Effect of Tungsten on the Pitting and Crevice Corrosion Resistance of Type 25Cr Super Duplex Stainless Steels

Eirik B. Haugan,* Monika Næss,* Cristian Torres Rodriguez,* Roy Johnsen,* and Mariano Iannuzzi‡,*,**

ABSTRACT

The oil and gas industry regularly uses Type 25Cr super duplex stainless steels (SDSS) for components exposed to seawater and hydrocarbon environments in topside facilities, downhole, and subsea equipment. Much debate still exists concerning the effect of tungsten on pitting and crevice corrosion resistance, particularly in standardization committees. Whereas some researchers claim that tungsten has a strong synergistic effect with molybdenum when added above a certain threshold value, others argue that tungsten additions at the expense of molybdenum could lower corrosion resistance. The objective of this investigation was to examine the effect of tungsten on localized corrosion of two super duplex stainless steels: a low-W (modified UNS S32750) and a high-W (UNS S39274) grade. Both crevice-free and creviced samples were studied. Tests were conducted in 3.5 wt% NaCl or natural seawater with temperatures ranging from 20°C to 95°C. Various independent methodologies including cyclic potentiodynamic polarization, electrochemical critical pitting temperature testing per ASTM G150, and long-term open-circuit potential exposure in natural seawater were used. Results showed that, in the solution annealed condition, tungsten additions to super duplex stainless steels had a marked positive effect on pitting and crevice corrosion resistance, increasing critical crevice temperatures by as much as 30°C. These findings suggested that tungsten-containing SDSS had a corrosion resistance on par with super austenitic stainless steel grades containing 6 wt% molybdenum. A new parametric definition of the pitting resistant equivalent is proposed to reflect the alloy’s localized corrosion resistance and to support standardization efforts in the materials oil and gas community.

KEY WORDS: critical crevice temperature, cyclic potentiodynamic polarization, oil and gas, seawater, super duplex stainless steels

INTRODUCTION

Duplex and super duplex stainless steels (DSS and SDSS, respectively) are steels composed of a two-phase ferritic-austenitic microstructure, the components of which are both stainless, i.e., they contain more than 10.5 wt% to 12 wt% chromium. Although the ferrite content of DSS and SDSS can vary between 35 vol% and 55 vol%, manufacturers balance the steels close to the ideal 50-50 ferrite-to-austenite ratio. DSS are ferritic-austenitic stainless steels with 22 wt% Cr and have a corrosion resistance on par with austenitic grades of similar Cr content. Examples of DSS include UNS S32205 and UNS S31803. In contrast, SDSS are defined based not only on their chromium content but also on the alloy’s pitting resistant equivalent (PRE). In this regard, the PRE is an empirical formula that attempts to correlate the complex beneficial effect of the main alloying elements using a simple compositionally derived “pitting index.” While NORSOK M-001 defines PRE based on Cr, Mo, and N
(Equation [1]).\textsuperscript{4,7} ISO 21457\textsuperscript{5} includes tungsten in the PRE expression (Equation [2]).

\[
\text{PRE}_N = \text{Cr} + 3.3 \times \text{Mo} + 16 \times \text{N} \quad (1)
\]

\[
\text{PRE}_{N,W} = \text{Cr} + 3.3 \times (\text{Mo} + 0.5 \times W) + 16 \times \text{N} \quad (2)
\]

In Equations (1) and (2), the sub-index "N" indicates that the original PRE expression suggested by Lorentz and Medawar\textsuperscript{8} was modified to include nitrogen, while the sub-index "W" in Equation (2) shows that the PRE formula also includes tungsten.\textsuperscript{9} In Equations (1) and (2), all values are given in wt%.

SDSS are, thus, defined as ferritic-austenitic stainless steels with 25 wt% Cr and a PRE \( \geq 40 \).\textsuperscript{4-5,7} The high Cr, Mo, and N content makes SDSS resistant to most oxidizing environments,\textsuperscript{10-12} but are only considered seawater resistant by NORSOK M-001 and ISO 21457 up to 20°C because of crevice corrosion concerns.\textsuperscript{13-14} The most common SDSS for subsea connection systems, piping, and tubing are UNS S32750, UNS S32550, and UNS S32760, which are all treated as equivalent\textsuperscript{13} in NORSOK M-001,\textsuperscript{4} NORSOK M-630,\textsuperscript{7} and ISO 21457.\textsuperscript{5} Table 1 summarizes the nominal composition of the most common DSS and SDSS used in oil and gas production.

DSS and SDSS combine high strength and localized and stress corrosion cracking (SCC) resistance at a competitive cost given the relatively small Ni content when compared to austenitic stainless steels.\textsuperscript{3} Table 2 compares mechanical properties of DSS and SDSS vs. selected austenitic stainless steels and nickel-based alloys to illustrate the advantages of DSS and SDSS over other alloys commonly used in the oil and gas industry.

At the time of this writing, there is much debate in the oil and gas community in Norway and in ISO standardization committees as to whether W, a ferrite stabilizer,\textsuperscript{15} has a beneficial effect on localized corrosion resistance of SDSS. There is still no agreement on whether the PRE definition should include W in future revisions of the NORSOK M-001 specification.\textsuperscript{4} Opponents to the inclusion of W and alignment with ISO 21457\textsuperscript{5} argue that manufacturers could add W at the expense of Cr and Mo, creating some concerns regarding the seawater resistance of the resulting material. This reasoning, however, appears unjustified solely on a simple price comparison between the main alloying elements. W, a rare metal primarily used for high-temperature applications, is more expensive than both Cr and Mo;\textsuperscript{16-18} tungsten additions are, therefore, intentional rather than a cost-saving strategy.

Although the effect of W on localized corrosion resistance has not been studied to the same extent as other alloying elements such as Cr, Mo, and N,\textsuperscript{11} researchers agree that there exists an optimal W concentration; outside this range, W is either ineffective or detrimental.\textsuperscript{19-20} Tomashov, et al.,\textsuperscript{21} were the first to investigate the effect of W additions to an austenitic 18 wt% Cr-14 wt% Ni alloy. They reported that tungsten had a minor influence on localized corrosion resistance in 0.1 M NaCl at room temperature when compared to, e.g., Mo; however, tungsten had a net beneficial effect at around 3 wt%. Anh, et al., examined the influence of W and Mo on pitting potentials of laboratory-grade Fe-29 wt% Cr alloys.\textsuperscript{22} The authors reported a linear relationship between pitting potentials (E_p) measured in 4 M MgCl₂ at 80°C and the alloy’s W and Mo contents. When normalized to their corresponding atomic weights, the authors concluded that W and Mo increased E_p in almost the same amount.

Kim and Kwon\textsuperscript{19} showed that a W-to-Mo weight ratio of 2 gave the best localized corrosion resistance. A W-to-Mo ratio of 2 contrasts, for example, with the W-to-Mo weight ratio of UNS S39274 and UNS S32760, which are approximately 0.70 and 0.25, respectively. Ogawa and coworkers\textsuperscript{20} found a maximum in pitting corrosion resistance at 2 wt% W, both in the base metal and the heat affected zone (HAZ) of a welded SDSS, but with a W-to-Mo weight ratio of 0.72, in line

\begin{table}[h]
\centering
\caption{Nominal Composition of Typical DSS and SDSS\textsuperscript{47}}
\begin{tabular}{|c|c|c|c|c|c|}
\hline
UNS & Cr (wt\%) & Mo (wt\%) & Ni (wt\%) & N (wt\%) & Other (wt\%) \\
\hline
S32205 & 22 to 23 & 3.0 to 3.5 & 4.5 to 5.6 & 0.14 to 0.20 & Mn\textless{}1.2, S\textless{}0.001, P\textless{}0.030, C\textless{}0.020, Si\textless{}0.8 \\
S32750 & 24 to 26 & 3.0 to 5.0 & 6.0 to 8.0 & 0.24 to 0.32 & Mn\textless{}1.2, S\textless{}0.015, P\textless{}0.035, C\textless{}0.030, Si\textless{}0.8 \\
S32550 & 24 to 27 & 2.9 to 3.9 & 4.5 to 6.5 & 0.10 to 0.25 & Cu\textless{}1.50-2.50, Mn\textless{}1.5, S\textless{}0.03, P\textless{}0.04, C\textless{}0.030, Si\textless{}1.0 \\
S32760 & 24 to 26 & 3.0 to 4.0 & 6.0 to 8.0 & 0.20 to 0.30 & W: 0.5-1.0, Cu: 0.5-1.0, Mn\textless{}1.0, S\textless{}0.01, P\textless{}0.03, C\textless{}0.030, Si\textless{}1.0 \\
\hline
\end{tabular}
\end{table}

\begin{table}[h]
\centering
\caption{Mechanical Properties and CPT and CCT of Conventional Stainless Steels and Nickel Alloys\textsuperscript{66-67,75}}
\begin{tabular}{|c|c|c|c|c|c|}
\hline
Property & 22Cr DSS & 25Cr SDSS & UNS S31603 & UNS S31254 & UNS N06625 & UNS N07752 \\
\hline
SMYS or R_p (0.2\% offset) (MPa) & 450 & 550 & 205 & 310 & 415 & 920 \\
Tensile Strength, R_m (MPa) & 620 & 750 & 515 & 675 & 825 & 1,268 \\
Min. Elongation to Failure, A (%) & 25 & 25 & 35 & 35 & 30 & 30 \\
\hline
\end{tabular}
\end{table}
with that of UNS S39274. Although the author did not specify the actual ratio in the article, Szkodarska-Smialowska mentions a particular W-to-Mo ratio in which a synergistic effect leads to peak corrosion performance.

Kim, et al., have analyzed the effect of tungsten additions to various austenitic and duplex laboratory-grade stainless steels, including Type 25Cr SDSS, on repassivation kinetics and SCC resistance in different environments. Whereas Ni had a detrimental effect, results suggested that W additions to a 22.92 wt% Cr-6.18 wt% Ni-2.11 wt% Mo-0.07 wt% W (i.e., a W-to-Mo ratio of 0.033) and to a 17.92 wt% Cr-14.04 wt% Ni-2.05 wt% Mo-4.16 wt% W (i.e., a W-to-Mo ratio of 2.03) alloy improved repassivation kinetics as determined by the scratch test method in 1 M magnesium chloride (MgCl₂) and 1 N sulfuric acid (H₂SO₄) + 3.5% Cl. Tungsten in solid solution also seemed to improve SCC resistance noticeably in boiling 35% MgCl₂.

It is still unclear whether W is enriched in the passive film as WO₃ or if dissolved W anions (i.e., tungstate, WO₄²⁻) inhibit the electrolyte inside pits and crevices. Bui and coworkers studied the effect of tungsten additions to a base 16 wt% Cr-14 wt% Ni alloy, as well as the influence of dissolved tungstate. They concluded that both W additions to the base alloy and dissolved tungstate ions increased pitting potential and reduced the critical current for passivation in 0.2 M HCl. Additionally, the authors suggested that the direct formation of WO₃ at the surface in neutral NaCl solutions was responsible for the improved localized corrosion resistance. Kim, et al., proposed that W contributed to the stability of the passive film, leading to a decrease in the critical current density to reach passivation in simulated pit-like solutions, as well as to an increase in pitting potential in neutral NaCl electrolytes. Kim, et al., attributed the improved passivity to a surface Mo accumulation in W-containing stainless steels, which was determined by auger emission spectroscopy.

Independently of the effect of W in solid solution, researchers disagree about the possible retardation or acceleration of the precipitation kinetics of deleterious intermetallic compounds and tertiary phases during, e.g., welding operations. Ogawa, et al., Kim, et al., and Kim and Kwon, and, more recently, Park and Lee studied the precipitation kinetics in DSS without and with W additions. The authors concluded that, during welding or isothermal heat treatments, W retards the formation of σ-phase for the less detrimental χ₁-phase and possibly Cr₂N in the HAZ. Ogawa, et al., showed that W additions above 2 wt% lowered impact toughness and localized corrosion resistance. Kim and Kwon also investigated the effect of isothermal heat treatments at 850°C as a function of different W-to-Mo ratios. In accordance with Ogawa, et al., the authors reported that W retarded the formation of σ-phase, favoring χ₁-phase precipitation. A 3 wt% W-1.5 wt% Mo alloy showed the highest resistance to embrittlement induced by aging. This alloy also gave the best SCC resistance in boiling MgCl₂ and localized corrosion resistance to chloride-containing electrolytes. Similarly, Park and Lee found that substituting, in part, Mo with W retarded σ-phase precipitation in the HAZ. Tungsten-containing weldments had better pitting corrosion resistance than those containing exclusively Mo, with an optimal composition of 2.2 wt% Mo-2.2 wt% W.

Jeon, et al., recently studied the retardation of σ-phase precipitation when substituting Mo for W on a hyper-duplex stainless steel, i.e., a DSS with a PRE₅₅₃₅₃≥ 50. The authors reported that W strongly favored the precipitation of χ₁-phase, improving the overall pitting corrosion resistance. They supported their findings based on both thermodynamic modeling and experimental measurements. Moreover, the authors showed that adding W to the base alloy not only retarded σ-phase formation but also reduced the total volume fraction of tertiary phases. The authors proposed that the preferential precipitation of the χ₁-phase during the early stages of aging depletes Mo and W along grain boundaries, reducing the driving force for σ-phase formation. The main criticism to Jeon, et al., is that the authors focused on relatively long, i.e., more than 600 s, temperature holding times, which are not representative of the temperature profiles experienced during welding. A similar criticism can be made about the investigation by Kim and Kwon discussed earlier.

In clear contrast, work on Mo- and W-alloyed SDSS weld metals by Nilsson, et al., and computational simulations by Wessman and Pettersson suggested that partial substitution of Mo by W caused a more rapid growth of intermetallic phases. Nilsson, et al., concluded that high-W (i.e., 2.16 wt% W) SDSS had a faster σ-phase precipitation kinetics than W-free and low-W counterparts. Likewise, Wessman and Pettersson indicated that partial substitution of Ni by copper (Cu) appeared to retard σ-phase formation but could accelerate chromium nitride precipitation rates. In both studies, the investigators focused on short, i.e., less than 60 s, temperature holding times, which may better represent the transformations occurring during welding.

**Electrochemical Techniques in Localized Corrosion Research**

Electrochemical techniques are valuable tools to quantify the effect of microstructure on localized corrosion performance. Anodic cyclic potentiodynamic polarization (CPP) testing can be used to evaluate the pitting and crevice corrosion behavior of an alloy under various metallurgical conditions. During CPP, the working electrode is first scanned forward in the anodic direction at a given scan rate and reverted...
once the current reaches a certain value.\textsuperscript{34} For most stainless steels in halide solutions, CPP provides two main parameters: (i) the pitting potential \( E_p \) (or crevice potential, \( E_{p,\text{crev}} \) if using creviced samples) and (ii) the repassivation potential \( E_{RP} \) (or \( E_{RP,\text{crev}} \) if using creviced specimens).\textsuperscript{35} Whereas \( E_p \) is a measure of the alloy’s resistance to pit initiation, \( E_{RP} \) has been shown to correlate well with the alloy’s overall resistance to localized corrosion.\textsuperscript{36} Dunn, et al.,\textsuperscript{35} and Sridhar and Cragnolino\textsuperscript{36} suggested that, above a certain critical charge density value, \( E_{RP} \) becomes independent of the current density at scan reversal. For UNS S31600 and UNS N08825, \( E_{RP} \) became independent of prior pit growth for deep pits. In this regard, deep pits were associated with a critical charge density of 10 C/cm\(^2\).\textsuperscript{36} Therefore, the authors concluded \( E_{RP} \) could be used as a reliable estimator of localized corrosion resistance when the charge density criterion is met. \( E_{RP} \) has also been used in parametric models to predict long-term corrosion performance.\textsuperscript{34,36-37}

While some studies have investigated the corrosion behavior of SDSS as a function of W content using various electrochemical and immersion techniques, the effects of W on repassivation kinetics, crevice corrosion initiation, and long-term performance remain unclear. The objective of this investigation was to compare the localized corrosion resistance of two SDSS: (i) a conventional low-W grade (i.e., a modified UNS S32750) and (ii) a commercial high-W SDSS (i.e., UNS S39274). The scope of the work was to establish the critical pitting temperature (CPT) and the critical crevice temperature (CCT) using CPP testing, temperature ramping or a fixed anodic potential, and long-term open-circuit potential (OCP) exposure in natural seawater.

**EXPERIMENTAL PROCEDURES**

**Materials**

Samples were cut from (i) a low-W modified UNS S32750 forged bar with an outer diameter of 30 mm, taken from an actual production run used to manufacture blind dowel pins and (ii) a 100 mm by 150 mm S32750 forged bar with an outer diameter of 30 mm, and had an average surface area of 16.9 cm\(^2\). For both crevice-free and creviced tests, samples were cut into 3-mm-thick disks that were 30 mm in diameter and had an average surface area of 16.9 cm\(^2\). A small, 2 mm hole drilled close to the perimeter of the samples served as the sample holder. The test specimens were suspended using a 200 \( \mu \)m platinum wire that acted as an electric connection.

**Sample Preparation**

Samples were polished down to 600 grit SiC paper using ethanol as a lubricant. Samples were subsequently rinsed in acetone, deionized (DI) water, and ethanol and cleaned in an ultrasonic bath for 300 s. A subset of samples was pickled according to NORSOK M-630 recommendations.\textsuperscript{7} Test coupons were immersed in a solution of 20% nitric acid (HNO\(_3\)) and 5% hydrofluoric acid (HF) at 60°C for 300 s, followed by a thorough DI-water rinse. Special safety procedures for handling HF were followed. All samples were stored in a desiccator for 24 h before testing, which is often referred to as “passivation.”\textsuperscript{7,36}

**Cyclic Potentiodynamic Polarization Testing**

CPP testing was conducted on crevice-free and creviced specimens in accordance with ASTM G61.\textsuperscript{35} For creviced samples, a spring-loaded crevice assembly was used.\textsuperscript{39} Flat polytetrafluoroethylene (PTFE) crevice formers were used as described by Steinsmo, et al.,\textsuperscript{40-41} and Høydal, et al.\textsuperscript{42} The titanium bolt was electrically isolated using a heat-shrinkable tube. The crevice assembly was mounted using a torque of 2 N·m. More details can be found elsewhere.\textsuperscript{42}

A deaerated 3.5 wt% NaCl (0.62 M NaCl) pH 8.0 solution was used as electrolyte. The solution pH was left unadjusted and monitored before and after testing. Tests were conducted at eight different temperatures: 25, 30, 40, 50, 60, 70, 80, and 90°C. The temperature was controlled using a regulating hot plate. The actual solution temperature was continuously monitored during testing and kept within ±1°C.

Cyclic anodic polarization curves were obtained using a conventional three-electrode array. A saturated calomel electrode (SCE) was used as reference electrode. The reference electrode was placed in a separate compartment kept at room temperature and connected to the electrochemical cell using a salt bridge. The test solution was purged for 1 h using high-purity nitrogen gas before immersing the samples in the solution. Nitrogen purging was maintained for the duration of the anodic polarization. Upon immersion,

**TABLE 3**

<table>
<thead>
<tr>
<th>UNS</th>
<th>C</th>
<th>Si</th>
<th>Mn</th>
<th>P</th>
<th>S</th>
<th>Cr</th>
<th>Mo</th>
<th>Ni</th>
<th>W</th>
<th>Cu</th>
<th>N</th>
<th>Co</th>
<th>PRE(_N)</th>
<th>PRE(_{N,W})</th>
</tr>
</thead>
<tbody>
<tr>
<td>S32750</td>
<td>0.02</td>
<td>0.32</td>
<td>0.56</td>
<td>0.019</td>
<td>0.0004</td>
<td>25.74</td>
<td>3.31</td>
<td>6.92</td>
<td>0.55</td>
<td>0.20</td>
<td>0.267</td>
<td>&lt;0.05</td>
<td>40.93</td>
<td>41.84</td>
</tr>
<tr>
<td>S39274</td>
<td>0.016</td>
<td>0.20</td>
<td>0.80</td>
<td>0.045</td>
<td>0.003</td>
<td>25.2</td>
<td>3.20</td>
<td>6.40</td>
<td>2.20</td>
<td>0.52</td>
<td>0.28</td>
<td>–</td>
<td>40.24</td>
<td>43.87</td>
</tr>
</tbody>
</table>
samples were left at the corrosion potential ($E_{corr}$) for 1 h before polarization. Samples were subsequently polarized in the anodic direction at a scan rate of 600 mV/h (0.167 mV/s). Scan reversal occurred once the current density reached 5 mA/cm². The test was completed when the hysteresis loop closed or upon reaching OCP.

**ASTM G150—Electrochemical Critical Pitting Temperature**

A subset of crevice-free UNS S32750 and UNS S39274 specimens was used to determine the potential independent CPT using a complimentary approach as described in ASTM G150. Samples were polished to 600 grit paper, pickled, and passivated as outlined in the Materials section. Tests were performed in 3.5 wt% NaCl with a pH = 8.0. Samples were polarized to an applied potential, $E_{App}$, of 600 mV$_{SCE}$ for 5 min before and during temperature ramping. The solution temperature was ramped at a rate of 1°C/min from 20°C to 95°C, or until a maximum current density of 150 μA/cm² was reached, whichever occurred first. The CPT was determined as the temperature at which $i = 100$ μA/cm².

**Long-Term Open-Circuit Exposure**

Creviced UNS S32750 and UNS S39274 specimens were prepared using the method described by Kivisäkk and Novak, Steinsmo, et al., and Heydal, et al., were exposed to filtered natural seawater obtained directly from the Trondheim fjord at NTNU’s seawater laboratory. The main advantage of the spring-loaded crevice assemblies used herein is that no significant drop in applied force has been observed even after long-term testing at elevated temperatures.

The initial temperature was set to 60°C, which was above the CCT determined by CPP for UNS S32750. A temperature of 60°C was also shown to be an adequate choice for SDSS by Johnsen and Vingsand. Both temperature and $E_{corr}$ were monitored during exposure. If no drop in $E_{corr}$ was observed after 750 h, the temperature was increased to 70°C and kept at that temperature for another 400 h or until a sudden decrease in $E_{corr}$ was detected, whichever occurred first. If no drop in $E_{corr}$ was measured after a total exposure time of 1,150 h, then the temperature was increased to 80°C. This process was repeated until crevice corrosion initiation occurred. More details about the experimental setup can be consulted elsewhere.

**Characterization**

A group of samples was first polished and etched following ASTM A923 recommendations to determine the absence of deleterious phases before testing. Test specimens were etched in a 40 wt% sodium hydroxide (NaOH) solution using 1.5 V for 30 s to 40 s. After etching, the test coupons were rinsed in acetone followed by air drying. Samples were then examined using confocal and scanning electron microscopes.

After testing, samples were rinsed in DI-water and stored in a desiccator. Samples were analyzed in the optical microscope to determine the presence of pits and the absence of crevice attack at the connection point. Samples were gently cleaned using a 3-μm diamond suspension to remove corrosion products and reveal sub-surface pits. Creviced samples exposed at $E_{corr}$ were also analyzed with the scanning electron microscope.

**RESULTS**

**Materials Characterization**

Figures 1 and 2 show both optical and secondary electron images of as-received and etched SA UNS S3270 and UNS S39274, respectively. Whereas UNS S32750 had an equiaxed fine grain structure, characteristic of forged products, UNS S39274 had elongated grains aligned in the rolling direction, typical of rolled plates. Both alloys had mechanical properties that exceeded the minimum requirements of ASTM A182 and NORSOK M-630 standards, as well as UNS S32750 and UNS S39274 specifications (Table 4). Charpy V-notch (CVN) and ASTM G48 method A results, as well as optical microscopy, suggested that the material was free of deleterious phases. The volume fraction of austenite was 47 vol%, whereas the volume fraction of ferrite was 53 vol%, as determined by optical microscopy in accordance with ASTM A923 and ferritescope.

Differences in microstructure introduced by the manufacturing process could affect corrosion performance, in particular, resistance to hydrogen stress cracking. Nevertheless, in this work, the localized corrosion resistance was assumed to be primarily controlled by alloy composition, in line with, e.g., the work by Sendriks and Newman. This hypothesis is justified based on the adequate average austenite and ferrite volume fractions, which were close to the ideal 50% in both cases, the lack of deleterious intermetallic compounds and tertiary phases evidenced by optical and scanning electron microscopy (Figures 1 and 2), as well as ASTM G48 method A quality control testing.

**Anodic Polarizations**

The shape of the anodic polarization curves depended on the composition, the microstructure of the test specimen, the type of coupon (i.e., crevice-free or creviced), and the temperature of the solution.
Nevertheless, all polarization plots could be grouped into three distinct cases: (i) curves showing no hysteresis, (ii) curves showing high $E_p$ or $E_{Crev}$ if using crevice formers and little hysteresis, and (iii) curves showing large positive hysteresis loops. Figure 3 illustrates the three cases. The inflection seen in curves

![Image](a)

![Image](b)

![Image](c)

**FIGURE 1.** (a) and (b) Optical and (c) secondary electron images of solution annealed UNS S32750 showing an equiaxed fine grain structure characteristic of forged products, austenite (light) and ferrite (dark). Samples were free of deleterious intermetallic compounds and third phases. Specimens were etched in a 40 wt% NaOH solution using 1.5 V for 30 s to 40 s.

![Image](a)

![Image](b)

![Image](c)

**FIGURE 2.** (a) and (b) Optical and (c) secondary electron images of solution annealed UNS S39274 showing elongated grains aligned in the rolling direction, characteristic of rolled plates, austenite (light) and ferrite (dark). Samples were free of deleterious intermetallic compounds and third phases. Specimens were etched in a 40 wt% NaOH solution using 1.5 V for 30 s to 40 s.

<table>
<thead>
<tr>
<th>TABLE 4</th>
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<tbody>
<tr>
<td>Actual Mechanical Properties of Tested Materials</td>
</tr>
<tr>
<td>UNS</td>
</tr>
<tr>
<td>---------</td>
</tr>
<tr>
<td>S32750</td>
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<tr>
<td>S39274</td>
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showing no hysteresis or high $E_P$ or $E_{Crev}$ and little hysteresis was, primarily, associated with the oxygen evolution reaction caused by water oxidation and transpassive dissolution at the transpassive potential, $E_{Trans}$. The presence of small pits concurrent with oxygen evolution translated into small hysteresis loops. Given that the total current is the sum of pit propagation and water oxidation, it was not possible to discern whether the 10 C/cm² critical charge density criterion proposed by Dunn, et al., was met in those cases. In contrast, curves displaying significant positive hysteresis loops were always associated with pitting or crevice, if using crevice formers, corrosion.

Although pitting corrosion was accompanied by uniform dissolution, especially at higher temperatures, it is reasonable to assume, that, for crevice-free samples, most of the current was associated with pitting corrosion. Integration of the current density vs. time curve confirmed that the deep pit condition was exceeded in all cases. Thus, $E_{RP}$ values corresponded to a lower-bound critical potential that could be used to estimate the conditions for crevice corrosion initiation.

To be consistent with Sridhar and Cragnolino, $E_P$ was determined at the inflection point of the $E$ vs. $i$ plot, while $E_{RP}$ values were identified as the potential, in the backward scan, where the current density reached 2 $\mu$A/cm². Likewise, the passive current density, $i_{pass}$, was measured as the mean current density in the passive region. A similar approach was followed to determine $E_{Crev}$ and $E_{RP,Crev}$ for creviced specimens.

**Critical Pitting Temperature, Protection Temperature, and Critical Crevice Temperature Based on Cyclic Potentiodynamic Polarization Testing**

*Crevice-Free Samples* — CPT values were obtained at the inflection point of $E_{Trans}$ or $E_P$ vs. temperature diagrams. The CPT was calculated as the mean temperature between the temperature of the last $E_{Trans}$ and the first $E_P$ potential. Because the temperature step was 10°C, there was an average ±5°C margin of error. Similarly, the protection temperature ($T_{Prot}$) was taken at the inflection point of $E_{Trans}$ or $E_{RP}$ vs. temperature curves, calculated following the same procedure. Figure 4 illustrates $E_{Trans}$, $E_P$, and $E_{RP}$ vs. temperature maps for UNS S39274, whereas Figure 5 compares average $E_{Trans}$ or $E_{RP}$ potentials as a function of temperature for UNS S32750 and UNS S39274. CPT and $T_{Prot}$ values are summarized in Table 5.

Whereas the CPT of UNS S32750 was 65°C to 75°C for the as-polished condition and 75°C for pickled surfaces, the CPT of UNS S39274 was 85°C regardless of surface condition. There was some degree of uncertainty in establishing CPT for as-polished samples, as the transition was more gradual than in the other cases. Tests in the 60°C to 75°C range using a smaller
temperature interval could give a better approximation. Likewise, the mean TProt of UNS S32750 was 55°C, whereas that of UNS S39274 was approximately 85°C. Because, for any given test condition, the potential dropped from the transpassive region to measurable EP or ERP values at the same temperature, min. and max. CPT and TProt results shown in Table 5 were identical. The only exception was the CPT of UNS S32750 in the polished and passivated case, where the drop in critical potential occurred at different temperatures.

No clear differences were observed between polished and passivated and pickled and passivated samples. Nevertheless, pickled and passivated surfaces led to less scatter. A more detailed discussion on the differences between the two surface preparation methods is presented elsewhere.  

Creviced Samples — To verify the validity of CCT values estimated using TProt on crevice-free specimens, i.e., CCT|ERP, CPP was also conducted on creviced coupons. Given the small difference in EP and ERP between as-polished and passivated and pickled and passivated samples, which was in line with the findings discussed by Szklarska-Smialowska and DeForce, the remaining tests were performed on polished and passivated samples only.

There was no clear inflection in ECrev as a function of temperature. Therefore, no reliable CCT could be inferred from ETrans or ECrev vs. temperature diagrams (Figure 6[a]). Nonetheless, a clear inflection point was observed in the crevice repassivation, ERP, vs. temperature curves (Figure 6[b]). As shown in Table 6, the average critical crevice repassivation temperature

| Microstructure | Surface Finish | CPT (°C), ASTM G61 | Mean TProt or CCT|ERP (°C) ASTM G61 | CPT (°C), ASTM G150 |
|----------------|----------------|--------------------|------------------|-----------------|
|                |                | min. | max. | min. | max. | min. | max. |
| UNS S32750     | Polished (600 grit SiC) | 65   | 75   | 55   |      |      |
| UNS S32750     | Pickled (NORSOK M-630)  | 75   | 75   | 63.5 | 65.2 |
| UNS S39274     | Polished (600 grit SiC) | 85   | 85   | 80   |      |      |
| UNS S39274     | Pickled (NORSOK M-630)  | 85   | 85   | 95   | >99  |

FIGURE 4. (a) ETrans or EP and (b) ETrans or ERP vs. temperature for crevice-free UNS S39274 with pickled surfaces. Lines added to aid visualization. Each data point represents an average of two or three independent CPP tests.

FIGURE 5. CCT|ERP comparison between crevice-free UNS S32750 and UNS S39274 pickled samples showing the direct effect of W on localized corrosion resistance. Lines added to aid visualization. Each data point represents an average of two or three independent CPP tests.

TABLE 5
Summary of CPT and CCT Values as Determined by CPP or ASTM G150 as Indicated
The E_app = 600 mV_SCE value was chosen to simulate the potential reached in chlorinated seawater systems with 0.5 ppmw residual chlorine. A similar potential was used by Johnsen and Vingsand to study the long-term crevice corrosion resistance of SDSS, including UNS S32750, and nickel-based alloys.

Long-Term Open-Circuit Potential Exposure in Natural Seawater

Figure 8 compares E_corr vs. time plots of creviced UNS S32750 and UNS S39274 specimens, respectively. The temperature increase profile is overlapped in the figures. Photomicrographs are also included to illustrate the presence of crevice corrosion after testing, which nucleated at the periphery of the crevice former in all cases. In Figure 8, a sudden drop in E_corr from values reflecting a passive state to active, i.e., negative, potentials determines the initiation of crevice attack. As shown by Newman and Galvele, the crevice (or pitting) potential of the material controls the sharp decrease in E_corr observed during long-term OCP exposure.

UNS S32750 suffered crevice corrosion at 60°C and tests were stopped with no further temperature rise. The induction time, i.e., the time for crevice corrosion initiation, varied between 416 h and 656 h. In contrast, UNS S39274 did not suffer crevice corrosion up to 80°C. Samples were exposed for a total of 1,416 h to 1,626 h before crevice corrosion initiation. The induction time at 80°C was between 312 h and 520 h.

Figure 9 shows the morphology and the extent of the crevice corrosion attack after long-term OCP exposure in natural seawater. As seen in Figure 9, UNS S32750 suffered selective dissolution of the austenite phase, which was later confirmed by EDS analysis. In contrast, the high-W UNS S39274 grade did not present
clear evidence of selective phase dissolution. Additionally, the depth of the crevice corrosion attack observed in UNS S39274 specimens was shallower on average than in UNS S32750. Selective phase dissolution is common in DSS.61-62 The preferential attack depends primarily on the partitioning of alloying elements to austenite or ferrite,61,63-64 which in turn defines the critical localized corrosion potential of the phase (and the local PRE).65 The phase with the lowest critical localized corrosion potential dissolves first and determines the alloy’s overall pitting or crevice potentials.1,10,60-61,63,65

**DISCUSSION**

The CPT of UNS S32750 as determined by CPP testing was 5°C to 15°C lower and 15°C to 24°C lower than CPT values reported by the alloy manufacturer66-67 in 6 wt% and 10 wt% FeCl₃, respectively. Measured CPT values were also 10°C to 32°C lower than CPT values reported by the alloy manufacturer obtained by ASTM G150 testing in 1 M NaCl (Table 2). Likewise, CPT values found by CPP and ASTM G150 testing at 600 mV_SCE were 12°C to 22°C and 22°C to 24°C lower, respectively, than results reported by Deng, et al.58 Deng and coworkers determined a mean CPT of 87.6±0.4°C for UNS S32750 using temperature ramping at a constant applied potential of 750 mV_SCE in 1 M NaCl. CPT values obtained by ASTM G61 testing were, however, in close agreement with Steinsmo, et al.,40 for the same alloy, but evaluated potentiostatically in seawater at 600 mV_Ag/AgCl. The authors reported CPT values for SA UNS S32750 of 76°C,40 which was just 1°C higher than the CPT obtained by CPP, but between 10°C and 13°C higher than those measured herein as per ASTM G150.67 ASTM G61 results were, too, in agreement with Tsapralis, et al.69 Tsapralis and coworkers obtained a CPT = 71°C for UNS S32750 in 10 wt% FeCl₃, determined using zero resistance amperometry (ZRA) at a temperature scan rate of 0.5°C/min. This value was 6°C to 8°C higher than the CPT values obtained herein as per ASTM G150.67
UNS S39274 had a CPT as determined by CPP and ASTM G150 that was between 15°C and 20°C and between 20°C and >30°C, respectively, higher than that of UNS S32750 depending on surface condition and test method. It is also remarkable that one of the specimens did not suffer pitting corrosion during the potentiostatic hold at 600 mV\textsubscript{SCE} exposure even after reaching the boiling point. These results lie within the experimental error reported by Kim and Kwon.\textsuperscript{19} Results were also in close agreement with Ogawa, et al.\textsuperscript{20} Ogawa, et al.,\textsuperscript{20} measured CPT as a function of W content in a 10 wt% FeCl\textsubscript{3} solution at the OCP. Ogawa reported a CPT of at least 80°C for 25 wt% Cr-7 wt% Ni-3 wt% Mo samples containing between 2 wt% and 4.5 wt% W. Ogawa, et al., suggested that the CPT of samples with more than 2 wt% W exceeded the highest temperature used in their investigation, which was close the acceptable temperature limit for testing in a 6 wt% to 10 wt% FeCl\textsubscript{3} electrolyte.\textsuperscript{18,70}

In addition to W, the two materials used in this investigation also differed in their Cu content. The modified UNS S32750 alloy had a Cu content that was about 60% lower than that of UNS S39274. Researchers have found that Cu additions to SDSS improve pitting potential and overall localized corrosion resistance to halide environments.\textsuperscript{61,64} It may be questioned, therefore, whether the difference in Cu could be responsible for the outstanding resistance of UNS S39274. Garfias-Mesias and Sykes\textsuperscript{64} and Garfias-Mesias, et al.,\textsuperscript{61} conducted extensive investigations on Cu-modified SDSS using a variety of electrolytes and electrochemical techniques. The authors have shown that the extent to which Cu improves localized corrosion resistance strongly depends on the partitioning of the main alloying elements during solution annealing. However, the highest CPT measured for a 25.92 wt% Cr-3.19 wt% Mo-0.20 wt% N-1.62 wt% Cu (PRE\textsubscript{N} = 39.65) was 64°C, much lower than the 85°C of UNS S39274. A low-Cu alloy with a PRE\textsubscript{N} = 40.23 wt% and 0.56 wt% Cu had a CPT of 61.8°C. Therefore, it seems reasonable to assume that the difference in localized corrosion performance should be attributed primarily to W.
The Repassivation Potential as an Estimator of Crevice Corrosion Resistance

According to Sridhar and Cragnolino, the ERp becomes independent of prior pit growth only above a certain pit depth. In line with the interpretation first given by Galvele in his seminal publication on localized corrosion, a growing pit can be visualized as a particular form of crevice corrosion. Nevertheless, it is worth mentioning that, for stainless steels, some researchers argue this may be an oversimplified interpretation. Using Galvele’s analogy, the ERp indirectly estimates the ease with which a crevice can grow stable. Given that the deep pit condition was met or exceeded above the inflection point in ERp vs. temperature curves. Tprot was used as an estimator of the CCT. Although ERp and Tprot cannot provide mechanistic information regarding crevice corrosion kinetics, both parameters relate to conditions leading to crevice corrosion initiation. CCT values indirectly estimated by ERp (i.e., CCT|ERp) vs. temperature curves were 15°C to 25°C lower than the CPT of UNS S32750 and independent of surface finish. In contrast, the CCT|ERP of UNS S39274 was virtually identical to CPT and 25°C to 30°C higher than that of UNS S32750, suggesting W favored pit repassivation kinetics as proposed by Bui, et al. and Lumsden and Szklarska-Smialowska (Figure 5). The effect of W on facilitating repassivation was also evidenced by the difference in CCRT between UNS S39274 and UNS S32750, shown in Figure 6.

CCT|ERp values of UNS S32750 were almost 25°C lower than values reported by Deng, et al., using cyclic temperature ramping at an applied potential of 750 mVAg/AgCl in 1 M NaCl, but 5°C higher than reported CCT values obtained by conventional ASTM G48 method D. Nevertheless, results were in reasonable agreement with CCTs reported by Høydal, et al. Høydal and coworkers investigated crevice corrosion of UNS S32760 as a function of temperature using flat PTFE rods as crevice formers. UNS S32760 is an SDSS. The researchers have found excellent agreement between tests conducted using crevice-free assemblies (e.g., PTFE, ceramic, or PTFE-covered ceramic) and torque strongly affect crevice corrosion results. Nevertheless, it is interesting to highlight the close agreement between critical crevice temperatures indirectly estimated by Tprot, i.e., CCT|ERP, and the crevice corrosion resistance observed in long-term OCP exposure. Thus, ERp, as determined by CPP testing on coupons without crevice formers, was a good estimator of crevice corrosion resistance. These findings are also in line with the recent work by Kappes, et al., and Martinez and collaborators on SDSS. The researchers have found excellent agreement between tests conducted using crevice-free specimens exposed to simulated crevice-like solutions following Galvele’s approach and results of potentiodynamic-galvanostatic-potentiodynamic (PD-GS-PD) testing on creviced coupons.

Despite its potential, the use of ERp to gauge crevice corrosion resistance has limitations. For cases where only little hysteresis is observed and when pitting is concurrent with oxygen evolution and transpassive dissolution, the system may not meet the deep pit condition (Figure 3[b]). Therefore, ERp values, as determined by ASTM G61 testing, cannot be used to establish immunity to localized corrosion.
Alternative electrochemical techniques such as the PD-GS-PD and potentiodynamic-potentiostatically-potentiodynamic (PD-PS-PD) methods and ZRA could give valuable information about the influence of test conditions on CPT and CCT.

A Parametric Pitting Resistant Equivalent Definition

UNS S39274 CPT and CCT values were higher than those of UNS S32750 by as much as 30°C (Figures 5, 7, and 8). CPT and CCT values were, in fact, comparable to those of super austenitic stainless steels with a PREN above 43. Yet, UNS S39274 has a specified minimum yield strength (SMYS) that almost doubles that of UNS S31254. The results presented herein suggest that W in solid solution plays a decisive role in increasing localized corrosion resistance when added to SDSS at about 2.2 wt%. Moreover, the higher CCRT measured by CPP on creviced samples also suggested that W additions led to a faster repassivation kinetics than that of low-W or W-free SDSS, results that are in line with the effect of W additions to Ni-based alloys.

Adding W to the PRE formula as detailed in ISO 21457 translates to a PREN,W of 43.87 (Table 3), similar to that of super austenitic stainless steel alloys. Adding W to the PRE formula as detailed in ISO 21457 when W is added within the limits established by, e.g., ASTM A182 for UNS S32750 and a high W-containing UNS S39274. The following conclusions were drawn based on the evidence presented herein:

- When added at about 2.2 wt%, W had a marked beneficial effect on pitting and repassivation potentials, as well as pitting and crevice repassivation kinetics.
- In the solution annealed condition, the CCT as determined by various independent techniques was up to 30°C higher for UNS S39274 than for UNS S32750.
- CCT values as estimated by ERP were in reasonable accord with results of long-term OCP exposure and literature ASTM G48 method D results, suggesting that ERP measured using coupons without crevice formers could be used to estimate crevice corrosion resistance of SDSS.
- Based on the evidence presented herein, it seems reasonable to accept the inclusion of W in the NORSOK M-001 and ISO 21457 PRE formula with a multiplying factor as detailed in ISO 21457 when W is added in the 1.5 wt% to 2.5 wt% range, maintaining a PREN = 40 as seawater resistance threshold for SDSS.

ACKNOWLEDGMENTS

We thank Atle H. Qvale (GE O&G), Alexander Fjeldly (GE O&G), Leif Brattås (GE O&G), Anders Wiktorsson (GE O&G), and Mariano A. Kappes and Ricardo M. Carranza (Instituto Sabato, Buenos Aires, Argentina) for their guidance, support, and technical discussions.

This work was funded by General Electric, Co. (Oil and Gas, Norway) and conducted at NTNU’s Department of Engineering Design and Materials (IPM).

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