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Propagation of Intergranular Corrosion Defects in AA 2024-T351 Evaluated by a Decrease in Mechanical Resistance

Céline Larignon, a Joël Alexis, b Eric Andrieu, a Grégory Odemer, a and Christine Blanc a, b, c, *

French

This study deals with propagation kinetics of intergranular corrosion defects. It complements previous works focused on the development of a procedure called “T-C (Tensile Test for Corrosion).” This procedure was previously applied on samples of 2024 alloy submitted to a continuous immersion test in 1 M NaCl solution. The aim of the present study was to determine if a modification in the way in which the samples are corroded impacts the ability to use this procedure. Therefore, the applicability of the T-C procedure to cyclic corrosion tests was evaluated. Intergranular corrosion defects developed during cyclic corrosion tests were characterized by using a statistical approach. The geometry of the tensile specimens was optimized to allow the T-C procedure to be applied in conditions expected as adequate. Results showed that, for cyclic corrosion tests, due to the branched morphology of the intergranular corrosion defects and the effects of hydrogen embrittlement, the T-C procedure cannot provide an estimation of the mean depth of the intergranular corrosion defects. However, depending on the cyclic corrosion tests, it allowed the maximal depth of the intergranular corrosion defects or the thickness of the corrosion-induced damage zone (defects + hydrogen affected zone) to be determined.

High strength aluminum alloys such as 2xxx series alloys and especially AA2024-T351 are widely used in the aircraft industry because of their good mechanical properties and their low density. However, these alloys are known to be susceptible to localized corrosion such as pitting corrosion,1–3 intergranular corrosion,4,5 exfoliation,6,7 and stress corrosion cracking.8,9 Intergranular corrosion is considered a major threat to the structural integrity of aircraft structures but, today, intergranular corrosion propagation and its impact on structural integrity is still uncertain and therefore it is not easy to include these data in the damage tolerance concept. Therefore, it is of interest to increase the knowledge concerning the kinetics for the propagation of intergranular corrosion defects. A first step for such a predictive approach of the lifetime of aircrafts is to provide data about the kinetics at least under controlled conditions, i.e. at the laboratory scale. However, to be reached, this objective requires the development of an experimental approach allowing characterization of the intergranular corrosion damage and providing data concerning both the intergranular corrosion defects themselves (size, distribution) and a possible volume damage associated with these defects. Indeed, Kamoutsi et al. provided clear evidence of corrosion-induced hydrogen embrittlement in aluminum alloy 2024 and showed that the decrease of the mechanical properties for a corroded sample included both the effect of the corrosion defects and that of hydrogen.10,11

Many authors have used a variety of methods to study intergranular corrosion damage growth and propose propagation laws. For example, Zhang and Frankel used the foil penetration technique on a 2024-T3 aluminum alloy to measure the time required for the fastest growing corrosion sites to penetrate through foils of various thicknesses.13,14 They observed strong anisotropy in the growth kinetics of damage, with slower propagation observed in the short transverse (ST) direction compared with either the long transverse (LT) or the rolling/longitudinal (L) directions. According to these authors, this kinetic anisotropy is directly related with the microstructural anisotropy, which is determined by both grain size and aspect ratio. In a later report, they continued this investigation by developing a statistical model for intergranular corrosion propagation to explain the differences observed in corrosion growth kinetics and the impact of the microstructure.15 Huang and Frankel developed an approach based on visual observations of serial cross sections along the corrosion propagation direction to determine the kinetics of sharp intergranular corrosion cracks in AA7178, monitoring not only the growth rates of long intergranular corrosion cracks but also the rates of the fastest growing sites.16

The methods described above have several limitations, including the fact that neither the random nature of the intergranular corrosion defect propagation nor the three-dimensional morphology of the corrosion defects are taken into account. Additionally, the main parameter responsible for the damage has not yet been identified, with the fastest growing defects or the mean depth of all corrosion defects as two possibilities. Furthermore, only data concerning the corrosion defects are obtained without considering the presence of a possible volume damage. Augustin et al. characterized the intergranular corrosion of the 2024-T351 aluminum alloy after continuous immersion in 1 M and 3 M chloride media by using optical microscopy observations of corroded sample cross sections.17 These observations allowed determination of a statistical value of the mean depth of the intergranular corrosion defects but this method was time consuming because it requires observation of a large number of cross sections to provide representative results. Thus, in the same work, the authors explored another method to estimate the depth of the corrosion defects. This method, called the Tensile Test for Corrosion (T-C) procedure, was based on the evolutions in the mechanical response, particularly the load to failure and the flow stress, between corroded and non-corroded tensile specimens. Because, with the exposure conditions tested (continuous immersion in 1 M and 3 M NaCl solution), the intergranular corrosion defects were statistically distributed along the gauge length of the tensile specimen, the authors assumed that the corroded zone of the sample could be simplified to the development of a mechanically non-load-bearing zone. This approach supposed that the load-bearing zone behaved as a thinner tensile specimen provided that the metallurgical state of the bulk material remained unaffected by the corrosion processes, e.g. hydrogen embrittlement. With this assumption, this method allowed for easier and more rapid determination of the geometric mean depth of the corrosion attack than optical microscopy observations. As this method yielded good results for chloride concentrations of 1 M and 3 M and for immersion times up to 168 h, the authors tested the procedure for a larger range of concentration from 0.5 M to 5 M and for longer immersion times up to 3000 h to define the applicability limits of this new method.18 A 5 M concentration was close to the saturation concentration; it could be considered as representative of the chloride content of the electrolyte trapped inside the corrosion defects under alternate immersion-emersion corrosion tests.19,20 The results of this second study indicated that the T-C procedure was relevant for NaCl concentrations from 0.5 M to 3 M and for immersion times no longer than 120 h.

However, the previous studies were based on continuous immersion tests, while aircraft structures are cyclically exposed to corrosive environments, with the plane exposed to a corrosive environment on the tarmac but not during a flight. Furthermore, in flight, some parts of

*a Université de Toulouse, CIRIMAT, UPS/CNRS/INPT, 31030 Toulouse Cedex 04, France
b Université de Toulouse, LIGP, ENIT, 65016 Tarbes Cedex, France

E-mail: christine.blanc@ensiacet.fr
the structure drop to roughly $-50^\circ C$, while other parts remain at room temperature. Larignon et al. showed that cyclic exposure affected the morphology of intergranular corrosion defects and generated a significant hydrogen embrittlement effect.

Based on these previous results, the present paper focuses on the application of the T$_2$C procedure to cyclic corrosion tests that correspond to corrosion conditions closer to those actually encountered in the aircraft industry. The aim was to assess the effect of both the intergranular corrosion defects and the hydrogen-enriched zone on the mechanical response of the corroded samples to evaluate the interest of the T$_2$C procedure for such corrosion tests. The first part of the manuscript is devoted to characterization by optical microscopy of the intergranular corrosion defects due to cyclic corrosion tests. Based on those results, the second part of the manuscript aims to determine the optimal thickness of the tensile specimen to detect the non-bearing zone generated by the corrosion defects. In the third part of the study, the applicability of the T$_2$C procedure for cyclic corrosion tests is studied.

**Experimental**

**Material.**— Experiments were performed on a 2024 T351 aluminum alloy supplied in the form of a 50 mm thick rolled plate. The chemical composition of the alloy is presented in Table I. The T351 temper consists of a solid-solution heat-treatment at 495°C ($\pm 5^\circ C$) followed by water quenching, straining (1 to 2%) and tempering in ambient conditions for four days. The microstructure of this material consists of grains elongated in the longitudinal direction (L). Average grain sizes in the longitudinal, long transverse (LT) and short transverse (ST) directions were 700, 300 and 100 $\mu$m, respectively. Observations using an optical microscope (OM), a scanning electron microscope with a field emission gun (SEM-FEG) and transmission electron microscopy (TEM) showed both intragranular precipitates, such as Al-Cu-Mn-Fe and Al$_3$CuMg particles, and intergranular precipitates (mainly Al$_3$CuMg and Al-Cu-Mn). No precipitate-free zone (PFZ) was observed along the grain boundaries in the as-received plate. These observations are in good agreement with the literature.4

**Intergranular corrosion tests.**— Both continuous immersion tests and cyclic corrosion tests (alternate immersion-emersion tests) were performed (Table II). For all the corrosion tests, the electrolyte was a 1 M NaCl solution prepared with Rectapur chemicals dissolved in distilled water. Tests were carried out on both cubic samples (10 mm edge) to observe and characterize the intergranular corrosion defects and on tensile specimens (Figure 1) to test the applicability of the T$_2$C procedure. The thickness of the tensile specimens in the L direction was a variable used to improve the applicability of the T$_2$C procedure (see the results and discussion part); it was varied from 1.2 mm to 12 mm. All samples were extracted from the core of the plate. For tensile specimens, if no more details are given in the manuscript, both the LT-ST and ST-L planes of the tensile specimens were exposed to the electrolyte; the mechanically bearing section was the LT-L plane with corrosion defects propagating mainly in the L direction which was the most critical direction for corrosion propagation. For cubic samples, the LT-ST plane was exposed to the electrolyte in order to observe and quantify the intergranular corrosion defects propagating in the L direction. Other planes were masked using a varnish. Independent of the corrosion tests, the same size of the samples was used and the same planes were exposed to the electrolyte. Before corrosion tests, the samples were degreased in ethanol, mechanically polished with different grit SiC papers up to 4000 grit grade, then rinsed in distilled water and air-dried. All corrosion tests were conducted in a laboratory room with controlled temperature ($T = 22^\circ C$). The hygrometry was measured during the testing period (51% relative humidity). Moreover, to avoid temperature fluctuations of the electrolyte due to the size of the laboratory room, the temperature of the electrolyte was also controlled and maintained at 25 ± 2°C by using a thermostat-cryostat. Two cyclic corrosion tests were performed, both consisting of three 24 h cycles. Each cycle consisted of an 8 h immersion at room temperature in the electrolyte followed by a 16 h emersion in air at the temperature of the laboratory room i.e., 22°C for the Cyclic Room (CR) test and at −20°C for the Cyclic Freezing (CF) test. The duration of both the immersion and emersion phases was chosen to be compatible with a work day. Furthermore, additional experiments using a microbalance were carried out to evaluate the time the electrolyte trapped inside the corrosion defects needed to evaporate during the air exposure periods at room temperature (CR tests). Results showed that it could be assumed that, even if corrosion defects could remain wet during the air exposure periods, the electrolyte evaporation was sufficient enough to partially dry the corrosion products and generate

<table>
<thead>
<tr>
<th>Alloying element</th>
<th>Al</th>
<th>Cu</th>
<th>Mg</th>
<th>Mn</th>
<th>Fe</th>
<th>Si</th>
<th>Ti</th>
</tr>
</thead>
<tbody>
<tr>
<td>% weight</td>
<td>4.464</td>
<td>1.436</td>
<td>0.602</td>
<td>0.129</td>
<td>0.057</td>
<td>0.030</td>
<td></td>
</tr>
</tbody>
</table>

**Figure 1. Geometry of the tensile specimens (all dimensions in mm).**

**Table I. Chemical composition of the 2024 T351 aluminum alloy (wt%).**

**Table II. Description of the corrosion tests.**

<table>
<thead>
<tr>
<th>Exposure conditions</th>
<th>Continuous immersion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Continuous immersion in 1 M NaCl (25 ± 2°C)</td>
<td></td>
</tr>
<tr>
<td>Duration 24 h: CI 24 h</td>
<td></td>
</tr>
<tr>
<td>Duration 72 h: CI 72 h</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Exposure conditions</th>
<th>Cyclic corrosion test = alternate immersion (1 M NaCl at 25 ± 2°C) and emersion periods</th>
</tr>
</thead>
<tbody>
<tr>
<td>3 × (24 hour cycles)</td>
<td></td>
</tr>
<tr>
<td>a cycle = 8 hour immersion / 16 hour dry period</td>
<td></td>
</tr>
<tr>
<td>Cumulative immersion time = 3 × 8 = 24 h</td>
<td></td>
</tr>
<tr>
<td>Duration of the cyclic corrosion test = 3 × 24 = 72 h</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Temperature during the dry period</th>
<th>Temperature during the dry period</th>
</tr>
</thead>
<tbody>
<tr>
<td>$22 ± 1^\circ C$</td>
<td>$-20 ± 1^\circ C$</td>
</tr>
<tr>
<td>CR tests</td>
<td>CF tests</td>
</tr>
</tbody>
</table>
a concentration increase of the chemical species trapped inside the corrosion defects. Concerning the CF tests, taking into account the thermal diffusivity of aluminum alloys and the depth affected by the electrolyte penetration, it was assumed that the electrolyte trapped inside the intergranular corrosion defects was frozen immediately at the beginning of the air exposure period at −20°C due to the small size of the samples. Therefore, a 16 h immersion period was long enough to strongly influence the corrosion behavior of the sample. Finally, for the CF tests, since no electrochemical processes could occur at −20°C, the effective corrosion duration was assumed to last 24 h. During the immersion periods, the samples were left at their open circuit potential and hanged in the corrosion cell using a thin nylon yarn. During the air exposure periods, the samples were hanged by using the nylon yarn in a sample glass enclosure placed in the laboratory room for the CR tests and, for the CF tests, the samples were hanged in a container maintained at −20°C. Continuous immersion tests were also performed; they lasted 24 h (CI 24 h) or 72 h (CI 72 h), which corresponded to the cumulative time of immersion for the cyclic corrosion tests or to the global duration of the tests respectively. After the corrosion tests, both cubic samples and tensile specimens were thoroughly rinsed with distilled water, cleaned using an ultrasonic bath and air-dried.

Microscopic observations. — After the corrosion tests, four cross sections (10 mm in length) were cut from the cubic samples along the ST-L plane (perpendicular to the plane exposed to the corrosive environment). Each cross section was abraded with SiC grit paper, then polished down to 1 µm diamond paste and observed with an Olympus PMG3 optical microscope. All intergranular corrosion defects propagating in the L direction, i.e. perpendicularly to the ST direction, were counted and their depth was measured. Taking into account the cumulative length of corroded surface observed in the ST direction, i.e. 40 mm, and the average grain size in this direction, estimated at 100 µm, almost 400 grain boundaries were analyzed in one cubic sample. For cyclic corrosion tests, due to the high number of corrosion defects (see the Results section), only one cubic sample was analyzed for each cyclic corrosion test. However, since the ratio of corroded grain boundaries was lower for continuous immersion tests than for cyclic corrosion tests (see the Results section), two cubes (i.e., 80 mm = 800 grain boundaries) were analyzed for continuous immersion tests. Three parameters were considered as relevant to describe the intergranular corrosion morphology and were statistically determined: the ratio of corroded grain boundaries, the average and maximal depth of intergranular corrosion defects. The ratio of corroded grain boundaries corresponded to the ratio between corroded grain boundaries emerging in the ST-LT plane and the total number of grain boundaries observed (in%). The depth of an intergranular corrosion defect was measured as the distance between the sample surface and the deepest point along the corrosion defect path.

Tensile tests. — Tensile tests were carried out on an Instron testing machine equipped with a 50 kN load cell for the first set of tensile tests (optimization of the tensile specimen geometry, i.e. study of the influence of the thickness on the mechanical response of the corroded samples after continuous immersion tests) and with a 5 kN load cell for the second set of tests, i.e. those concerning the tensile specimens submitted to cyclic corrosion tests. The rate of displacement was set so that the strain rate was close to 10⁻³ s⁻¹. The mechanical variables considered in this study are the load and the strain to failure, as the effective bearing section of the corroded specimen was a priori unknown. Eight un-corroded tensile specimens were tested to establish the reference mechanical properties of the material.

Results and Discussion

Characterization of the intergranular corrosion defects after cyclic corrosion tests. — Figure 2 presents SEM micrographs of intergranular corrosion defects observed on 2024 alloy samples after a CR test (Figure 2a), a CF test (Figure 2b) and a CI 24 h test (Figure 2c) respectively. Comparison between the three micrographs shows that, after cyclic corrosion tests, the intergranular corrosion defects are more branched than after continuous immersion tests. They also seemed to be more numerous. To obtain a quantitative description of the intergranular corrosion defects induced by cyclic corrosion tests, each corrosion defect was counted and their depth in the L direction was measured. The distribution of the depth values (frequency versus depth for regular intervals of depth) was plotted in Figure 3 for both CR and CF tests. For the CR tests (Figure 3a), the depths of the intergranular corrosion defects were quite homogeneously distributed up to 250 µm but some corrosion defects were deeper than the others: 4% of the defects measured were deeper than 250 µm. For the CF tests (Figure 3b), almost 34% of the defects were less deep than 20 µm and the frequency had an exponential-type decrease as the depth values increased. It can also be noticed that no really deep defects were observed as the deepest defect did not penetrate deeper than 250 µm. The high proportion of short defects indicated that this type of exposure tended to promote the initiation of intergranular corrosion defects rather than their propagation. It is known that a geometric mean calculated from a set of values is less dependent on the extreme values than an arithmetic mean of the same set of values. The distributions given in Figure 3 tended to show that the mean depth of the intergranular corrosion defects could be calculated on the basis of either an arithmetic mean or a geometric mean without observing
a large discrepancy between the two mean values. To check this result, the cumulative probability vs. the logarithm of the depth of the corrosion defects was plotted in Figure 4 for both CR (Figure 4a) and CF tests (Figure 4b). Independent of the cyclic corrosion tests, the cumulative probabilities varied linearly with the logarithm of the depth of the corrosion defects but this linear relationship between cumulative probability and the logarithm of the depth was better for CF test than for CR test. This was well-correlated with the presence, for CR tests, of a few very deep intergranular corrosion defects (more than 250 \(\mu m\)). These results showed that it was better to calculate both an arithmetic and a geometric means of the depth values to take into account the presence of very deep intergranular corrosion defects, in particular for CR tests. However, it was assumed that the values should not be very different and, for some figures of the manuscript, only the geometric means of the intergranular corrosion defect depths were reported. Furthermore, to analyze the applicability and the interest of the T2C procedure to cyclic corrosion tests and to determine the information provided by such an approach, it was necessary to provide relevant data concerning the corrosion defect morphology. Therefore, to ensure that observing four cross sections of 10-mm length provided representative data on the corrosion morphology, the ratio of corroded grain boundaries and the arithmetic and geometric mean depths were plotted against the number of corroded grain boundaries observed (Figure 5). For both the CR and CF tests, no significant change in the ratio of corroded grain boundaries was observed from the 400 grain boundaries observed (Figure 5a) and all mean depth values were stabilized too (Figure 5b). As a consequence, the statistical data obtained from the observation of the four cross sections were considered as representative of the corrosion processes. As expected, Figure 5b exhibits a difference between the arithmetic (111 \(\mu m\)) and geometric (72 \(\mu m\)) mean depths for CR test which could be explained, as said previously, by the presence of a few very deep defects (Figure 3a). The geometric mean depth tended to minimize the influence of very deep defects. The difference between the two mean depth values was lower for CF tests as suggested by the distribution (Figure 3b). Furthermore, comparison of the values obtained for both CF and CR tests showed that the ratio of corroded grain boundaries was higher for CF tests than for CR tests (for 400 grain boundaries observed) which confirmed that CF tests promoted the initiation of intergranular corrosion defects. Both arithmetic and geometric mean depths were higher for CR tests than for CF tests. For CF tests, the high proportion of short defects could explain the low mean depth values and indicated once more that this type of corrosion tests tended to promote the initiation rather than the propagation of the intergranular corrosion defects.

Table III summarizes the statistical results obtained for the two cyclic corrosion tests. Results for continuous immersion tests (CI 24 h and CI 72 h) are also given for comparison as well as results obtained by Augustin et al. for a CI 24 h test in 1 M NaCl. First, concerning the ratio of corroded grain boundaries, one third of the grain boundaries were corroded with CR tests and almost half with CF tests. These values were significantly higher than the ratio measured for CI 24 h and CI 72 h tests. Second, the maximal depth of the intergranular corrosion defects was measured at 430 \(\mu m\) for the CR test, with regard to a maximum of 246 \(\mu m\) for the CF test compared to 313 \(\mu m\) for a CI 24 h test. This confirmed that CR tests seemed to promote the propagation of intergranular corrosion defects as well as the initiation while for, CF tests, mainly the initiation of corrosion defects seemed to be promoted. Results from Augustin et al., after a CI 24 h test were in good agreement with the results obtained in the present study. The
differences could be explained by the fact that results were related to two different rolled plates; furthermore, as suggested by Augustin et al., the measurements of corrosion depth are always subjective and depend strongly on the quality of the surface preparation before the OM observations. Augustin et al. showed that it was difficult to obtain relevant data concerning the corrosion depth by analyzing corroded surfaces. For example, they showed that the observation of four cross sections led them to measure a maximal depth of 150 µm for the intergranular corrosion defects while, by polishing step by step a corroded surface, they observed defects as deep as 600 µm. The results led them to assume that the determination of the kinetics of propagation for intergranular corrosion required the development of a new method.

In the present study, the objective is to determine the applicability of the T-C procedure proposed by Augustin et al. These authors indicated that this procedure required that the intergranular corrosion defects were statistically distributed along the gauge length of the tensile specimen. If this condition is validated, the thickness of the non-load-bearing zone, estimated by the T-C procedure, can be estimated a good approximation of the mean geometric depth of the intergranular corrosion defect provided, as said previously, that the material is not affected by a volume damage, e.g. hydrogen embrittlement. Results from the statistical analyzes (Figures 3, 4, and 5) showed that, for both CR and CF tests, the ratio of corroded grain boundaries was high which was compatible with the hypotheses concerning the statistical distribution of the corrosion defects along the gauge length of the tensile specimens. Furthermore, during cyclic corrosion tests, the medium exposed was a 1 M NaCl solution, the cumulative time of immersion was 24 h and the whole duration of the cyclic corrosion tests was 72 h, all of these parameters were in the framework of the T-C procedure application. Thus, it could be assumed that the T-C procedure may be a useful method to evaluate the propagation of intergranular corrosion during cyclic corrosion tests. In their study, Augustin et al. used 1.5 mm-thick tensile specimens (Figure 1) for intergranular corrosion defects developed during continuous immersion tests. As there were numerous short defects in the CF samples with a mean depth value lower compared to CI 24 h, CI 72 h and CR tests, the thickness of tensile specimens had to be adapted to allow the detection of a minimum of a 50 µm thick non-bearing zone. It had to be chosen also to allow the analysis of deeper defects because, for CR tests, the geometric mean depth calculated was 72 µm which was higher compared to continuous immersion tests. Therefore, an optimization of the thickness of the tensile specimens was necessary.

Optimization of the geometry of tensile specimens for application of the T-C procedure to cyclic corrosion tests.— This portion of the study aims to identify an optimal geometry, i.e. optimal thickness, for the testing of tensile specimens using the T-C procedure for cyclic corrosion tests. The T-C procedure, as defined by Augustin et al., assumes that the strength to failure or ultimate tensile stress, \( \sigma_{\text{fail}} \), remained constant for a given metallurgical state of the material. Therefore, the authors proposed to calculate the thickness of the corroded, i.e. non-bearing, zone (Figure 6), named \( x(t) \), where \( t \) is the time of immersion in chloride solution before the tensile test, using the following equation:

\[
x(t) = \frac{a}{2} \left( 1 - \frac{L_{\text{fail}}(t)}{L_{\text{fail}}} \right)
\]

where \( a \) is the initial thickness of the tensile specimen, \( L_{\text{fail}} \) is the maximum load to failure of a non-corroded tensile specimen, and \( L_{\text{fail}}(t) \) is the maximum load to failure measured for a tensile specimen pre-corroded for time \( t \).

As said previously, in their work, Augustin et al. used tensile specimen (Figure 1) with a thickness, \( a \), equal to 1.5 mm. In the present work, to optimize the geometry of the tensile specimens, the same tensile specimens as those used by Augustin et al. were tested but their thicknesses, \( a \), varied from 1.2 to 12 mm (Figure 1). In the present work, the tensile specimens were immersed in a 1 M NaCl solution for 24 h to place them under the T-C conditions and to enable comparison with the results obtained by Augustin et al. Corroded specimens were then subjected to tensile testing to evaluate their residual mechanical properties. In this exposure condition, i.e., 24 h continuous immersion in a 1 M NaCl solution, Augustin et al. calculated a geometric mean depth for the intergranular corrosion defects of 65 µm (from optical observations of corroded samples), compared with a value of 110 µm for the thickness of the non-bearing zone estimated by the T-C procedure. The ST-L faces of the tensile specimens had not been masked from the electrolyte but, the authors ignored in the T-C procedure the corrosion propagation in the LT direction because of the very small surface area exposed in the ST-L plane compared with that exposed in the LT-ST plane. This assumption may not be valid for every tensile specimen (thickness from 1.2 to 12 mm) tested in the present work, and the \( x(t) \) value probably over-estimates the real thickness of the non-bearing zone, at least for some tensile specimens. Thus, in the present work, an alternative method based on the T-C procedure was also used to evaluate the thickness of the non-bearing zone. This method, similar to the T-C procedure, calculated the thickness of the non-bearing zone assuming that propagation in the LT direction was the same as that in the L direction (this meant that the ST-L plane was not protected from the electrolyte). The thickness of

Table III. Statistical analysis of the data concerning intergranular corrosion morphology for cyclic corrosion tests. Results concerning continuous immersion tests are also given for comparison, as well as results from Augustin’s work.¹⁷

<table>
<thead>
<tr>
<th>Corrosion test</th>
<th>CI 24 h</th>
<th>CI 72 h</th>
<th>CR</th>
<th>CF</th>
<th>CI 24 h from Augustin’s work¹⁷</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ratio of corroded grain boundaries%</td>
<td>16</td>
<td>19</td>
<td>35</td>
<td>46</td>
<td>7</td>
</tr>
<tr>
<td>Arithmetic mean depth (µm)</td>
<td>69</td>
<td>72</td>
<td>111</td>
<td>53</td>
<td>90</td>
</tr>
<tr>
<td>Geometric mean depth (µm)</td>
<td>52</td>
<td>54</td>
<td>72</td>
<td>32</td>
<td>65</td>
</tr>
<tr>
<td>Maximal depth (µm)</td>
<td>313</td>
<td>380</td>
<td>430</td>
<td>246</td>
<td>150 (600)</td>
</tr>
</tbody>
</table>

Fig. 6. Schematic of a section of a pre-corroded tensile specimen, with hatched areas corresponding to the non-bearing zone. The \( x(t) \) value varies with time of immersion \( t \). Figure from reference 17.
Figure 7. Schematic of the section of a pre-corroded tensile specimen, with grey areas corresponding to the non-bearing zone. The d(t) value varies with the time of immersion t.

the non-bearing zone, named d(t), where t is the time of immersion in chloride solution before tensile testing, was calculated using the following equation:

$$d(t) = \frac{1}{4} \left[ a + b - \sqrt{(a - b)^2 + 4 \frac{L (t)_{fail}}{L_{fail}}} \right]$$  \[2\]

where b is the width of the tensile specimen and a, L_{fail} and L_{fail}(t) are the same variables used in the T-C procedure as reported in Figure 7. As Zhang and Frankel\textsuperscript{13} observed, the propagation in the LT direction is slower than that in the L direction, thus ensuring that the values of d(t) underestimated the real value of the thickness of the non-bearing zone. However, the calculated x(t) and d(t) values surrounded the real value. Figure 8 displays the x(t) and d(t) values calculated in the present work for various thicknesses of tensile specimens corroded for 24 h in 1 M NaCl solution. For comparison, the geometric mean depth of intergranular corrosion defects measured in the present work, from optical observations of corroded samples (52 µm, Table III) is indicated. Furthermore, the data from Augustin’s work,\textsuperscript{17} i.e. geometric mean depth measured by optical microscopy for the corrosion defects (65 µm) and the thickness estimated by the T-C procedure for the non-bearing zone (x(t) value equal to 110 µm from Equation 1) for samples after 24 h of continuous immersion, are also given.

For tensile specimen thicknesses below 1.5 mm, x(t) values were higher than both values from Augustin et al. (estimated thickness of the non-bearing zone and geometric mean depth) and the geometric mean depth measured in the present work despite the fact that negligible propagation of the corrosion was observed in the LT direction for these tensile specimen thicknesses. The d(t) values for these tests were approximately equal to the geometric mean depth estimated by Augustin et al., but as noted previously, d(t) is assumed to underestimate the real values. This lack of agreement between the thickness of the non-bearing zone determined by tensile testing compared with the geometric mean depth determined by optical observation can be understood by considering that when taking the 700 µm grain size of the material in the L direction into account, no more than two grains lay in the thickness of the tensile specimen. As a result, an excessively thin tensile specimen would be too sensitive to deep defects, meaning that notch effects would also have to be considered and then the tensile specimen would have to be considered a damaged structural element rather than a homogeneously strained sample. Moreover, considering the results obtained during the CR test and especially the mean and maximal depths of the observed intergranular corrosion defects, for an initial tensile specimen thickness below 1.5 mm, the remaining bearing zone would be too thin, meaning that the results would be too sensitive to the deepest defect rather than to the mean depth of the intergranular corrosion defects. This result showed that the thickness of the tensile specimens had to be adapted to the ratio between the grain size of the specimens and the thickness of the non-bearing zone in order to get a mechanical response characteristic of a polycrystal. However, when dealing with the shape of the intergranular corrosion defects, one had to consider that these defects were not very extended in the LT direction and therefore, the notch effect was not significant except in the case of very thin specimens, i.e. thinner than 1.5 mm.

For tensile specimen thicknesses in the range of 1.5 mm to 5.5 mm, the geometric mean depths of the intergranular corrosion defects were close to the estimated thickness values for the non-bearing zone. This was mainly true for the d(t) values because, with this tensile specimen thickness, it was better to consider the propagation of the corrosion defects both in the L and in the LT directions. The results suggested that the geometry of these tensile specimens was suitable for the T-C procedure by using the d(t) values if the ST-L plane was not protected by a varnish.

Tensile specimen thicknesses over 6 mm resulted in non-acceptable values. Between tensile specimen thicknesses of 6 and 9 mm, both the x(t) and d(t) values were too low, underestimating the real geometric mean depth of intergranular corrosion defects. This can be explained by the fact that under the corrosion condition studied here, i.e., 24 h of continuous immersion, the geometric mean depth of intergranular corrosion was approximately 52 µm, representing a decrease in the mechanical load bearing zone of only 2% for a 6 mm thick tensile specimen and 1.3% for a 9 mm specimen. This reduction in the bearing zone is too small to be detected by the tensile test. For tensile specimen thicknesses greater than 9 mm, the results were not representative as the x(t) values were much higher than the geometric mean depth observed by both Augustin et al.\textsuperscript{17} and those of the present work. Moreover, for the thicker tensile specimens, the estimated x(t) value was negative, which is a non-realistic value.

In addition, for all tests, the error in the values estimated from equation 1 was calculated as follows:

$$\Delta x = \frac{1}{2} \left( 1 - \frac{L (t)_{fail}}{L_{fail}} \right) \Delta a + \left[ \frac{a}{2} L_{fail} \right] \Delta L(t)_{fail}$$

$$+ \left[ a \frac{L(t)_{fail}}{2L_{fail}^2} \right] \Delta L(t)_{fail}$$  \[3\]

The results showed that the error was less than 10 µm for tensile specimen thicknesses below 1.5 mm and approximately 30 µm for thicknesses of approximately 3 mm. The error reached 80 µm for thicknesses of up to 5 mm and was greater than 100 µm for thicker specimens. These data indicate that tensile specimen thicknesses in the range of 1.5 to 3 mm seemed to be the most suitable for the evaluation of the intergranular corrosion defect depth by assessing the decrease in the mechanical properties.

To explain and confirm the fact that tensile specimen thicknesses in the range of 1.5 to 5.5 mm are suitable, a rough calculation was performed. First, all tensile tests were assumed to be reproducible as
the differences in the measured mechanical properties and particularly in the ultimate tensile stress between several identical tests do not vary by more than 10 MPa. Thus, to ensure that a measured decrease in the load to failure is due to the growth of a non-bearing zone rather than due to a lack of accuracy for the load cell, the difference in apparent maximal stress between a non-corroded and a pre-corroded tensile specimen ($\Delta \sigma$) must be greater than 10 MPa. Keeping the same notation as in equation 1, the problem was formulated as follows:

$$2x(t) L_{fail} = a (L_{fail} - L_{fail}(t))$$

$$2x(t) \frac{L_{fail}}{a^3b} = \frac{L_{fail}}{ab} - \frac{L_{fail}(t)}{ab}$$

where $L_{fail}/ab = \sigma_{fail}$ is the ultimate tensile stress of a non-corroded sample and $L_{fail}(t)/ab = \sigma_{fail}(t)$ is the apparent ultimate tensile stress of a corroded sample. Thus, the decrease in apparent ultimate tensile stress can be written as:

$$\Delta \sigma = \frac{2x(t)\sigma_{fail}}{a} > 10 \text{ MPa}$$

Thus:

$$a < \frac{2x(t)\sigma_{fail}}{10 \text{ MPa}}$$

The characterization of the corrosion induced by cyclic corrosion tests showed that the non-bearing zone must be at least 50 $\mu$m thick to be detected. As previously mentioned, eight non-corroded tensile specimens were tested to determine the reference mechanical properties, yielding an estimated $\sigma_{fail}$ of the uncorroded material of 450 MPa. Taking these data into account, the maximal thickness of the tensile specimens must be 4.5 mm to provide a relevant value for the thickness of the non-bearing zone. These results are similar to those deduced from the data presented in Figure 8. However, tensile tests revealed good results for thicknesses close to 5.5 mm. This difference is not surprising given that the decrease in ultimate tensile stress with a non-bearing zone of 50 $\mu$m and a tensile specimen thickness of 5.5 mm was approximately 8 MPa, close to the 10 MPa difference estimated previously.

To conclude, these results indicated that to obtain relevant results, tensile specimens tested by the T$_2$C procedure had to be in the range of 1.5 to 4.5 mm thick. In the following part of the present study, the thickness of the tensile specimens was fixed at 2 mm. With 2 mm-thick tensile specimens, it seemed more appropriate to use the $d(t)$ values if the ST-L plane of the specimens were not protected with a varnish to take into account the propagation of the intergranular corrosion defects in the LT direction. However, in the following, for all tensile specimens submitted to a cyclic corrosion test before tensile testing, the surfaces in the ST-L direction were protected with a varnish to consider only the propagation of the corrosion defects in the L direction. Thus, the decrease in the bearing area was essentially related to the propagation of intergranular corrosion on the large sides of the section, i.e. in the L direction. Therefore, the $x(t)$ value was used to estimate the thickness of the non-bearing zone.

**Applicability of the T$_2$C procedure to cyclic corrosion tests.**—Figure 9 presents the results of tensile tests for a non-corroded sample (NC) and corroded specimens CR and CF. For comparison with the results obtained for continuous immersion tests, the load/elongation curve of a pre-corroded CI 24 h specimen is also presented. As expected, the mechanical properties decreased for all the corrosion tests compared to a non-corroded sample. Furthermore, for both the CI 24 h and the CR samples, the global shape of the tensile curve remained unchanged compared with the curve for the non-corroded specimen, with a homothetic shift in the flow stress. With this result, it could be assumed that the corrosion damage was statistically distributed along the gauge length and that the corroded zone could be considered a non-load bearing zone. This result was less convincing for the CF sample for which the plastic domain was very short; however, obviously, a significant shift in flow stress was observed for both CR and CF samples compared to CI 24 h sample with the greatest effect observed for the CF samples despite a lower geometric mean depth for the intergranular corrosion defects for CF samples compared to CI 24 h and CR samples (Table III). Furthermore, a significant decrease of the elongation to failure was observed for both CR and CF samples compared to non-corroded and CI 24 h samples. Based on these tests, the thickness of the non-bearing zones was estimated by the T$_2$C method for both CR and CF samples (Table IV). As said previously, because the ST-L plane was protected by a varnish, the $x(t)$ value was calculated. As assumed from the load/elongation to failure curves, the thickness of the non-bearing zone for the CF test was higher than that estimated for the CR test. Furthermore, for both CF and CR tests, the thickness of the non-bearing zone was very different from both the geometric and arithmetic mean depth calculated from optical observations of the intergranular corrosion defects. For the CR test, the $x(t)$ value was three times higher than the geometric mean depth, and for the CF test, the ratio between the $x(t)$ value and the geometric mean depth was about ten. It was also observed that even though the arithmetic mean depths calculated for the CR and CF tests were higher than the geometric mean depths, the results of the T$_2$C procedure were not in agreement with the arithmetic mean depths either. These results indicated that the T$_2$C procedure was not suitable for the estimation of the mean depth of intergranular corrosion defects during cyclic corrosion tests, as all results over-estimated the real mean depth of the intergranular corrosion defects.

Because the defects observed after cyclic corrosion tests exhibited a more branched morphology that those developed during continuous immersion, deeper defects were assumed to have a greater impact on the decrease in the mechanical properties than those formed during continuous immersion tests, which could contribute to explain the differences between the thickness of the non-bearing zone from

<table>
<thead>
<tr>
<th>Corrosion test</th>
<th>CR</th>
<th>CF</th>
</tr>
</thead>
<tbody>
<tr>
<td>$x(t)$ ($\mu$m)</td>
<td>217</td>
<td>308</td>
</tr>
<tr>
<td>Geometric mean depth ($\mu$m)</td>
<td>72</td>
<td>32</td>
</tr>
<tr>
<td>Arithmetic mean depth ($\mu$m)</td>
<td>111</td>
<td>53</td>
</tr>
</tbody>
</table>
shows that the analysis of 400 grain boundaries was enough to cal-
high, 36%. Almost 15% of the intergranular corrosion defects were
deeper than 200 micrometers and all of the defects were branched, leading to a
spreading of the corrosion affected zone. Indeed, independent of the
corrosion tests, the anodic reaction corresponded to aluminum oxi-
ation, and, for a short immersion time, the cathodic reactions most
likely corresponded to oxygen reduction on the alloy surface and,
when the intergranular corrosion propagated, due to the local acidifi-
cation at the tip of the corrosion defect, to H⁺ reduction inside the
corrosion defect. However, oxygen reduction on the walls of the corro-
sion defects was also a possibility. After a long immersion time, some
depth corrosion defects were observed, with the accumulation of cor-
rosion products inside these defects and on the surface of the sample.
These defects could be rapidly considered an occluded zone charac-
terized by a chloride-enriched electrolyte and by H⁺ reduction as the
major cathodic reaction. For CR tests, partial evaporation of the elec-
trolyte during the emersion period at room temperature led rapidly to
a very high chloride concentration inside the intergranular corrosion
defects. This promoted their propagation based on an autocatalytic
process since the more the aluminum ions were produced, the more the
chloride ions penetrated into the corrosion defects generating a more
aggressive solution. However, the corrosion rate was limited by the
oxygen reduction kinetics due to oxygen transport to the sample sur-
face where the reaction took place. Corrosion products accumulated
inside the corrosion defects so that there was rapidly little connection
between the tip of the corrosion defects and the external electrolyte.‖
Therefore, during the air exposure periods and, for deep corrosion
defects during the immersion period, the main reaction was H⁺ re-
duction inside the corrosion defects and the propagation rate of the
corrosion defects mainly depended on local chemical conditions.‖
In this case, there was no limitation by H⁺ transport processes, which led
to extensive propagation of nearly all the corrosion defects compared
with continuous immersion tests (Table III). Furthermore, due to the
high chloride concentration inside the corrosion defects, new inter-
granular corrosion defects initiated leading to a high ratio of corroded
grain boundaries. Because of the high amount of hydrogen produced
during corrosion processes, a corrosion-induced hydrogen embrittle-
ment could be expected for CR samples as suggested by Kamoutsi
et al.●11●● Larignon et al.●10●● performed hydrogen content measure-
ments by using a Horiba EMGA-621W and a Bruker Galileo HMAt
Instrumental Gas Analyzers (IGA). They showed that the hydrogen
content was 8 ppm, 11 ppm and 25 ppm for non-corroded, CI 24 h
and CR samples respectively. The result showed that hydrogen could
penetrate inside the corroded material and mainly for the CR sam-
pies, in agreement with the kinetics of the corrosion processes, but the
amount of hydrogen trapped inside the material was limited because
only diffusion processes at room temperature could occur to explain
the hydrogen ingress inside the material for CR samples. On the con-
trary, for CF samples, the electrochemical processes were limited due
the emersion period at negative temperatures leading to mean depth

Figure 10. The rupture surfaces of tensile specimens pre-corroded under (a) CR and (b) CF tests. The black area corresponds to brittle surface rupture.

\[ x(t) = \frac{S_b}{2b} \]  

Figure 11 presents the results obtained for CI 24 h, CR and CF samples. Results showed that calculations of the mean depths of the intergranular corrosion defects from surface rupture observations did not show consistent results for CR and CF tensile specimens even though these values were lower than the values estimated for the thickness of the non-bearing zone using the T₂C method.

For CR tests, the ratio of corroded grain boundaries was relatively high, 36%. Almost 15% of the intergranular corrosion defects were deeper than 200 micrometers and all of the defects were branched, leading to a spreading of the corrosion affected zone. Indeed, independent of the corrosion tests, the anodic reaction corresponded to aluminum oxi-
for the intergranular corrosion defects lower than for CR samples and even CI 24 h samples. However, due to their volume expansion during the emersion period at negative temperatures, the electrolyte and/or the corrosion products trapped inside the corrosion defects generated mechanical stresses strong enough to create a strain state at the tip of the intergranular corrosion defect that could promote hydrogen transport. \(^{16,20}\) Hydrogen/dislocations interactions were studied by different authors. For example, Zohidi and Meletis showed that, in the presence of a crack, hydrogen penetration in front of the crack tip can occur mainly by stress assisted diffusion and dislocation transport. \(^{24}\) It was assumed that, for CF tests, this phenomenon was added to a classical hydrogen diffusion process which could occur only during the immersion period at room temperature since the hydrogen diffusion coefficient was very low at \(-20\)°C. Larignon et al. measured a hydrogen content between 50 and 90 ppm for CF samples which corroborated this assumption. \(^{19,20}\) Further work performed by the same authors provided evidence of the presence of hydrogen inside the CF samples. \(^{21-23}\)

Taking all of these data into account, the applicability of the T\(_C\)C procedure to cyclic corrosion tests can be discussed. Kamoutsi et al., who provided clear evidence of corrosion-induced hydrogen embrittlement in aluminum alloy 2024, \(^{11,12}\) showed that, to consider the possible effect of hydrogen embrittlement as a volume effect, the corrosion layer should have been removed on the tensile specimens and then tensile specimens be constructed to differentiate the effect of corrosion from the effect of hydrogen on ductility. However, for the CR tests, due to the hydrogen content, it was assumed that hydrogen embrittlement could contribute to the mechanical response of the corroded sample but was not the major phenomenon; the discrepancy between the mean depth of the intergranular corrosion defects measured by optical microscopy and the thickness of the non-bearing zone from the T\(_C\)C procedure was mainly attributed to the size of the intergranular corrosion defects following CR tests. However, 90% of the intergranular corrosion defects observed penetrated less than 220 µm, close to the thickness of the non-bearing zone estimated by the T\(_C\)C procedure (217 µm). Therefore, for the CR test, the results of the T\(_C\)C procedure were more closely related to the maximal depth of the majority of the intergranular corrosion defects than to the mean depth of all defects. Moreover, the few very deep defects, i.e., those deeper than 250 µm, do not have a significant influence on the decrease in the mechanical properties as Augustin et al. suggested. \(^{27}\)

For CF tests, the large proportion of very short defects led to a low mean depth of the intergranular corrosion defects compared to the estimation of the non-bearing zone thickness from the T\(_C\)C procedure. In addition, neither the T\(_C\)C procedure nor the study of the rupture surface resulted in agreement with the mean depths of intergranular corrosion defects from optical observations. Previous results about hydrogen embrittlement \(^{14,25}\) led to the assumption that, for CF tests, the T\(_C\)C procedure did not allow estimation of the propagation kinetics (mean depth or maximal depth) of intergranular corrosion defects because the decrease of the mechanical properties took into account both intergranular corrosion propagation, which is visible by optical observation, and a hydrogen affected zone that cannot be seen under the microscope. Furthermore, the large proportion of brittle fracture on the rupture surfaces could be attributed to corrosion-induced hydrogen embrittlement. For CF tests, the T\(_C\)C procedure seemed to provide an estimation of the damaged thickness of the tensile specimens including the maximal depth of the intergranular corrosion defects and hydrogen affected area. Thus, the T\(_C\)C procedure could be a good method to estimate the damage thickness to take into account for gauging calculations during the design of structural parts. However, as suggested by Kamoutsi et al., \(^{11,12}\) to differentiate the effect of the corrosion defects from that of hydrogen embrittlement, heat-treatment should be applied to generate hydrogen desorption but this required evaluation of the effect of the desorption treatment on the mechanical properties of the alloy.

**Conclusions**

This study examined the applicability of the T\(_C\)C procedure for cyclic corrosion tests and provides a critical analysis of this methodology previously proposed to determine the mean depth of the intergranular corrosion defects on the basis of continuous immersion tests. The T\(_C\)C procedure was based on the analysis of the mechanical behavior of corroded samples compared to non-corroded specimens.

(+) Cyclic corrosion tests were alternate immersion-emersion with emersion periods at room temperature for CR tests and at negative temperatures, heat-treatment being performed after corrosion tests. They generated intergranular corrosion defects more branched than after continuous immersion. After CR tests, the intergranular corrosion defects were also deeper than those after continuous immersion while, for CF tests, they were shorter. Differences in morphology were explained by taking into account the electrochemical processes. For both CR and CF tests, the ratio of corroded grain boundaries was higher compared to continuous immersion suggesting that the intergranular corrosion defects could be statistically distributed along the gauge length of the tensile specimen. Therefore, T\(_C\)C procedure was at first expected to be applicable for cyclic corrosion tests.

(-) The geometry of the tensile specimens used for the T\(_C\)C procedure, i.e. their thickness, was optimized on the basis of intergranular corrosion defects developed during continuous immersion tests. Results clearly indicated the major parameter to take into account in such an approach.

The T\(_C\)C procedure was applied to cyclic corrosion tests. For CR tests, the T\(_C\)C procedure was assumed to provide a good estimation of the maximal depth of the intergranular corrosion defects. For CF test, this procedure was found to be inadequate due to a hydrogen embrittlement effect. However, it allows calculation of the thickness of a corrosion damaged zone including both intergranular corrosion defects and the hydrogen embrittled zone.

**References**