P2.1 and S4 PEO coatings.](image)

PEO coatings based on aluminate and silicate electrolytes showed the typical peaks at 9.6° and 19.9°, which corresponded to the characteristic (003) and (006) reflections of LDH intercalated with NO\(_3^-\) [32,48,49]. Additionally, the presence of ZnO and Al\(_2\)Si\(_2\)O\(_5\)(OH)\(_4\) characteristic peaks in the S4-LDH XRD pattern came from LDH chemical composition (Table 2) and silicate electrolytes, respectively (Table 1).

In the particular case of PEO coating developed in phosphate electrolyte, there were no peaks detected in that range. This could be attributed to the formation of non-crystalline phases, or to only a small amount that could not be detected at the selected scan rate (Figure 7).

Figure 8 highlights a detailed morphology of the selected PEO-LDH coatings, which reveals the importance of PEO coating composition. The characteristic flake-like LDH structure can be clearly observed for the A3.1-LDH coating, in which LDH flakes are covering the whole PEO coating (Figure 8a,b). This is also observable in the S4-LDH coating, but in this case is more heterogenous (Figure 8e,f). These results are in accordance with XRD patterns that showed the presence of these characteristic reflections (Figure 7). However, for the P2.1-LDH coating there is a drastic decrease of the density of LDH-like flakes (Figure 8c,d), which is in accordance with the XRD patterns (Figure 7).
Consequently, a highly heterogeneous LDH layer was achieved on the microdischarges until the end of the treatment (Figure 2b) resulted in greater thickness and density of the P2.1 coatings. Consequently, a highly heterogeneous LDH layer was achieved in comparison with the S4 PEO coating. This justifies the absence of peaks in XRD patterns of the P2.1-LDH (Figure 7).

This fact is in strong agreement with a high dependence on the availability of Al(OH)$_2^+$ cations necessary to form Zn-Al-LDH. According to previous studies [40,56], Zn-Al LDH synthesis can be explained via the following chemical reactions:

\[
\text{Al}_2\text{O}_3 + 3\text{H}_2\text{O} \rightarrow 2\text{Al(OH)}_3
\]

\[
\text{Al(OH)}_3 + \text{NH}_4^+ \rightarrow \text{Al(OH)}_2^+ + \text{NH}_3\cdot\text{H}_2\text{O}
\]

\[
\text{Zn(OH)} + \text{Al(OH)}_2^+ + 2\text{NO}_3 \rightarrow \text{LDH} - \text{NO}_3
\]

As mentioned before, the in situ growth method was used in the present work and, consequently, Al(OH)$_2^+$ cations were an essential requirement to form the LDH layers. Due to the porous structure of PEO coatings and their compositions, two sources can provide Al(OH)$_2^+$ cations: (i) the aluminium metal matrix (due to the electrochemical interactions with LDH solutions), and PEO coating thickness [57].

Firstly, in consideration of the PEO coating cross-section (Figure 3) and planar view micrographs after LDH treatment (Figure 8), it can be concluded that the amount of LDH flakes on the selected PEO coatings was highest for A3.1, and lowest for the P2.1 and S4 coatings. This could firstly be explained by thickness, and secondly by the chemical composition of the PEO coating surfaces. Due to the low thickness of the A3.1 PEO coating (~1 µm) (Figure 3a,b), the migration capacity of Al(OH)$_2^+$ cations from the aluminium metal matrix towards the coating surface was ensured. However, the P2.1 (Figure 3c,d) and S4 (Figure 3e,f) PEO coatings showed higher thickness values (~1.5–2 µm), and, consequently, the migration capacity of Al(OH)$_2^+$ was reduced. With respect to the chemical composition of PEO coating surfaces, the aluminium content decreased in the order A3.1 > P2.1 > S4 (Figure 3, EDS analysis table) due to the presence of aluminate species in the A3.1 electrolyte. Further, greater charge passed during the P2.1 treatment, and intense sparking from the onset of the microdischarges until the end of the treatment (Figure 2b) resulted in greater thickness and density of the P2.1 coatings. Consequently, a highly heterogeneous LDH layer was achieved.
in comparison with the S4 PEO coating. This justifies the absence of peaks in XRD patterns of the P2.1-LDH (Figure 7).

3.5. Corrosion Resistance of PEO + LDH Coatings

In order to evaluate the effect of LDH formation on selected PEO coatings, corrosion resistance was measured by electrochemical impedance spectroscopy (EIS) for 1 h of immersion in 3.5 wt.% NaCl solution at room temperature (Figure 9).

![Scatter diagram of impedance modulus at 10^-2 Hz; selected PEO-LDH coatings on AA1050 alloy.](image)

As can be seen in Figure 9, PEO coatings with and without LDH treatment showed high similitude of the impedance modulus.

It is important to note that, in this work, the corrosion resistance of different PEO coatings was strongly connected with PEO coating porosity, because the lack of pores restricts the Cl^- penetration and prevents its detrimental effects [58]. For this reason, A3.1 showed the lowest corrosion resistance due to the combination of pores and heterogeneities in comparison with the P2.1 and S4 coatings. The highest corrosion protection exhibited by the S4 PEO coating (∼5 × 10^6 Ω·cm^2) may be attributed to its sparse surface porosity due to the formation of a glassy silica-rich layer.

In this work, the results showed similar behavior of PEO-LDH compared with PEO coatings without any clear improvement after the post-treatment. This could be attributed to several factors, for instance: (i) LDH flake resistance was negligible compared with that of the PEO coating, and (ii) PEO coating lost some of its barrier properties during the formation of LDH, somewhat retracting from the possible small beneficial effect of the LDH layer.

It should be noted that no studies of LDH formation on flash PEO coatings (at <5 min anodizing time) have been reported so far. However, based on this work and few works carried out with non-flash PEO of aluminium, some conclusions can be drawn regarding the effects of LDH post-treatments. For instance, when an LDH layer is not loaded with inhibitors, the corrosion resistance remains unchanged or degrades slightly, as has been shown in [37]. On the other hand, when LDH is intercalated with an inhibitor (e.g., vanadate ions), an improvement in corrosion resistance is observed with immersion time due to an active protection effect [37,38].

The present findings highlight that the development of LDH-container layers did not deteriorate the corrosion resistance of flash PEO coatings, which has a potential for added active protection functionality. Therefore, the first stage of active protection system building can be considered
successful. The second stage would consist of loading the LDH scaffold with corrosion inhibitors that would ensure enhanced corrosion protection.

In conclusion, these results are highly relevant for understanding the relation between coherent and uniform LDH layer formations on flash PEO coatings, which is the first step in achieving active protection systems through the incorporation of green corrosion inhibitors into the LDH layer.

4. Conclusions

The following can be summarized from this preliminary study of LDH growth on flash PEO coatings:

- Flash PEO coatings with ~1–2 µm thickness and ~2–5 KW·h·m⁻²·µm⁻¹ energy consumption were generated on a commercially pure aluminum alloy. Low energy consumption was ensured through relatively high electrolyte conductivity and a transition of the anodizing regime from constant current to constant voltage control.
- The first stage of the active protection system was successfully completed on flash PEO coatings via the development of an LDH layer. LDH coating is continuous and well defined when the PEO layer is thin (~1 µm), and the LDH formation is further facilitated when additional Al(OH)₂⁺ cations are lixiviated from the coating.
- Corrosion resistance of inhibitor-free flash PEO/LDH coatings is mainly determined by the low porosity of the PEO layer. Formation of the LDH layer does not compromise the corrosion resistance of flash PEO coatings. Loading of the LDH scaffold with corrosion inhibitors is necessary in order to achieve an enhanced corrosion protection.

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