Investigation of Stress Corrosion Cracking of Austenitic, Duplex and Super Duplex Stainless Steels under Drop Evaporation Test using Synthetic Seawater

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Stress corrosion cracking (SCC) of UNS S31603 austenitic stainless steel (ASS), UNS S32205 duplex stainless steel (DSS) and UNS S32750 super duplex stainless steel (SDSS) was investigated. SCC tests were carried out at 110 °C for 500 h under drops of synthetic seawater (DET, drop evaporation test). Two loading conditions were investigated: 50 % and 100 % of the experimental yield strength of each steel. DSS and SDSS specimens showed no susceptibility to SCC under loading of 50 % of their yield strength, contrary to ASS, but all steels fractured at the highest load. SCC nucleated under the salt deposit formed on the surface of all specimens. SCC propagation was mainly transgranular, but SCC propagation of DSS also featured crack ramification in the austenite phase. In addition, SDSS also presented crack propagation along the ferrite/austenite interfaces. Transgranular cleavage fracture was also observed in all fractured specimens, but DSS also presented ferrite/austenite interfacial brittle fracture, while SDSS also featured intergranular brittle fracture.

Keywords: Stress Corrosion Cracking, Drop Evaporation Test, Stainless Steels.

1. Introduction

In Brazil, ninety-five percent of the petroleum is extracted from offshore wells, requiring a large range of engineering components and materials, which should present high corrosion resistance to marine environments. In this sense, ASS, DSS and SDSS are usually selected for the manufacturing of offshore pressure vessels1, due to their high corrosion resistance and good mechanical properties. These offshore components are usually submitted to service conditions combining tensile stress and chloride environments, which may induce their premature failure by stress corrosion cracking (SCC), even under tensile values below the yield strength2-4. A comparison of the general properties and characteristics of the ASS and DSS and SDSS is shown in Table 1. Although DSS and SDSS are more expensive than ASS, their values of fracture toughness, yield strength and pitting resistance equivalent number (PREN) are comparatively much higher5. Despite their wide and growing usage of DSS and SDSS components, they might be subjected to various types of mechanical and environmentally induced fractures during their life cycle, especially when these components are exposed to temperatures in the range of 300 °C to 900 °C, which might promote the precipitation of stable and deleterious phases4-5.

Stainless steels (SSs) may suffer SCC when three critical conditions occur simultaneously: microstructural susceptibility, corrosive environment and tensile stress. The weight of each one of these critical parameters is different for each class of SS4,6-9. Environments containing chloride ions, however, are especially harmful to the strength of the passive film of SSs, promoting the pitting corrosion and increasing the susceptibility to SCC in these steels1-3,5-7. Additionally, the presence of other cations, such as magnesium and calcium, might further increase the aggressivity of the chloride environments and the susceptibility of SSs to SCC10. Actually, the SCC of SSs might occur even in caustic solutions, when the surfaces of the component present regions of pitting or crevice corrosion11-12. SCC might also happen when the anodic reaction takes place underneath an occluded region of the surface, causing a significant decrease of the solution’s pH11-12. Although pitting and crevice corrossions are not prerequisites for the nucleation of SCC in SSs, they can act as stress raisers and solution acidifiers, promoting, therefore, the nucleation of SCC2,5-7,10-14.

The stable crack propagation of ASS, DSS and SDSS usually presents a ductile behavior under non aggressive environments, but the action of SCC promotes a brittle behaviour in these steels, leading to the formation of intergranular or cleavage-like transgranular brittle fractures15. It is interesting to observe that the exposure to a corrosive environment might decrease the fracture toughness of ASS from \( K_{IC} \) equals to 72 MPa.m\(^{1/2}\) to \( K_{ISC} \) (acidic solution) equals to 53 MPa.m\(^{1/2}\). There are at least four atomic-level
mechanisms to explain the stable brittle crack propagation of metals and alloys caused by SCC:

1. The tensile decohesion of the atomic bounds of the metallic material ahead of the crack tip (cleavage) is promoted by the adsorption of ions of the corrosive solution on the metallic surfaces near the crack tip. This adsorption is followed by surface diffusion of these ions into the region ahead of the crack tip, causing local embrittlement due to the decohesion of the atomic bounds (see Figure 1);

2. The creation of vacancies on the metallic surfaces near the crack tip is caused by the removal of the elements of the crystal lattice by the corrosive solution. These vacancies preferentially diffuse towards the region ahead of the crack tip due to the presence of a stress gradient. The metallic ion surface mobility is promoted by the presence of contaminants in the corrosive solution. The stable step-like brittle propagation of the crack takes place when these vacancies reach the crack tip (see Figure 2);

3. The formation of a brittle surface film on the metallic surfaces near the crack tip locally reduces the fracture toughness, promoting the stable step-like brittle propagation of the crack. According to this model, the formation of the brittle film is induced by the environment, which controls the kinetics of the stable crack propagation. In this sense, the next step of the stable brittle propagation of the crack will proceed when another layer of brittle film is formed ahead of the crack tip (see Figure 3);

4. The shear strain in the region located ahead of the crack tip is promoted by the adsorption of atomic hydrogen on the surfaces of the crack (see Figure 4). These hydrogen atoms diffuse into the FCC lattice, preferentially into the region located ahead of the crack-tip. The presence of interstitial hydrogen atoms in this region locally increase the plasticity of the metal. In the case of FCC iron, the hydrogen atom (atomic radius equals to 0.48 Å)

| Property                         | UNS S31603 | UNS S32205 | UNS S32760 |
|----------------------------------|------------|------------|------------|
| Price (R$/kg)                   | 18.5 to 21.4 | 20.6 to 22.4 | 29.4 to 31.8 |
| Elastic modulus (GPa)            | 190.0 to 205.0 | 195.0 to 205.0 | 190.0 to 200.0 |
| Yield strength (MPa)             | 170.0 to 310.0 | 460.0 to 510.0 | 515.0 to 565.0 |
| Fracture toughness (MPa.m(1/2))  | 53 to 72    | 144 to 195 | 150 to 195  |
| Sea water corrosion resistance   | Good       | Good       | Excellent   |
| Pitting resistance equivalent number (PREN) | 22.6 to 27.9 | 30.5 to 37.0 | 37.9 to 45.7 |
| Pitting and crevice corrosion (µm) | Medium (20 to 30) | High (30 to 40) | Very high (> 40) |
| CO₂ footprint (kg/kg)            | 5.3 to 5.9  | 4.4 to 4.9  | 5.1 to 5.6  |

Figure 1. SCC mechanism: tensile decohesion of the atomic bounds due to the presence of adsorbed ions in the crystal lattice ahead of the crack tip.

Figure 2. SCC mechanism: cracking propagation of the crack tip due to the surface mobility induced by corrosive environment.

Figure 3. SCC mechanism: formation of a brittle surface film at the crack tip.
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2. Materials and Methods

The specimens used for the drop evaporation tests were machined from 12.7 mm thickness plates and their microstructures are shown in Figure 5-a to 5-c. Figure 5-a features the microstructure of the UNS S31603 ASS specimen, showing equiaxial austenitic grains with small amount of ferrite strips, following the rolling direction of the plate. The center of this Figure shows an unetched region, due to the work hardening of the ASS plate. Figures 5-b and 5-c feature the microstructures of UNS S32205 DSS and UNS S32750 SDSS, respectively, revealing a lamellar microstructure composed of ferrite (dark region) and austenite (white region) phases with elongated grains, indicating the rolling direction of the plate (KOH electrolytic attack was used to identify the ferrite phase).

The chemical compositions of the SS used in the present investigation are shown in Table 2. Carbon and sulfur contents were measured by the combustion method using a LECO analyzer (model CS844) and the remaining elements were measured by wavelength dispersive X-ray fluorescence spectrometry (Rigaku WDXRF, model RIX 3000). The chemical compositions of the SSs are in accordance with the requirements of ASTM A240 standard.

Tensile mechanical tests were performed using a 10 t MM-004 tensile testing machine, according to the ABNT NBR ISO 6892 standard and using cylindrical specimens with diameter of 5 mm. The results are shown in Table 3, indicating that the tensile properties of the SSs are in accordance with the requirements of the ASTM A240 standard. Table 3 also features the investigated tensile stresses during DET for each SS.

DET specimens (10 mm x 25 mm x 178 mm) were machined along the perpendicular plane of the rolling direction. A schematic representation of the mechanical loading assembly using “four point bending” is shown in Figure 6 along with the identification of its geometrical parameters. The roughness of the top surface of the specimens (exposed to the dripping during the DET) was measured by a KOSAKA rugosimeter to guarantee that the roughness values (Ra) of the specimens were lower than 0.7 µm. The maximum tensile stress on the surface of the specimens after bending was calculated by Equation 1. Two mechanical loading conditions were investigated: a) maximum tensile stress equal to 50 % of the experimental yield strength at room temperature (see Table 3); b) maximum tensile stress equal to 100 % of the experimental yield strength at room temperature (see Table 3).

\[
\sigma = \frac{12 \cdot E \cdot t \cdot y}{3 \cdot H^2 - 4 \cdot A^2}
\]

Where \(\sigma\) is the tensile stress upon the maximum bending strain (MPa); \(E\) is the elastic modulus (MPa); \(t\) is the thickness of the specimen (mm); \(y\) is the bending height between the superior loading point and the center of the specimen (mm); \(H\) is the distance between the upper loading points (mm); and \(A\) is the distance between the upper and lower loading points (mm).
Pre-loaded specimens were placed in chambers with heating resistances controlled by a proportional-integral-derivative controller (PID, NOVUS® model N1030). Synthetic seawater used was prepared according to the ASTM D1141 and ASTM D1193 standards, featuring a pH equals to 8.2 and chemical composition described in Table 4 38-39.

DET was carried at 110 ºC for 500 h and the dripping rate of synthetic sea water was kept equal to (10 ± 1) drops.

**Table 2.** Chemical composition of UNS S31603 ASS, UNS S32205 DSS and UNS S32750 DSS *.

| Steel            | C     | Si    | Mn    | P     | Cr    | Ni    | Mo    |
|------------------|-------|-------|-------|-------|-------|-------|-------|
| UNS S31603 (ASS) | 0.019 ± 0.001 | 0.44 ± 0.02 | 1.34 ± 0.01 | 0.035 ± 0.002 | 17.1 ± 0.1 | 10.1 ± 0.1 | 2.16 ± 0.01 |
| ASTM A240™<br>UNS S31603 | < 0.03 | < 0.75 | < 2.00 | < 0.045 | 16.0 to 18.0 | 10.0 to 14.0 | 2.0 to 3.0 |
| UNS S32205 (DSS) | 0.017 ± 0.002 | 0.38 ± 0.01 | 1.44 ± 0.02 | 0.026 ± 0.002 | 22.3 ± 0.3 | 5.79 ± 0.07 | 3.08 ± 0.03 |
| ASTM A240™<br>UNS S32205 | < 0.03 | < 1.00 | < 2.00 | < 0.030 | 22.0 to 23.0 | 4.5 to 6.5 | 3.0 to 5.0 |
| UNS S32750 (SDSS) | 0.015 ± 0.001 | 0.38 ± 0.02 | 1.44 ± 0.05 | 0.029 ± 0.002 | 24.9 ± 0.2 | 6.92 ± 0.06 | 3.76 ± 0.04 |
| ASTM A240™<br>UNS S32750 | < 0.03 | < 0.80 | < 1.20 | < 0.035 | 24.0 to 26.0 | 6.0 to 8.0 | 3.0 to 5.0 |

*Sulphur contents are not show in the table, but they were in accordance with the requirements of ASTM A240™.

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**Figure 5.** Microstructure of the investigated SSs: (a) UNS S31603 ASS; (b) UNS S32205 DSS; (c) UNS S32750 SDSS. Optical microscope. Electrolytic etching: oxalic acid for ASS and KOH for DSS and ASS.
Table 3. Mechanical properties of UNS S31603 ASS, UNS S32205 DSS and UNS S32750 SDSS at room temperature and the values of maximum tensile stress used for DET for conditions (a) and (b).

| Property                        | ASS (MPa) | ASTM A240 (MPa) | DSS (MPa) | ASTM A240 (MPa) | SDSS (MPa) | ASTM A240 (MPa) |
|--------------------------------|-----------|-----------------|-----------|-----------------|-----------|-----------------|
| Yield strength                 | 278 ± 2   | > 170           | 473 ± 3   | > 450           | 589 ± 3   | > 550           |
| Ultimate tensile stress        | 629 ± 4   | > 485           | 749 ± 4   | > 655           | 877 ± 4   | > 795           |
| Elongation (%)                 | 51 ± 1    | > 40            | 35 ± 1    | > 25            | 37 ± 1    | > 15            |
| DET, condition (a): 50% of the yield stress (MPa) | 139       | -               | 236       | -               | 295       | -               |
| DET, condition (b): 100% of the yield stress (MPa) | 278       | -               | 473       | -               | 589       | -               |

Table 4. Saline concentration of synthetic sea water according to ASTM D114138-39.

| Compounds   | Concentration (g/L) |
|-------------|---------------------|
| NaCl        | 24.53               |
| MgCl₂       | 5.20                |
| Na₂SO₄      | 4.09                |
| CaCl₂       | 1.16                |
| KCl         | 0.695               |
| NaHCO₃      | 0.201               |
| KBr         | 0.101               |
| H₃BO₃      | 0.027               |
| SrCl₂       | 0.025               |
| NaF         | 0.003               |

A salt deposit in the form of a “volcano” was formed on the surfaces of all DETed specimens. The surfaces on the dripping region, however, did not feature the presence of any deposit (see Figure 7).

Figures 8-a to 8-c show the general aspects of the DETed specimens after rinsing with water. Figure 8-a shows an ASS specimen tested at 50% of its yield stress (139 MPa) which exhibits some dark oxidized regions on the surface of Quanta 3D FEG dual beam microscope was also used as focused ion beam (FIB) microscopy in order to characterize the microstructure under the surface of SCC (cross-sectional examination).

3. Results

A salt deposit in the form of a “volcano” was formed on the surfaces of all DETed specimens. The surfaces on the dripping region, however, did not feature the presence of any deposit (see Figure 7).
the specimen without any evidence of cracking during the visual examination. Additionally, its dripping zone does not show any signs of oxidation. DSS specimen tested at 50 % of its yield stress (236 MPa) and SDSS specimen tested at 50 % of its yield stress (295 MPa) did not show evidences of cracking during the visual examination. However, all specimens tested under tensile loading of 100 % of their yield strength presented cracking or fracture during visual inspection. The general view of a DSS specimen after testing at 100 % of its yield stress (473 MPa) is shown in Figure 8-b, where a 45° crack is observed (see yellow arrow). A SDSS specimen after testing at 100 % of its yield stress (589 MPa) is observed in Figure 8-c, where multiple cracks can be observed (see yellow arrow). Both crack nucleation sites coincide with the position of the salt layer. Additionally, the dripping zones located in the center of the specimens do not show any signs of oxidation. A closer examination of the dark oxidized regions of all the tested specimens, located away from the salt layer, showed corrosion products, such as pitting and localized-corrosion, indicating that temperature of 110 °C is above the critical limit for the pitting corrosion.

Contrary to the visual examination, the metallographic examination of an ASS specimen tested at 50 % of its yield strength (139 MPa) showed ramified cracking, which was nucleated in a corrosion pit under the salt deposit (see Figure 9-a). Furthermore, the metallographic examination of this sample

Figure 8. Visual inspections of specimens tested at 110 °C for 500 hours after the cleaning. (a) UNS S31603 ASS tested at 50 % of the yield stress (139 MPa); (b) UNS S32205 DSS tested at 100 % of the yield stress (473 MPa); and (c) UNS S32750 SDSS tested at 100 % of the yield stress (589 MPa). Visual inspection.

Figure 9. Metallographic examination of the crack propagation, cross section after DET at 110 °C. (a) UNS S31603 ASS tested at 50 % of the yield stress (139 MPa); (b) UNS S32205 DSS tested at 100 % of the yield stress (473 MPa); (c) UNS S32750 SDSS tested at 100 % of the yield stress (589 MPa). Optical microscope.
indicated multiple cracks and the longest one showed a depth of approximately 3 mm. The cracks were transgranular and featured intense ramification after a crack depth of 600 µm and the primary crack preferentially propagated normal to the surface of the specimen. Metallographic examination of a DSS specimen tested at 100 % of its yield strength (473 MPa) presented a 45º crack propagation followed by a horizontal crack deflection near the plate’s center, where the alloy exhibited a more prominent work-hardened microstructure (see Figure 9-b). The crack propagation of the DSS was comparatively less ramified than the ASS (see Figure 9-a), but their secondary cracks were comparatively much deeper (maximum depth of 2500 µm). Metallographic examination of a SDSS specimen tested at 100 % of its yield strength (589 MPa), see Figure 9-c, exhibited multiple nucleation sites and, comparatively, more intense crack ramification. The secondary cracks were comparatively deeper than the DSS (maximum depth of approximately 4000 µm).

Figure 10-a presents the crack propagation of an ASS specimen tested at 100 % of its yield stress (278 MPa), presenting transgranular propagation and crack deflection at grain boundaries. Figure 10-b presents the crack propagation of a DSS specimen tested at 100 % of its yield stress (473 MPa), showing transgranular propagation and preferential crack ramification inside the austenite phase. Figure 10-c presents the crack propagation of a SDSS specimen tested at 100 % of its yield stress (589 MPa), showing transgranular and interfacial propagation and preferential crack ramification inside the austenite phase. The cracks of the DSS and SDSS propagated preferentially at 45º of the rolling direction and parallel to the maximum shear stress of the bended specimens, while the crack of ASS propagated normal to the rolling direction.

Figures 11-a to 11-c show the microfractographic examination of the exposed crack surfaces. The fracture of an ASS specimen tested at 100 % of its yield stress,
Figure 11. Microfractographic examination of the exposed surface of the cracks after DET at 110 °C. (a) UNS S31603 ASS tested at 100 % of the yield stress (278 MPa); (b) UNS S32205 DSS tested at 100 % of the yield stress (473 MPa); (c) UNS S32750 SDSS tested at 100 % of the yield stress (589 MPa). SEM.

278 MPa, is shown in Figure 11-a, which exhibits a brittle fracture with stretch marks and few secondary cracks, typical of transgranular cleavage. The fracture of a DSS specimen tested at 100 % of its yield stress (473 MPa) is shown Figure 11-b, which features cleavage fracture and intense presence of secondary cracks and interfacial brittle fracture (see flat facets), resulting in transgranular and intergranular brittle fracture. The fracture of a SDSS specimen tested at 100 % of its yield stress (589 MPa) is shown Figure 11-c, which features transgranular cleavage with more intense secondary cracking. Additionally, the presence of interfacial and grain boundary brittle fractures (see comparatively smaller flat facets) can also be observed.

Figures 12-a to 12-c display the images obtained by the FIB cross section examination near the SCC surfaces. Figure 12-a shows the microstructure under a SCC surface (ASS specimen tested at 100 % of its yield stress - 278 MPa), presenting angular deflection of the SCC at a grain boundary (see solid arrow) and a small area of intense plastic deformation near the SCC surface (see dotted arrow). Figure 12-b shows the microstructure under the SCC surface (DSS specimen tested at 100 % of its yield stress - 473 MPa), exhibiting various secondary cracks with transgranular and interfacial propagation and crack ramification (see solid arrows). Additionally, a small area of intense plastic deformation is also observed near the SCC surface (see dotted arrow). Figure 12-c presents the microstructure under the SCC
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4. Discussion

The formation of a thick salt deposit associated with the SCC in all DSS and SDSS specimens tested at 110 °C (see Figure 7) is in accordance with previous results of Turnbull and Hinds\(^{30-31}\) and Steinsmo and Drugli\(^{32}\). Turnbull and Hinds\(^{31}\), for instance, investigated the susceptibility of 22 Cr and 25 Cr DSSs to SCC under evaporative seawater conditions using a modified DET. The use of flat specimens allowed the seawater to accumulate on the dripping zone, producing a hollow-damp salt deposit, whose edges advance along the gauge length during the test (see Figure 13).

In another investigation, the authors\(^{41}\) showed that during DET at 110 °C, the solution found at the center of the specimens (dripping zone) presented a pH value of around 11, much higher than the original pH of the synthetic seawater solution (pH equal to 8.2), indicating the occurrence of a cathodic reaction in this region (see Figure 14-a). Additionally, the temperature of the DET specimens in the vicinities of the dripping zone was at least 10 °C lower than the remaining regions, due to the cooling action of the synthetic solution.
Figure 13. Scheme of the salt deposit formed on all the DETed specimens tested at 110 °C for 500 h. The SCCs were nucleated under the salt deposit, approximately 22 mm away from the center of the dripping region, by crevice corrosion. 

The formation of salt deposit on the samples tested at 110 °C promoted the crevice corrosion of the plate under the deposit (see Figure 7), followed by the preferential attack (ferrite phase and austenite/ferrite interfaces), the formation of a corrosion pit and, finally, the nucleation and growth of SCC, confirming previous results. The salt deposit was identified by Pereira as a mixture of CaSO₄, NaCl, Mg(OH)₂, MgCl₂, which is hygroscopic, creating a corrosive wet interface between the salt deposit and the surface of the specimen. The observation of occasional corrosion and pitting beneath the salt deposit is in accordance with the observation of previous work, indicating that the combination of aggressive solution under the salt deposit (with pH values between zero and one) and microplastic deformation was responsible for SCC of the SSs tested above 105 °C (this temperature coincides with the evaporation temperature of the seawater). FIB results indicated the presence of small regions of plastic deformation below the SCC surfaces of ASS and DSS specimens, as suggested by previous investigation on SCC of DSSs.

ASS samples showed multiple nucleation sites even for DET at 110 °C and loading of 50 % of its yield stress (139 MPa), see Figure 9-a, indicating that its SCC threshold stress at 110 °C is above 139 MPa. DSS and SDSS samples presented fractured DET specimens at 110 °C and loading of 100 % of the yield strength (473 MPa and 589 MPa, respectively), see Figures 9-b and 9-c, showing that their threshold stress for SCC is between 50 % and 100 % of their yield strength (see Table 3).

Additionally, ASS specimens tested at 100 % of its yield stress (278 MPa) showed transgranular crack propagation and crack deflection at grain boundaries, while the crack propagation of UNS S32205 DSS specimens tested at 100 % of the yield stress (473 MPa) also revealed preferential crack ramification inside the austenite phase. SDSS specimens tested at 100 % of yield stress (589 MPa) featured additional ferrite/austenite interfacial crack propagation (see Figures 10-a to 10-c). The cracks of the DSS and SDSS preferentially propagated at 45° of the rolling direction (parallel to the maximum shear stress of the bended specimens), while ASS cracking was perpendicular to the tensile stress. Microfractographic examination of ASS specimens tested at 110 °C with 100 % of the yield stress (278 MPa) exhibited transgranular brittle fracture, while DSS specimens tested at 110 °C with 100 % of the yield stress (473 MPa) also featured austenite/ferrite interfacial brittle fracture. SDSS specimens tested at 100 % of the yield stress (589 MPa) also featured intergranular brittle fracture (see Figures 11-a to 11-c).

DETed ASS specimen tested at 100 % of its yield stress (279 MPa) presented SCC deflection at the grain boundaries and plastic deformation of the austenite grains near the SCC surface (see Figure 12-a). DETed DSS specimen tested at 100 % of its yield stress (473 MPa) additionally exhibited ramified secondary cracks with transgranular and interfacial propagation (see Figure 12-b). SDSS specimen tested at 100 % of its yield stress (589 MPa) showed the influence of interfaces and grain boundaries on the crack deflection, promoting a step-like crack propagation (see Figure 12-c). These observations are in agreement with previous result.

Finally, the SDSS specimens showed the lowest SCC susceptibility at 110 °C in seawater, followed by DSS and finally by ASS specimens.

5. Conclusions

The results of SCC sweater DET of ASS (UNS S31603), DSS (UNS S32205) and SDSS (UNS S32750) performed at 50 % and 100 % of the yield stress at 110 °C for 500 hours indicated that:

- SCC presented preferential nucleation sites under the salt deposit. The formation of the salt deposit at 110 °C promoted the anodic reaction, resulting in the formation of an aggressive wet interface...
between the deposit and the surface of the specimens, leading to crevice corrosion, preferential attack of the ferrite phase, pit formation and SCC nucleation and growth.

- The tensile stress of 139 MPa was above the SCC threshold stress for the ASS at 110 °C. The tensile stress values of 473 MPa (DSS) and 589 MPa (SDSS) were above the SCC threshold stresses at 110 °C;

- SCC propagation of ASS and DSS was mainly transgranularly. DSS specimen also showed preferential crack ramification inside the austenite phase, while

- SCC of DSS and SDSS specimens preferentially propagated at 45° of the rolling direction (parallel to the maximum shear stress), while the SCC of ASS propagated normal to the rolling direction.

Figure 14. (a) Scheme showing the relative position of the cathodic (dripping zone) and the anodic reactions (under the salt deposit); (b) Temperature distribution along the upper surface of the specimen at 110 °C. The temperature in the center of the dripping region is roughly 10 °C lower than the testing temperature. Points located at -25 mm and +25 mm represent the usual SCC nucleation sites.
SDSS specimen additionally presented crack propagation along the ferrite/austenite interfaces.

- Transgranular cleavage microfractography was observed for all SSs tested at 110 °C. DSS also featured areas of flat facets, indicating ferrite/austenite interfacial brittle fracture, while SDSS also featured intergranular brittle fracture.

- SDSS specimens showed the lowest SCC susceptibility during DET at 110 °C in seawater, followed by DSS and finally by ASS specimens.

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