The repassivation response from single cycle anodic polarization: The case study of a sensitized Al-Mg alloy

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The repassivation behaviour of artificially aged 5083-H111 Al-Mg alloy in near neutral NaCl solutions was investigated by means of single cycle anodic polarization. Artificial aging was carried out at 150 °C during different times up to 360 h. The characterization of the microstructure and composition by XRD, SEM/EDS and metallographic analyses indicated most favoured grain-boundary β(Al13Mg5) phase precipitation for 168 h of isothermal treatment time. Nitric acid mass loss test (NAMLT) and microhardness measurements indicated the highest degree of sensitization (DoS) and developed strength for 168 h as well. For all the aging conditions, the potential drop at high currents during the reverse scan of the cyclic polarization was detected at the pit transition potential $E_{\text{pp}}$ with similar but higher values than the corrosion potential of β phase ($E_{\text{corr}} \approx -900$ mV vs SCE). The associated current density $i_{\text{pp}}$ and the steepness of the potential drop increase as the amount and contiguity of β phase precipitates along grain boundaries. The onset and sustenance of metastable conditions that limit the simultaneous repassivation of all the corroded surfaces are driven by the anodic dissolution of β(Al13Mg5). Metastable processes that limit the hydration of metal ions while producing high local hydrogen concentration are more likely to involve the formation/decomposition of reactive hydride intermediates. $E_{\text{pp}}$ corresponds to the mixed electrode potential at which both metal dissolution and hydrogen evolution occur beyond some distance into the cavity, while $i_{\text{pp}}$ evaluates the cathodic activity of the corroding surfaces. The effective anodic charge transfer coefficient $\alpha_{\text{ad}}$ determined from the steepness of the potential drop estimates the contribution of $\text{Cl}^-$ electromigration in response to local electrodeposition processes.

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1. Introduction

Single cycle anodic polarization has been widely used for investigating the repassivation behaviour of Al and alloys in chloride containing solutions [1–17]. A potential drop at high currents during the potential scan into the negative direction has been frequently detected [1–4,6–10,12–17]. This characteristic, illustrated for an Al-Mg-Si alloy in Fig. 1 [10], manifests a transition to more occluded local processes that limit the simultaneous repassivation of all the corroded surfaces [1,8]. The transition supported by the critical pit chemistry has been recognized [1–10]. The potential at the inflection (Fig. 1), designated as the pit transition potential $E_{\text{pp}}$ according to Yasuda et al. [2], corresponds to the thermodynamic driving force of Al dissolution on freshly created (filmed) surface rather than on initially passive surface as for the pitting potential $E_{\text{p}}$ [3,10]. The associated current density $i_{\text{pp}}$ is proportional to the rate at which hydrolysis equilibrium is reached at a critical saturation concentration of Al$^{3+}$ [10], being represented by the following general equation to account for the contribution of OH$^-$ (pH > 7):

$$2\text{Al}^{3+} + \text{H}_2\text{O} + \text{OH}^- \leftrightarrow 2\text{Al(OH)}^{2+} + \text{H}^+ \quad (1)$$

Both $E_{\text{pp}}$ and $i_{\text{pp}}$ define the onset of metastable conditions that hinder the repassivation processes. The steepness or gradient of the potential decrease with current density at $E < E_{\text{pp}}$ (Fig. 1) is proportional to the local acidity removal for full hydrolysis to be reached at the protection potential $E_{\text{prot}}$ [10].

The response to repassivation in aftermath of microstructural effects (e.g. impurities, precipitate phase, solute-depleted region) [6,10,12] and of mechanical deformation (residual [13] and applied [9,16,17] stresses) is not fully clear. The not distinguishable inflection detected for commercially pure Al 1050 exhibiting either etched or crystallographic pits [10] suggests that the presence of a salt layer in mass-transport limited repassivation is not mandatory. From studies with alloyed aluminium [6,10], $E_{\text{pp}}$ does not change if the critical feature is the electrochemical behaviour of a precipitate phase, as for Al 6082-T6 which intergranular corrosion is governed by the selective dissolution of MgSi [10]. Conversely, $E_{\text{pp}}$ varies if the critical feature is a solute depleted region as in the case of Cu-containing alloys [6,10]. The shift of $E_{\text{pp}}$ to less negative values with the amount of corrosion [10] is consistent with the selective dissolution of Cu-rich particles producing less active Cu-depleted zones between the

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trolling processes of IGC and IGSC propagation along the grain boundary between the β phase particles. The anodic dissolution of the β phase aiding the sustenance of the acidified pit-like chemistry for grain boundary diffusion of hydrogen and embrittlement has been considered in the IGSC mechanism of Al-Mg alloys [21–23].

In this paper, the effect of [β(Al,Mg)] precipitation at grain boundaries on the repassivation behaviour of a high content of Mg (≥ 4 wt%) 5083-H111 Al alloy in 0.6 M NaCl (pH 6.5) is investigated by means of single cycle anodic polarization. The alloy specimens were isothermally treated at 150 °C in air during different times and characterized by XRD, SEM/EDS and metallographic analyses. The DoS and the mechanical properties were evaluated by conventional NAMLT and microhardness measurements.

2. Experimental part

The starting material was a commercial wrought sheet (thickness 1.5 mm) of 5083-H111 Al alloy with nominal composition (wt %) 4.3 Mg–0.6 Mn – 0.3 Fe – 0.2 Si - bal. Al (Aaviometal Spa., Italy), present in the laboratory since 2006 (aged ≈ 10 years aged at room temperature). Metallic specimens of dimensions 20 × 30 mm were wet ground up to 1200 grit using SiC papers, then polished with 4000 grit (3 μm) 3 M polishing paper, and cleaned with ethanol in an ultrasonic bath. Thermal treatment was carried out at 150 °C in an open to air oven during different times (from 24 to 360 h). The composition and microstructure of the untreated and isothermally treated specimens were evaluated by XRD, SEM/EDS and metallographic analyses. The diffractograms were collected using Philips PW 1830 diffractometer with CuKα radiation from 5 to 80° of 2θ. XRD data were analyzed with Diffrac plus EVA software (Master database, Bruker). SEM/EDS analysis of freshly-prepared specimens with surfaces polished up to 0.05 μm grain size of colloidal alumina (water-based dispersion) was performed at LEO 1430 scanning electron microscope (SEM) equipped with EDS spectrometer at the chamber pressure of 8 × 10^-6 torr and 20 keV accelerating voltage. Another set of specimens was used for metallographic analysis, following a reported procedure [24]. Briefly, the surfaces were chemically etched by immersion in (NH₄)₂S₂O₇ (10 g/100 mL) at room temperature during 30 min and thereafter examined at Nicon Eclipse MA200 metallographic microscope coupled with JVC-C1380 digital photocamera. Nitric acid mass loss test (NAMLT) [19] was typically used for evaluating the susceptibility to IGC in terms of the degree of sensitization (DoS) in mg/cm² of mass loss, being the traditional ranges of IGC susceptibility defined as: resistant (DoS < 15 mg/cm²), intermediate (15 mg/cm² < DoS < 25 mg/cm²) and susceptible (DoS > 25 mg/cm²). Because of the faster dissolution than the matrix of the β phase in concentrated nitric acid, the mass loss increases due to grain fall-out with the amount and connectivity of β phase at grain boundaries. The interdependence between the exposure conditions (temperature, time and environment) and interrelated variables such as metallurgical, electrochemical and stress states, have been underlined in recent reviews reporting most of the relevant works on this argument and new experimental findings [20–23]. The critical temperature range 100–200 °C (DoS > 25 mg/cm²) is determined by the appreciable diffusion kinetics and volume fraction of β-phase [20]. However, long-time exposure at ambient temperatures promotes sensitization due to the high equilibrium β-phase volume fraction. Nucleation and growth of β phase at other lattice defects such as at the interface between the matrix and intragranular second-phase particles and at dislocations may occur [20,21]. The apparent β-phase continuity parameter, defined as the ratio between the average nearest neighbour distance (NDD) and the average diameter (D) of β-phase at grain boundaries, has been indicated as more adequate than DoS for predicting IGSC susceptibility [21]. A key aspect concerns the rate-con
solution. The potential scan was reversed at the preselected current density limit of 2.5 mA/cm² (I_{rev}, Fig. 1) and terminated when the current became cathodic. The polarization curves were replicated at least twice for a given treatment condition. Representative specimens were cleaned with MilliQ water and ethanol in an ultrasonic bath, then sectioned, mounted in a cold-working resin, and finally polished up to 0.05 μm grain size of colloidal alumina (water-based dispersion), for cross-section corrosion morphology analysis at SEM.

Graphical and quantitative analyses of the experimental data were carried out using OriginPro 2016 (OriginLab, Northampton, MA), as reported elsewhere [10].

### 3. Results and discussion

XRD, SEM/EDS and metallographic analyses of thermally treated Al 5083-H1111 specimens indicated good agreement with reported microstructural changes of Al-Mg alloys due to sensitization [20]. The main secondary phases identified by XRD were Mg₂Si, Al₆(Fe,Mn) and β(Al₃Mg₂) (supplementary data, Fig. S1). The number of diffraction lines for the latter two phases was higher for 168 h of isothermal treatment at 150 °C. For this treatment time, larger Al₆(Fe,Mn) particles and less discontinuous precipitation of β phase along grain boundaries were observed on the etched surfaces at the metallographic microscope (Fig. 2). Similarly, SEM/EDS analysis (Fig. 3) indicated more significant nucleation and growth of β phase at the interface between Al matrix and either intragranular Al₆(Fe,Mn) or Mg₂Si particles. Accordingly, the isothermal treatment time of 168 h favours most the precipitation of β(Al₃Mg₂) phase along grain boundaries and at other lattice defects. The dimensions of Al₆(Fe,Mn) and Mg₂Si intermetallics decreased with prolonged thermal treatment (Fig. S2). In addition, the morphology of Mg₂Si particles changed from roughly spherical to elongated shapes (Fig. 3b,d). Because of the reduced solubility of silicon in high Mg content Al alloys [25], unbalanced Mg₂Si due to excess of Si was detected for all the treatment conditions. Long-time isothermal treatment is likely to bring about elongated grains of smaller size (Fig. 2), being not so well discerned due to the lessening of grain-boundary beta phase precipitates (Fig. S1). The variation of the degree of sensitization (DoS) in terms of mass loss with the treatment time (Fig. 4) reproduces rather well documented data for as-solutionized Al 5083 (see Fig. 4 in Ref. [20]). The closely equal DoS values for the untreated alloy (0 h) and that reported for as-received Al 5083 H (=9 mg/cm²) [23] suggest no significant effect of long-time aging under laboratory air environment on the susceptibility to IGC. The higher DoS for treatment times between 100 and 200 h (Fig. 4) correlates with the less discontinuous β phase precipitation along grain boundaries (Fig. 2c). The mechanical properties of the alloy changed with the thermal treatment time also. The higher microhardness for sensitization times between 24 and 168 h (Fig. 5) suggests significant precipitation hardening [26]. By considering the DoS results (Fig. 4), the uniform distribution of fine β phase particles in the case of 24 h of thermal treatment is likely to account for the strength development. The concomitant development of internal strains in the matrix [26] is expected to be significant for 100–200 h at 150 °C because of β phase precipitation at other than grain-boundary lattice defects (Fig. 3). For long-time exposure at 150 °C (≥240 h), the decrease of both microhardness and DoS indicates reversion (or remediation) of sensitization at the expense of a loss in strength [20,23], being consistent with the β phase dissolution in the solid matrix (Figs. S1 and 2). The further relief of internal

![Image](image_url)

**Fig. 2.** Surface images at the optical microscope of etched Al 5083-H1111 after aging at 150 °C during (a) 0 h; (b) 24 h; (c) 168 h; (d) 360 h. The aging time in (a) corresponds the untreated alloy (10-years aged at room temperature under laboratory conditions). Grain boundaries decorated with black [(Al₃Mg₂) precipitates, gray particles correspond to Al₆(Fe,Mn).
Fig. 3. Surface SEM/BSE images of as-polished Al 5083-H111 after isothermal treatment at 150 °C during (a,b) 24 h; (c) 168 h; (d) 360 h.

Fig. 4. Mass loss as a function of time at 150 °C for Al 5083-H111. The time of 0 h corresponds to the untreated alloy. Error bars represent the standard deviation of the mean mass loss values. The dashed line correspond to Dos = 25 mg/cm².

Fig. 5. Micro-hardness as a function of time at 150 °C for Al 5083-H111. The time of 0 h corresponds to the untreated alloy. Error bars represent the standard deviation.

The response to the positive potential scan is featured by a significant increase of the current density at similar pitting potentials ($E_{pp} = −740 \pm 10 \text{ mV vs SCE}$), regardless the extent of β phase precipitation (Figs. 2 and 3). Likewise, the inflection during the potential scan into the negative direction is detected at closely equal pit transition potentials ($E_{ppt} = −775 \pm 5 \text{ mV vs SCE}$). The corrosion potential of the β phase in 0.6 M NaCl is more negative ($E_{corr} ≈ −900 \text{ mV vs SCE}$) [21]). This phase can sustain high dissolution currents between pH 2–10 [27], while Al(OH)₃ is stable between pH 4–9 and Mg(OH)₂ at alkaline pH only. Accordingly, metastable processes that hinder the simultaneous repassivation of all the corroded surfaces are
driven by the electrochemical activity of the β phase. This fact supports further the close relation between $E_{\text{pp}}$ and $E_{\text{pot}}$ [3], being the former potential less reproducible as influenced by the initial surface state condition [10]. The smaller by about 40 mV $E_{\text{pp}}$ reproduces previous results with the untreated alloy [10] and is consistent with the thinner layer of the non-protective as-filmed surface during the first stages of repassivation.

Regardless the above, only the reverse curves change with the isothermal treatment time as the amount and contiguity of β phase precipitation along grain boundaries. That is, by increasing the isothermal treatment time up to 168 h, the inflection at $E_{\text{pp}}$ shifts to higher current densities $i_{\text{pp}}$ and the gradient or steepness of the potential drop increases with respect to the untreated alloy (Fig. 6a). The opposite trend is obtained for 240 and 360 h. Thus, the protection potential $E_{\text{pot}}$ changes non-monotonically between −850 and −900 mV (vs SCE), obtaining the more negative values for 100 and 200 h at 150 °C (Fig. 6). The variation of the steepness and of $E_{\text{pot}}$ with the thermal treatment time are better discerned from the corresponding reverse curves plotted as $E$ vs $\ln (i_{\text{pp}})$ (Fig. 6b). The less negative ill-defined inflection at about −0.74 V for 240 and 360 h (Fig. 6b) could be ascribed to the repassivation of superficial IG corroded sites due to surface spreading of β phase attack [21]. The $E_{\text{pot}}$ values are closely similar to the $E_{\text{pot}}$ of $\beta$(Al,Mg) phase in Al-Mg alloys between −850 and −900 mV vs SCE for sensitized Al 5083-H131 in 0.1 M NaCl (pH 8) [21], as-received Al 5083-H131 in 0.01 M NaCl (pH 4–8) [27], and sensitized Al-5 wt.% Mg microelectrodes in deaerated 0.6 M NaCl (pH 3.5) [28]. Accordingly, metastable processes during repassivation are driven and sustained by the anodic dissolution of the β phase.

The empirical relationships of $i_{\text{pp}}$ and of the steepness as a function of time at 150 °C are reported in Fig. 7a and b. The values of $i_{\text{pp}}$ and of the steepness do not significantly differ for long-time aging at room temperature and for 24 h at 150 °C, in spite of the different microstructure (Fig. 2), DoS (Fig. 4) and microhardness (Fig. 5). For isothermal treatment times between 100 and 200 h, higher $i_{\text{pp}}$ and steepness correlate with higher DoS (≈50 mg/cm²) and microhardness. For treatment conditions that favour the reversion of the sensitization (≥240 h at 150 °C), the DoS differs little from that of the untreated alloy (Fig. 4) but $i_{\text{pp}}$ values are much smaller (Fig. 7a) while the decrease of the steepness follows closely that of the microhardness (Figs. 7b and 5). These results indicate that matrix dissolution nearby grain boundaries is as important as the extent and contiguity of β phase at grain boundaries, in support to IG/IGSC propagation between discrete β phase particles as proposed by others [20,21] and refs. therein. The localization of anodic processes during repassivation is enhanced with β phase precipitation at lattice defects [17,20] and refs. therein. SEM analysis results of cross-sectioned corroded specimens strengthens the considerations above. Prevalent IG attack at the base of crevice like pits was observed for long time aging at room temperature and for 24 h at 150 °C (Fig. 8a). IGSC morphology manifested by IG attack over multiple grains was evident for 168 h (Fig. 8b). Exfoliation corrosion was the prevalent type of damage for prolonged artificial aging (≥240 h) (Fig. 8c), being ascribed to the more directional microstructure (Fig. 2d) [21].

![Fig. 6. (a) Average cyclic anodic polarization curves of Al 5083-H111 calculated from replicated curves in 0.6 M NaCl (pH 6.5). Error bars for 40 points of averaged E for a given log t are included. (b) $E - \ln (i_{\text{pp}})$ plots of the average reverse curves in (a). Numbers near the traces in (a,b) indicate the sensitization time (hours) at 150 °C. The time of 0 h corresponds to the untreated alloy.](image1)

![Fig. 7. Variation of the kinetic properties of repassivation with time at 150 °C: (a) mean $i_{\text{pp}}$, (b) mean steepness. The time of 0 h corresponds to the untreated alloy. Error bars represent the standard deviation.](image2)
Summing up present findings, the susceptibility to IGSC due to sensitization can be evaluated from the repassivation response in near neutral NaCl solutions. Metastable processes that hinder the growth of protective corrosion products are driven and sustained by the anodic dissolution of β(Al₃Mg₂) phase, being in accordance with its controlling role in the IGC and IGSC of Al-Mg alloys [20–23]. The $E_{\text{pp}} > E_{\text{re}} \approx E_{\text{cet}}$ of β phase supports the prevalence of a simultaneous dissolution of both Mg and Al [29]. The resulting Mg²⁺ and Al³⁺ chloride complexes should promptly interact with water molecules to form densely hydrated oxychloride products [30,31]. The growth of a protective (hydr)oxide film should readily overcome that of the halide salt layer with decreasing potentials. Similarly, $\text{H}^+$ reduction at (hydr)oxide covered walls should assist the dilution of the local aggressive environment. The concentration gradients between the occluded cavity and the outer bulk solutions should easily decrease if the geometrical constriction of localized attack is not significant. This is likely the case of 168 h of isothermal treatment (Fig. 8b), being consistent with the highest $i_{\text{pp}}$ (Fig. 7a) according to the reaction (1) and the highest DoS (Fig. 4), but not with the steeper potential drop at high currents (Figs. 6 and 7b). Accordingly, the anodic dissolution of β phase drives and sustains electrodissolution processes nearby grain boundaries through a reaction pathway that limits $\text{H}_2\text{O}/\text{OH}^-$ adsorption and $\text{H}^+$ removal during repassivation. A self-driving process could be the anodic dissolution of the β phase through hydrides intermediates that can actively deprotonate and thus be a source of hydrogen [28,29]. $E_{\text{pp}}$ can be considered as the mixed electrode potential at which both metal dissolution and hydrogen evolution occur beyond some distance into the cavity, while $i_{\text{pp}}$ reflects the catalytic activity of the corroding surfaces.

With the intention to supply a physical meaning to the steepness, high currents driving a potential drop during repassivation are related to mass transport processes involving metal electrodissolution and electromigration of $\text{Cl}^-$ through the highly conductive salt layer. The decrease of the potential below $E_{\text{pp}}$ corresponds to the variation of the electrostatic potential at some distance into the cavities, being driven by a flow of charge across the metal/solution interface proportional to $i \leq i_{\text{pp}}$. Assuming further that, for a stagnant electrolyte, the processes above depend only on the potential and their rate depends only on the concentrations of $\text{Al}^{3+}$ and $\text{Cl}^-$, the model developed by Engelhardt et al. [32] for calculating corrosion cavity propagation rates was used to estimate the effective anodic charge transfer coefficient $a_{\text{eff}}$ from the steepness (Fig. 4b) with Equation (2):

$$\ln \left( \frac{i}{i_{\text{pp}}} \right) = \frac{a_{\text{eff}} F}{RT} \left( E - E_{\text{pp}} \right)$$

where $E$ and $E_{\text{pp}}$ are expressed in V (vs SCE), $F = 9.65 \times 10^4$ J/Vmol, $R = 8.31$ J/molK, $T = 298$ K. Considering that $a_{\text{eff}} = a - \lambda$ [32], where $a$ is the anodic charge transfer coefficient and $\lambda$ the effective kinetic order of metal dissolution with respect to $[\text{Cl}^-]$, $a_{\text{eff}}$ allows to estimate the contribution of $\text{Cl}^-$ electromigration in response to local dissolution processes.

For all the investigated conditions, $a_{\text{eff}} > 0$ is consistent with the decrease of the dissolution rate as the potential shifts to more negative values (Fig. 6b). However, $a_{\text{eff}}$ decreases from 0.35 to 0.17 with artificial aging up to 168 h and then increases to 0.54 for 360 h. Accordingly, $a_{\text{eff}} \rightarrow 0$ and $a \rightarrow \lambda$ with β phase precipitation indicates enhanced electromigration of $\text{Cl}^-$ for compensation of the potential drop while the dissolution rate tends to change less with the cavity length. The potential drop at high currents during repassivation is, all in all, a
manifestation of the limited growth of the (hydr)oxide film over that of the salt layer.

Although it is not possible with the present data to propose a definite mechanism by which the aggressive local environment is sustained, some observations regarding the β phase acting as local anode and as local source of hydrogen are disclosed. The generation of low pH is not sustained by the hydrolysis of non-acidic Mg²⁺ (pKa = 11.4) \[31\]. Complexes of MgH and MgH₂ accounting for hydrogen embrittlement have been claimed because of the increased affinity between Mg and H in aluminium \[21\]. However, these hydrides would prevent further hydrogen diffusion and may readily hydrolyse as well \[29,33,34\]. Taking into account the predilection of Mg surface to reactively dissociate water and the low barrier of hydrogen atoms recombination on a defective MgO surface \[31,35\], as well as our recent experimental findings on hydrogen evolution during the anodic dissolution of Mg-Al alloys \[36\], oxodic hydrides could form according to the reaction (2):

\[
\text{Mg} + \text{H}_2\text{O} \rightarrow \{\text{Mg}^2\text{H}^- + \text{O}^2--\text{H}^+\} \rightarrow \text{MgO} + \text{H}_2 \uparrow
\]

The “solvated” hydrides may react with AlCl₃ to form metastable complex H-bridged networks of magnesium aluminiumhydrides (or alanates) of unit formula Mg(AlH₄)₂ \[37,38\]. Although this awaits verification through accurate studies, experimental evidence of magnesium alanate intermediates has been provided for Mg-Al alloys \[36\]. In addition, Mg(AlH₄)₂ is less stable at ambient temperatures, exhibits higher gravimetric hydrogen density and presents a more open structure for hydrogen mobility in comparison to MgH₂ \[38\]. Furthermore, the decomposition of Mg(AlH₄)₂ gives a stable mixture of MgH₂ and Al, alongside with H₂ evolution, being the former sub-products experimentally detected during corrosion of Al₃Mg₂ \[29\]. The anodic dissolution of β phase through reactive hydride intermediates like magnesium alanate would support the limited hydration of both Al³⁺ and Mg²⁺, the critical saturation concentration of Al³⁺ for production of fresh reactive surfaces nearby grain boundaries, and the build-up of high local concentration of hydrogen. The local generation of diffusible hydrogen species would assist the embrittlement and cracking between discrete β phase precipitates at grain boundaries.

4. Conclusions

The effect of the isothermal sensitization at 150 °C on the susceptibility to IGSC and IGSC of Mg-rich 5083-H111 Al alloy in near neutral NaCl is discerned from the repassivation response of single cycle anodic polarization. The analysis of the electrochemical potentials provided confirmation of the electrochemical behaviour of β(Al-Mg₃) phase as the critical feature in IG/IGSC phenomena of Al-Mg alloys. The onset and sustenance of metastable conditions that hinder the simultaneous repassivation of all the corroded surfaces are enhanced with the amount and connectivity of β phase at grain boundaries, in correlation with the increase of IG/IGSC susceptibility in terms of DoS (NAMLT). The anodic dissolution of β phase drives and sustains matrix dissolution nearby and along grain boundaries. The most likely metastable processes are related to the formation/decomposition of hydride intermediates that limit the growth of protective corrosion products while supply high local hydrogen concentration. E₄₆₅ is proposed as the mixed electrode potential at which both metal dissolution and hydrogen evolution occur beyond some distance into the cavity. The catalytic activity of the corroding surfaces is evaluated from i₉₄₅. The potential drop at high currents is related to Cl⁻ electromigration in response to local electrode dissolution processes. The contribution of anodic processes localization during repassivation can be estimated from the steepness through the effective anodic charge transfer coefficient αₑᵣ.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at https://doi.org/10.1016/j.electacta.2017.10.202.

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