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Influence of hydrogen on the propagation of intergranular corrosion defects in 2024 aluminium alloy

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ABSTRACT

To study the influence of hydrogen on the intergranular corrosion mechanism of a 2024 aluminium alloy, samples were hydrogen precharged by cathodic polarisation and then exposed to a NaCl solution. EBSD analyses and SEM observations showed that hydrogen increased the number of corroded interfaces and led to the embrittlement of low-angle grain boundaries which were not susceptible to corrosion without hydrogen precharging. The increase of the reactivity of the 2024 aluminium alloy in the presence of hydrogen gave a new insight into the intergranular corrosion mechanism: corrosion-induced hydrogen promoted the intergranular corrosion propagation and partially controlled the corrosion defect morphology.

1. Introduction

2024 Al alloy (AA 2024) is one of the most widely used Al alloys in aerospace industries. It is a structural-hardening alloy with Cu and Mg as major alloying elements. The formation of hardening precipitates (Guinier-Preston (GP) zones or θ-Al2Cu/S’-Al2CuMg phases depending on the metallurgical state) leads to high mechanical properties (yield strength and ultimate tensile strength) [1,2], but it is combined with a more largely heterogeneous microstructure, characterised also by the presence of intergranular precipitates, dispersoids and intermetallic coarse particles (IMC), that increases the susceptibility of the alloy to localised corrosion, such as intergranular corrosion (IGC) [3–5]. Numerous papers in the literature show that pits initiated preferentially on IMCs in AA 2024 [6–8]; concerning IGC, the mechanism is usually described as galvanic coupling between intergranular Cu-rich particles (θ-Al2Cu; S-Al2CuMg) and the adjacent matrix [9–12].

Nevertheless, even if the precipitation state is a first order parameter for IGC, the AA 2024 susceptibility towards IGC depends on other parameters as, for example the level of misorientation (θ) of the grain boundaries. The influence of this parameter on corrosion phenomena has been studied on various materials, such as stainless steels, Ni base alloys and Al alloys [13–16]. For AA 5182, Davenport et al. [17] have reported that high-angle grain boundaries (HAGB, θ > 20°) are susceptible to corrosion mechanisms, such as IGC due to the presence of β-Mg2Al3-phase precipitates. In another study performed on AA 2050, Guérin et al. have shown that, even if most of the interfaces in the alloy are low-angle grain boundaries (LAGB, θ < 10°), only HAGB (θ > 10°) are corroded after exposure to an aggressive NaCl solution [18]. Other authors, such as Luo et al. have explained the susceptibility to intergranular corrosion of AA 2024 T351 taking the grain-stored energy into account [19]. As grain-stored energy reflects the dislocation density, they proposed that a high level of dislocation density within individual grains made them more susceptible to preferential attack at the grain boundary.

However, the corrosion processes associated with the galvanic coupling between intergranular Cu-rich particles (θ-Al2Cu, S-Al2CuMg) and the adjacent matrix lead to the production of Al cations; the hydrolysis of Al cations generates protons. This induces an acidification of the electrolyte trapped inside the IGC defects. Further, previous studies showed the formation of a thin metallic Cu-nanoparticles layer at the interface alloy-IGC defect [20], that could be due to dissolution of the Cu-rich intergranular precipitates [19,21], and/or to the matrix dissolution [22]. This Cu-rich layer could protect the interior of the attacked grains from further corrosive environment in AA 2024 [19,23,24]. However, Cu could also provide effective cathodic support for oxygen and/or protons reduction on the walls of the IGC defects [24]. Bonfils et al. showed hydrogen (H) penetration inside the matrix during the corrosion processes with H amount reaching 20 ppm and even 120 ppm after 30 and 500 h of exposure to a 1 M NaCl solution, respectively [25]. Similarly, Larignon et al. have shown that H can be detected at HAGB and LAGB in corroded AA 2024 samples by scanning kelvin probe force microscopy (SKPFM) [26]. Lafouresse et al. have also
used SKPFM analyses on corroded AA 2024 in order to study H desorption kinetics from corrosion defects and have distinguished between H trapped at different locations [27]. Therefore, due to the large amount of H trapped inside the corroded materials, questions are being raised about its subsequent influence on the corrosion processes. Laignon et al. have clearly shown, for cathodically H-precharged AA 2024 samples, that H induced a shift towards more negative values of the corrosion potential [26]. Further, Lafouresse et al. have performed alternative current scanning electrochemical microscope (AC-SECM) measurements on H-precharged AA 2024 samples and have shown an increase of the local reactivity, i.e. higher local currents associated with a lower effective resistance, due to the H diffusion inside the samples [28].

Similarly, the influence of H on the oxygen reduction reaction in a martensitic stainless steel has been shown by Schaller et al. [29,30]. For other materials and especially for Ni base alloys, the influence of H localised at the grain boundaries on the propagation of intergranular cracks has been investigated. Barrows et al. have used molecular dynamics simulations for Ni base alloys to highlight the influence of H localised at the grain boundaries on the intergranular decohesion and also showed that the increase of the H coverage of the Σ3 grain boundaries influenced the crack tip propagation kinetics [31]. In another study concerning Ni3Al in a H2-containing environment, Takanashi et al. have observed brittle fracture nucleation for randomly oriented grain boundaries, except for LAGB [32]. On the contrary, for coherent Σ3 boundaries, no grain-boundary fracture was observed. The H diffusion along the grain boundaries has also been studied by Feaugas et al. in Ni base alloys [33,34]; they showed a fast diffusion and a high solubility of H along high-angle random boundaries. These authors also showed that H was trapped at special Σ3 grain boundaries, which might reduce the susceptibility of the metals to H embrittlement due to the low H diffusion [35]. The results have been confirmed by Bechtle et al. [36].

However, to the author knowledge, except for Lafouresse et al. [28] and Schaller et al. [29,30], there are very few publications that provide data concerning the influence of H on the electrochemical reactivity of the materials. However, if H is introduced around a corrosion defect and if it can modify the electrochemical reactivity of the material, then it could have an influence on the corrosion defect growth and then on its morphology. Therefore, to provide a better understanding of the IGC mechanism and the propagation kinetics of the IGC defects in AA 2024, the present work aims to study the influence of H on the propagation of the IGC defects. H-precharged samples were studied after exposure to a corrosive environment (5 mM NaCl solution) and the results were compared with those obtained for non-charged samples. Firstly, optical and transmission electron microscopic (OM and TEM) observations were used to characterise both the morphology and chemical composition of the IGC defects accurately. Then, electron backscatter diffraction (EBSD) analyses were performed to identify the corroded interfaces accurately for both non-charged and H-precharged samples.

2. Experimental procedure

2.1. Material

The material used for this study was a AA 2024 T351 provided by Airbus Group (France). The material was received as a 50 mm thick plate obtained by hot rolling; this step was followed by solution heat treatment at 495 °C (±5 °C), water quenching, straining and tempering in ambient conditions for 4 days to achieve the final T351 metallurgical state. The chemical composition of the material was the following: Al base, 4.46% Cu, 1.44% Mg, 0.60% Mn and 0.13% Fe, wt. %. The microstructure of this alloy corresponded to grains highly elongated in the rolling direction (longitudinal, L) with the average sizes of 700, 300 and 100 μm in the longitudinal (L), long transverse (LT) and short transverse (ST) directions, respectively.

2.2. H precharging

Cathodic charging in 10 mM H2SO4 solution was selected to introduce H inside the material since a large H amount can be inserted into pure Al with this method [37]. A Pt counter electrode and a saturated calomel reference electrode (SCE) were used. The samples were 1 cm³ cubes, the precharging duration was 40 h and the precharging potential was ~ 0.8 V vs SCE; these parameters allowed a sufficient amount of H to be introduced inside the sample in a reasonable amount of time. The precharging side was polished with diamond paste down to 3 μm with distilled water as a lubricant before precharging; it was chosen to be perpendicular to the elongated grain direction (L) [27,28]. Therefore, one of the long transverse (LT) - short transverse (ST) sides of the samples was exposed to the electrolyte; all the other sides were protected by a varnish. After cathodic precharging, the H amount was determined by gas fusion analysis using a Bruker G8 GALILEO ON/H Instrumental Gas Analyser (IGA); the H content was 19 ppm in agreement with previous results [27].

2.3. Corrosion tests

Corrosion tests were performed for both non-charged and H-precharged samples to make IGC defects grow along the L direction. The tests corresponded to continuous immersion of the samples (1 cm³ cubes) in a 5 mM NaCl aerated solution at room temperature (25 °C). A low chloride concentration was chosen because, as previously said, corrosion phenomena lead to H production. To be able to study the influence of H on the corrosion behaviour of AA 2024, it was essential that the H amount generated by the corrosion processes remained lower than the H introduced by cathodic precharging for H-precharged samples. The duration of the corrosion tests was also adapted to allow the influence of H introduced by cathodic precharging to be revealed. Some corrosion tests were performed in a more concentrated NaCl solution (1 M) and for long duration (24 h) for two non-charged samples to study the evolution of the IGC defect morphology. Only the LT-ST faces of the samples were exposed to the NaCl solution, the other faces being protected by a varnish [38,39]. Before exposure to the NaCl solution, the non-charged samples were abraded using SiC papers down to 5 μm; then, they were polished down to 3 μm using diamond paste with distilled water as a lubricant. For the H-precharged samples, only a very short polishing step with 3 μm diamond paste was done since they have been abraded then polished before the H-precharging. Unless it was short, this polishing step could lead to some H desorption. However, it corresponded to a perfectly defined protocol used for all samples so that it was assumed that this did not lead to misinterpretation of the results.

2.4. Characterisation of the IGC defects: preparation of the samples and description of the techniques

The evolution of the IGC defect morphology was first studied by using a MA200 NIKON OM. After the corrosion tests (6 h in a 5 mM NaCl solution), cross-sections (L-ST plane) of the corroded samples were prepared and observed to characterise the morphology of the corrosion defects for both non-charged and H-precharged samples. The results were obtained from the observations of 5 samples for each case (non-charged and H-precharged samples), which corresponded to 5 cm analysed in the ST direction for each case. In the following, mean values were given. For a more accurate description of the corrosion defects, transmission electron microscopic (TEM) observations were performed using a JEOl JEM2100 F equipped with a field-emission gun and an energy-dispersive X-ray spectrometer (EDX, SDD Bruker) operating at 200 kV at UMS Caistina. In order to obtain a thin sample in a localised region, i.e. one containing an IGC defect, the preparation was done using conventional lift off procedure using a FEI HELIOS Nanolab 600i dual beam FIB / Scanning Electron Microscope (SEM) [20]. Three corroded H-precharged samples were observed and representative
observations are shown in the following. The results were compared to those described in previous work [20] for corroded non-charged samples. EDX analyses were helpful in identifying the precipitates at the grain boundaries.

2.5. EBSD analysis

EBSD analyses were performed on three samples before H precharging and corrosion exposure using a JEOL JSM 7100 F TTIS LV field-emission SEM and an OXFORD INSTRUMENTS HKL EBSD system (Norlyx Nano EBSD detector - version 3.1 of the Aztec software). Samples were analysed at 20 kV with a 12.5 mA current. EBSD maps were performed on a 1.5 mm x 3 mm area delimited by micro-hardness indents with a step size of 1.5 µm in the LT-ST plane. This selected area was large enough to obtain representative results. Overall, approximately 1000 grains were analysed for each sample. The commercial orientation imaging software package Oxford Channel 5 was then used for the post-processing of the EBSD data. Prior to data analysis, in order to minimise measurement errors, all grains comprising less than 9 pixels were automatically removed from the maps. Further, a lower limit boundary misorientation cut-off of 3° was used to eliminate spurious boundaries caused by orientation noise. The analyses allowed the determination of the nature of the interfaces (Fig. 1a) in relation with their level of misorientation in agreement with Priester who defines a grain boundary as the interface between two crystals, with the same crystallographic structure, with a greater or lesser degree of misorientation [40]. The word 'interfaces' refers, in this work, to the grain boundaries independent of their level of misorientation. Usually, a rotation angle of 15° was defined as the transition angle from LAGB, i.e. subgrain boundaries, and HAGB, i.e. grain boundaries. For pure aluminium, the critical angle for the transition LAGB/HAGB was between 8.6° and 14.4° [41]. In this study, the critical misorientation angle to discriminate the subgrain boundaries and the grain boundaries was chosen to be equal to 10°. For grain boundaries, a distinction was made between coincidence site lattice grain boundaries (CSL), where the degree of fit (Σ) between the two structures of the two grains is described by the reciprocal of the ratio of coincidence sites to the total number of sites oriented, and random grain boundaries where the structure was observed as being disorganised. After the corrosion tests (24 h of immersion in 5 mM NaCl aerated solution), the selected area was analysed using SEM with the backscattered electron (BSE) detector to easily identify any IGC defects (Fig. 1b). The corrosion map obtained was superimposed on the previous EBSD map of the non-corroded material (Fig. 1a) in order to determine the nature of the corroded interfaces (Fig. 1c). This procedure was performed for three corroded non-charged samples and two corroded H-precharged samples; it permitted the determination of the misorientation level for each corroded interface (Fig. 1c).

3. Results and discussion

3.1. Influence of H on the IGC defect morphology

The morphology of the IGC defects was first investigated using OM after a 6 h exposure in a 5 mM NaCl aerated solution at room temperature (25°C). Fig. 2a and b) show OM micrographs of IGC defects formed on non-charged and H-precharged samples, respectively. For the non-charged sample (Fig. 2a), only a few IGC defects were detected (mean value = 6 defects/cm for a cross-section in the L-ST plane). Defects were thin and 86% of them were unbranched. For the H-precharged sample (Fig. 2b), IGC defects were more numerous (mean value = 28 defects/cm) and more branched: 26% of the IGC defects formed could become branched, but the level of ramifications observed was lower than for H-precharged samples. The evolution of the morphology of the IGC defects for the non-charged samples when the exposure to the NaCl solution was longer was assumed to be related to H diffusion all around the IGC defect, H being produced by the corrosion processes [25]. Therefore, the comparison of the IGC defect morphology for non-charged and H-precharged samples led us to assume a significant influence of H on the IGC defect propagation. Furthermore, the evolution of the IGC defect morphology with exposure time to NaCl solution seemed to confirm this hypothesis.

To go further in the characterisation of the IGC defects, conventional lift out procedure using a FEI HELIOS Nanolab 600i dual beam FIB / scanning electron microscope (SEM) was then used to obtain a thin sample in a specific region, i.e. near an IGC defect. Fig. 3 shows representative bright field (BF) TEM images showing a ramifications of an IGC defect formed for an H-precharged sample of AA 2024 T351 after a 24 h immersion in a 5 mM NaCl solution. A continuous precipitation was observed along the two walls of the IGC defects in the upper ramifications. The precipitates were identified as Al-Cu-Mn (71% Al - 18% Cu - 11% Mn, at. %) and Al-Cu-Mg (83% Al - 15% Cu - 2% Mg, at. %) by EDX analyses. The morphology of Al-Cu-Mg precipitates suggested that they were partially dissolved. This was in agreement

![Fig. 1. Methodology used to study the influence of H on the corrosion behaviour of AA 2024 alloy: a) EBSD map of the surface exposed (1.5 x 3 mm²) to the electrolyte before H precharging and corrosion tests; b) SEM image on BSE mode of the corroded surface (same zone) after immersion in a 5 mM NaCl solution during 24 h; c) EBSD map (Fig. 1a) superimposed to the corrosion map (Fig. 1b).](image-url)
with literature data, in particular the work of Birbilis et al. who studied at the nano-scale the dissolution of Al-Cu-Mg particles. The authors concluded that, even if many papers explained IGC considering mainly galvanic effects, their work suggested that the particles themselves dissolved and were the principle contributor to the localization of corrosion on this fine scale [42]. Considering Al-Cu-Mn precipitates, the work performed by Thompson et al. led to assume that the dissolution of the precipitates could not be excluded [21]. Fig. 3 also showed the formation of a Cu-rich layer at the matrix / IGC defect interface composed of Cu nanoparticles that should coincide with the first step of the Cu-rich layer formation as shown by Thompson et al. in their studies about anodisation of Al-Cu model alloys [43]. Overall, the morphology at the TEM scale of the IGC defects formed for the H-precharged sample was similar to that observed for a non-charged sample [20]. Therefore, the results suggested that H did not impact the IGC elementary mechanisms. However, it probably modified the corrosion kinetics, by increasing the corrosion susceptibility of grain boundaries and subgrain boundaries. This was consistent with OM observations (Fig. 2) that showed the increase of the IGC defect density and their ramification for H-precharged samples. To provide experimental proof about this hypothesis, EBSD analyses were therefore performed on H-precharged samples by comparison to non-charged samples.

3.2. H influence on the susceptibility to corrosion of the interfaces depending on their level of misorientation

The distribution of the levels of misorientation of the interfaces for non-corroded AA 2024 alloy was studied for three samples: similar results were obtained with the three samples. Therefore, for more clarity, Fig. 4a represents the results for one sample only. In the selected zone (1.5 × 3 mm²), more than 1000 grains were present, which was a good indication of the number of analysed interfaces. Subgrain boundaries constituted the majority of interfaces, i.e. 59% of the studied interfaces, and only 41% were grain boundaries (Fig. 4a). The dark line in Fig. 4a represents the distribution of the boundary misorientations of a set of grains spread randomly and having a cubic symmetry [44]. Some similarities between the experimental grain boundaries distribution and the theoretical one were observed for θ > 10°, but the theoretical distribution did not completely fit with the experimental results. This could be explained considering the shaping treatment during the plate manufacturing. The presence of subgrains could be linked to the recovery process during the rolling. Moreover, in order to characterise the nature of the interfaces accurately, the distribution of the coincidence site lattice (CSL) grain boundaries was also studied (Fig. 4b). Results showed that 93% of the grain boundaries were random grain boundaries and only 7% were CSL grain boundaries, with Σ3 interfaces representing the majority of the CSL (14%) (Fig. 4b). Therefore, in the following, for the CSL grain boundaries, only the Σ3 grain boundaries were considered.

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**Fig. 2.** Optical micrograph of IGC defects formed after a 6h exposure at E_corr in a 5 mM NaCl solution for a) non-charged and b) H-precharged samples of AA 2024 T351. Insert in Fig. 2a shows an IGC defect after 24 h of immersion in a 1 M NaCl solution for a non-charged sample.

**Fig. 3.** Bright field (BF) TEM images showing a ramification of an IGC defect in H-precharged AA 2024 T351 after a 24 h immersion in a 5 mM NaCl solution.
Then, as explained in the experimental section (Fig. 1), the levels of misorientation of the corroded interfaces were recorded to establish a relationship between the level of misorientation of an interface and its susceptibility to corrosion. For this analysis, the duration of the corrosion tests was 24 h in order to make enough IGC defects grow without introducing a too high H amount. For a non-charged sample, after the corrosion test, 22 corroded interfaces were identified on the corroded area (4.5 mm²); all of them were analysed. Clearly, the number of interfaces analysed for the corroded non-charged sample was significantly lower than for the non-corroded sample, but the results were assumed to be representative considering that the same analysis was performed for three samples and similar results were obtained (Table 1). Fig. 5 thus shows, as an example, the distribution of the levels of misorientation of these 22 corroded interfaces; in this figure, the distribution plotted for the non-corroded sample, with all the interfaces analysed in the selected zone of 4.5 mm², is reported for comparison. Results showed that grain boundaries corresponded to 95% of the corroded interfaces whereas they represented only 41% of all the interfaces in the non-corroded material; further, their susceptibility to corrosion increased with their level of misorientation with a larger number of corroded interfaces identified for the highest levels of misorientation. In particular, it was calculated that 67% of the corroded interfaces had a misorientation angle higher than 40° whereas those interfaces constituted only half of the total interface population. Finally, none of the 22 corroded interfaces observed were Σ3 CSL grain boundaries. The results were in perfect agreement with Guerin’s work performed for an AA 2050 alloy [18] and confirmed the strong reactivity of the grain boundaries with the highest levels of misorientation, which may be linked to their disarranged structure and correlated with their high energy [44]. For the Σ3 CSL grain boundaries, Kim et al. also showed that these interfaces had a good resistance to IGC [16].

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**Table 1**

Summary of the results obtained after the corrosion tests in a 5 mM NaCl solution. *For the corrosion tests in 3.1., results concerned IGC defects: when several corroded interfaces were connected together, they were considered as only one IGC defect. **For the corrosion tests in 3.2., each corroded interface was considered individually so that the concept of “ramification” (branched and unbranched) was not relevant in this part of the study. GB = grain boundaries and sGB = subgrain boundaries.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Non charged</th>
<th>H precharged</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1 2 3 4 5</td>
<td>1 2 3 4 5</td>
</tr>
<tr>
<td>Total number (TN) of IGC defects (defect/cm)</td>
<td>5 7 5 9 3 6</td>
<td>32 27 29 35 19 28</td>
</tr>
<tr>
<td>Number of IGC defects N1 (defect/cm) / (N1/TN) \times 100</td>
<td>0 0 2 2 0 1 / 14</td>
<td>5 8 10 6 7 7 / 26</td>
</tr>
<tr>
<td>Branched (N1)</td>
<td>5 7 3 7 3 5 / 86</td>
<td>27 19 19 29 12 21 / 74</td>
</tr>
<tr>
<td>Unbranched (N1)</td>
<td></td>
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</tr>
</tbody>
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**Fig. 4.** AA 2024 T351: a) Distribution of the levels of misorientation of the interfaces before H precharging and corrosion tests (non-corroded sample): the black curve represents the distribution of the boundaries misorientation of a set of grains spread randomly with a cubic symmetry [44] b) Distribution of the CSL grain boundaries before H precharging and corrosion tests.

**Fig. 5.** Distribution of the levels of misorientation of the corroded interfaces for non-charged and H-precharged samples after a 24 h exposure at Ecorr in a 5 mM NaCl solution. The results obtained for the non-corroded alloy are reported for comparison (all interfaces present in the samples were analysed).
low energy of the subgrain boundaries provided also an explanation to their higher resistance to corrosion \[16,18,19,45,46\].

Comparison of the results with those obtained for the H-precharged and corroded samples permitted to study the influence of H on the susceptibility to IGC of the AA 2024. In this case, results showed that the corrosion damage was significantly more extended in agreement with OM observations (Fig. 2b). Therefore, the EBSD analyses was restricted to an area of 2.3 mm², i.e. half-reduced area by comparison to the one studied for the corroded non-charged sample, in order to facilitate the identification of the corroded interfaces. However, even for this reduced area, the number of IGC defects was equal to 336, all being analysed in this work (Fig. 5). As for the corroded non-charged samples, the experiments were repeated, in this case twice, and similar results were obtained (Table 1). Therefore, for more clarity, only the results obtained for the 336 corroded interfaces from the same H-precharged corroded sample are presented here. For the H-precharged corroded sample, a total of 93% of the corroded interfaces were grain boundaries and 7% of them were subgrain boundaries. Therefore, the results showed that H enhanced the susceptibility to IGC of the AA 2024 with an increase by a factor of fifteen of the number of corroded interfaces. Further, in the presence of H, the most susceptible interfaces to IGC remained the grain boundaries. However, the percentage of subgrain boundaries among the corroded interfaces increased from 5 to 7 for corroded non-charged and corroded H-precharged samples, respectively. Clearly, the increase could be considered as low; however, considering the significant number of corroded interfaces for the corroded H-precharged sample, the results clearly showed the significant enhancement of the corrosion susceptibility for subgrain boundaries: 24 corroded subgrain boundaries for a 2.3 mm² analysed area for the corroded H-precharged sample to be compared to only 1 corroded subgrain boundary for a 4.5 mm² analysed zone for the corroded non-charged sample. Furthermore, as indicated before and as shown in Table 1, the results were reproducible. They are also consistent with results obtained after 6 h of immersion in the NaCl solution (Fig. 2 and Table 1). H was thus found to enhance the susceptibility to corrosion of all the interfaces in such an extent that interfaces, which did not corrode when H amount was low, became susceptible to corrosion for a larger H amount. The corrosion behaviour of E3 CSL grain boundaries confirmed the results since, for the corroded H-precharged sample, 3% of the 336 corroded interfaces were E3 CSL grain boundaries whereas, for the corroded non-charged sample, no E3 CSL grain boundaries were corroded. To further quantify the results, the mean level of misorientation of each distribution was calculated. For the non-corroded sample, the mean level of misorientation was 39°, considering here all the interfaces in the selected zone. As explained before, for the corroded samples, the distributions were restricted to the corroded interfaces. For the corroded non-charged sample, a shift of the distribution was observed towards the high degrees of misorientation compared to the non-corroded sample. The mean level of misorientation was equal to 41.5°, which led us to assume that interfaces with a high level of misorientation were more susceptible to IGC. As indicated previously, the result was obtained for a low number of corroded interfaces, but it was confirmed by the analysis of two other samples and was in agreement with literature data \[17,18\]. By comparison, the distribution of the corroded grain boundaries for the corroded H-precharged sample was more similar to that of the non-corroded alloy, with subgrain boundaries, but also grain boundaries with a low level of misorientation (10° < θ < 40°) that became more reactive leading to a decrease of the mean value of the level of misorientation to 37.5°. In that case, the result was obtained for a large number of corroded interfaces. As done by Guerin et al. \[18\], an IGC susceptibility index (I_{IGC}) was also calculated for the different types of interfaces (subgrain boundaries, random and E3 CSL grain boundaries). It was determined by the ratio of the proportion of corroded interfaces (\%I) divided by the proportion of the interfaces of the same type identified for the n on-corroded sample (\%n_{I}). The results are summarised in Table 2 and confirmed that grain boundaries are more susceptible to IGC than subgrain boundaries. However, in the presence of a large H amount, the subgrain boundaries susceptibility was increased and Σ3 CSL grain boundaries also became susceptible to corrosion.

3.3. New insight into the IGC mechanism for AA 2024

The results presented in the previous two paragraphs permitted to provide a new insight into the IGC mechanism for the AA 2024, highlighting the influence of H produced during the corrosion processes. Fig. 6a is a schematic representation of the AA 2024 T351 microstructure for a non-corroded sample. The most numerous interfaces are subgrain boundaries (59%). Fig. 6b shows the IGC initiation and propagation of short IGC defects for a corroded non-charged sample. The IGC defects initiate following the common mechanism described in the literature \[12\], where intergranular Cu-rich particles promote the corrosion processes due to a galvanic coupling with the adjacent matrix. The intergranular precipitates are mostly present at the grain boundaries, rather than at the subgrain boundaries, because the high energy of the grain boundaries promotes the precipitation phenomenon. As a consequence, the corrosion processes mostly concern the grain boundaries during the first time of exposure to the chloride media; therefore, the main part of the corroded interfaces are grain boundaries (Fig. 2a). However, the corrosion processes lead to the formation of a Cu-rich layer at the interface IGC defect/matrix, that promote the reduction of protons formed due to the cations hydrolysis. H diffused into the sample from the IGC defects where protons are produced as illustrated by SKPFM study \[26\], with the interfaces (both grain boundaries and subgrain boundaries) adjacent to the corroded interfaces acting as preferential diffusion paths. The H enrichment of those interfaces enhances their electrochemical reactivity (Fig. 6d), promoting their dissolution, as shown by AC-SECM measurements \[28\] so that IGC defects can progress laterally and become more and more branched when the duration of exposure to the chloride solution increased (insert in Fig. 2a). Further, Larignon et al. \[26\] showed a decrease of the corrosion potential of AA 2024 in the presence of H; therefore, a galvanic coupling could occur between the H-enriched zone and the adjacent material where the H-enriched zone is the sacrificial anode. This could also explain the propagation of the IGC defects along the subgrain boundaries connected to the corroded grain boundaries. Overall, the H enrichment of the material during the corrosion processes provides an additional explanation to the evolution of the morphology of the IGC defects with increasing duration of exposure to the chloride solution. Clearly, when H was artificially inserted inside the material before the corrosion processes by cathodic charging, the corrosion processes were enhanced more rapidly as shown in Fig. 6c with a stronger ramifications of the IGC defects (Fig. 2b).

4. Conclusions

The present work was focused on the influence of H on the intergranular corrosion (IGC) mechanisms of AA 2024 T351. The main results can be summarised as follows:

1. At the optical and scanning electron microscopic scale, the IGC damage was found to be more extended with a significant ramification of the IGC defects for corroded H-precharged samples. At the transmission electron microscopic scale, no obvious difference was observed between corroded H-precharged and corroded non-charged samples; in both cases, the formation of a Cu-rich layer was observed at the matrix-oxide interface. It was therefore assumed that H did not influence the elementary processes of IGC, but it had a significant impact on the local reactivity.

2. EBSD analyses showed, for a corroded non-charged sample, that the most corroded interfaces were those with the highest levels of misorientation (grain boundaries, θ > 40°). Furthermore, the
presence of H strongly increased the number of corroded interfaces and made subgrain boundaries ($\theta < 10^\circ$), grain boundaries with low misorientation angle ($10^\circ < \theta < 40^\circ$) and $\Sigma 3$ CSL grain boundaries susceptible to IGC.

3 The results were analysed considering previous work that showed that H could both lower the corrosion potential of AA 2024 and increase its reactivity probably due to the disturbance of the crystallographic lattice. They allowed to propose a new insight into the IGC mechanism for AA 2024 where the corrosion-induced H that was trapped inside the IGC defects could then modify the local reactivity of the material. The embrittlement of new interfaces, in which H had diffused, was assumed to explain the ramification of the IGC defects when the duration of exposure to the chloride solution was increased.

In this study, corrosion tests were performed under isothermal conditions and results are thus immediately applicable for structural components used in these conditions. However, when dealing with aerospace industry and real flying conditions, some structural parts can experience thermal cycles with exposure to temperatures $< 0^\circ$C. In those conditions, the solidification of the electrolyte trapped inside the IGC defect can generate stress concentration at the IGC defect tip, which should lead to larger H enrichment of the corroded alloy. Therefore, the H-enhanced corrosion mechanism proposed in this study might be helpful in predicting the lifetime of those structural components.

Data availability

The raw/processed data required to reproduce these findings cannot be shared at this time as the data also forms part of an ongoing study.

Acknowledgement

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### Table 2
Calculation of the IGC susceptibility index ($I_{SC}$) for the different types of interfaces for AA 2024 samples exposed to a 5 mM NaCl solution for 24 h. Non-charged and H-precharged samples are corroded samples here.

<table>
<thead>
<tr>
<th>Type of interfaces</th>
<th>Random and special (except $\Sigma 3$) grain boundaries</th>
<th>$\Sigma 3$ CSL grain boundaries</th>
<th>Subgrain boundaries</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Non charged $%_{nc}$</td>
<td>H precharged $%_{nc}$</td>
<td>Non charged $%_{c}$</td>
</tr>
<tr>
<td></td>
<td>40</td>
<td>40</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>95</td>
<td>90</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>2.38</td>
<td>2.25</td>
<td>0</td>
</tr>
</tbody>
</table>

Fig. 6. IGC Mechanism for AA 2024 T351. a) Microstructure of the non-corroded material with identification of the interfaces. Identification of the corroded interfaces for b) non-charged and c) H-precharged samples after immersion in a NaCl solution. d) Scheme of the H influence on the local reactivity.
References


