EFFECT OF PLASTIC STRAIN ON THE ELECTROCHEMICAL BEHAVIOR OF LEAN DUPLEX UNS S32304 STAINLESS STEEL*

Charles David¹
Fiona Rue²
Florent Krajcarz³
Saghi Saedlou⁴
Virginie Roche⁵
Ricardo Nogueira⁶

Abstract
This work aimed at evaluating the effect of pre-strain of Lean Duplex UNS S32304 on anodic dissolution and hydrogen evolution reaction rates in acidic chloride media. Interrupted tensile testing was employed in order to generate different deformation levels. Potentiodynamic experiments allowed to detect a decrease on the cathodic and anodic reaction rates as a consequence of plastic deformation. A non-linear evolution of corrosion rates was observed, probably as a consequence of the evolution of specific dislocation structures in the material. Strain also caused a slight ennoblement of the ferritic phase, while austenite became a little less noble. These behaviors are analyzed in terms of residual stresses, characterized by X-ray diffraction.

Keywords: Corrosion; Strain; Duplex; Stainless.

¹ PhD student, LEPMI – Université Grenoble-Alpes, APERAM Research Center, Isbergues, France.
² PhD, research engineer, APERAM Research Center, Isbergues, France.
³ PhD, research engineer, APERAM Research Center, Isbergues, France.
⁴ Engineer, Head of Solutions Dept, APERAM Research Center, Isbergues, France.
⁵ PhD, researcher, LEPMI – Université Grenoble-Alpes, Grenoble, France.
⁶ PhD, Professor, LEPMI – Université Grenoble-Alpes, Grenoble, France.
1 INTRODUCTION

Austenitic-ferritic stainless steels, also known as Duplex (DSS), possess high mechanical properties and very good corrosion resistance. Among these steels, Lean Duplex, which contain very few or no molybdenum and less nickel than standard Duplex, are suitable for use in environments where standard Duplex would be over-specified from a corrosion resistance standpoint and thus represent an attractive technical and economic choice.

Given its applications, Lean DSS might undergo plastic strain during forming or in-service use. Although the corrosion behavior of these materials has been well studied, the effect of prior deformation on their corrosion resistance is not well known.

To this day, only a few works on the impact of strain of stainless steel on corrosion resistance have been carried out. The majority of them address pitting corrosion. N. Renton and co-workers [1-2] observed that strain of Super Duplex induced an irregular evolution of pitting corrosion resistance in chloride medium, which they explain by the non-linear development of slip bands on the metal’s surface. L. Peguet et al [3], while studying the effect of cold work on pitting corrosion of ferritic and austenitic stainless steels, observed an increase of the dissolution rate with cold work. Y. Huang et al [4] observed the same trend in a study on austenitic SS. That would be due to an increase of dislocation density in the material after plastic strain, and eventual changes on the corresponding surface morphology.

Even if pitting corrosion is a major concern for stainless steels, these materials are also employed for their resistance to stress corrosion cracking (SCC). Knowing if and how pre-strain would impact on SCC would be important. Such an impact has been observed by S. Yamazaki et al [5] in their study of the effect of prior deformation on the crack growth rates on a nickel-based alloy. The authors pointed out that the evolution of crack growth rate with cold work is not only related to the increasing of stress concentration at the crack tip, but also to the enhancement of the metal’s oxidation kinetics. A high dislocation concentration near grain boundaries would result in a high oxidation rate.

The impact of prior deformation on oxidation rates but also on hydrogen evolution, which may cause embrittlement, has been addressed on different materials. Y. Cheng and co-workers [6-11] have concluded that strain would promote corrosion specially in the plastic region on high-strength steels [11]. That is in agreement with the mechanoelectrochemical interactions theory, which takes into account the activation of new dislocations in the material. In addition to that, their results show that, even if hydrogen evolution is also promoted by plastic strain, the impact on anodic dissolution is greater.

On the other hand, H. Krawiec et al, in their studies on different aluminum and titanium alloys, observe a predominant influence of strain on the cathodic reaction than on anodic dissolution [12-14]. The role of dislocations has been shown to be complex [14], since an increase of the number of dislocations enhances cathodic reaction up to a certain point, and then it decreases due to the formation of specific dislocation structures.

Dislocations have been proved to impact on dissolution of metal and hydrogen production in acidic media [15-20]. M. Sahal et al [15-16] observed that the dissolution rate of nickel in acidic medium depends on the dislocation density, and that taking into account the internal stresses generated by dislocation distribution is essential when correlating strain and anodic dissolution. Large et al’s [17] studies confirmed it, and also proved that hydrostatic stresses play a role on dissolution. Regarding hydrogen
production, El Alami et al [18-19] concluded that dislocations density and structures play a role on the adsorbing sites of the cathodic reaction on nickel. Yet, strain is not suspected to impact on corrosion resistance only in terms of cathodic and anodic kinetics. Residual stress, which can be generated by straining, may cause corrosion potential shifts. Some authors have studied the effect of compressive and tensile stresses on corrosion resistance [20-22]. Globally, tensile stresses would shift the corrosion potential towards less noble values, whereas compressive stress would make materials nobler.

Briefly, for different materials, strain has been shown to impact on anodic dissolution and/or on the cathodic reaction, both of which are substantial parameters when considering Duplex stainless steels. Shifts of corrosion potentials due to residual stresses have also been reported.

This work aims at evaluating the effect of pre-strain of Lean Duplex UNS S32304 on anodic dissolution and hydrogen evolution reaction rates in acidic media. Furthermore, residual stresses are characterized by X-ray diffraction in order to evaluate their role on electrochemical behavior.

2 MATERIAL AND METHODS

An industrial grade of Lean Duplex stainless steel UNS S32304 produced by Aperam was used in this study. It was cold rolled down to a thickness of 1 mm, annealed and pickled. The chemical content of main alloying elements is given in Table 1. Optical images acquired after chemical etching in Beraha's solution (0.27 g potassium metabisulfite, 20 mL HCl, 100 mL water) of the plate’s section (Figure 1a) and surface (Figure 1b) allowed to evaluate the morphology of austenitic (γ) and ferritic (α) zones. The average sectional band width was 1.34 µm for ferrite and 1.07 µm for austenite. On the plate surface, images revealed a phase composition of 47.8% α/52.2% γ. A measurement of the magnetic phase by sigmameter revealed a larger ferritic ratio in the bulk, 53.9% α/46.1% γ.

Table 1. Chemical composition of UNS S32304

<table>
<thead>
<tr>
<th>C ppm</th>
<th>Mn %</th>
<th>P %</th>
<th>N ppm</th>
<th>Si %</th>
<th>Ni %</th>
<th>Cr %</th>
<th>Cu %</th>
<th>Mo %</th>
</tr>
</thead>
<tbody>
<tr>
<td>159</td>
<td>1.52</td>
<td>0.029</td>
<td>917</td>
<td>0.45</td>
<td>4.73</td>
<td>23.43</td>
<td>0.38</td>
<td>0.47</td>
</tr>
</tbody>
</table>

Figure 1. Micrographies of the plate’s section (a) and of the surface (b) after etching
2.1 Interrupted tensile testing

Due to technical constraints of a parallel ongoing study, tensile specimens were rectangular coupons of 100 x 23 mm². In order to ensure that the strain was uniaxial and homogenous in such a type of specimen, in-situ image correlation was carried out. For that purpose, specimens were covered with a white painting layer and then sprinkled with black painting so as to form the correlation dots (Figure 2).

![Figure 2. Tensile testing specimen for image correlation before (a) and after (b) painting application.](image)

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**Figure 3.** Strain maps by in-situ image correlation for 10% and 20% total elongation

**Figure 4.** Strain profiles by in-situ image correlation for 20% total elongation

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The specimen was mounted on the tensile testing machine and centered so as the gage length was 60 mm. A two-camera dispositive was placed in front of the gage section in order to capture images of the specimen during the test. The assembly configuration did not allow the use of an extensometer. The tensile test protocol started by a preload of 2 N. Strain was controlled by the movement of the crosshead, with a constant rate of 0,5 mm/min up to fracture. Images were captured every 22 seconds. For every image that was capture, the correlation software calculated both axial (ε₁) and transverse (ε₂) strains. Figure 3 shows the strain maps for 10% and 20% total elongation. In addition to that, longitudinal and transverse strain profiles were drawn on the axial direction of the specimen at 20% total elongation rate, at the center and near the edges of the specimen (Figure 4).

It was concluded that both axial and transverse strains were sufficiently homogeneous and uniaxial over a 45 mm-length area in the center of the specimen until a 20% total elongation rate. The electrochemical experiments could therefore later be carried out on samples collected from that area.

For the fabrication of the pre-strained samples by tensile interrupted testing, the total elongation levels were: 0.5%, 1%, 3%, 5%, 10% and 20%. The crosshead displacement needed for each one of these elongation levels was known based on the image correlation results, but this time an extensometer was used in order to know the final plastic strain. Specimens were strained using the same protocol that for image correlation. Once the desired total elongation for each specimen was reached, the crosshead’s position was maintained for 5 minutes for deformation to accommodate, and then plastic strain was released down to zero load with a -0,5 mm/min crosshead displacement rate.

2.2 Electrochemical experiments

Two samples of approximately 21 x 23 mm² were collected from the center of each pre-strained specimen. They were then polished down to colloidal silica in order to avoid cold-working due to preparation. Surfaces were checked in SEM after polishing. Electrochemical experiments were carried out in a dual-vessel three-electrode assembly thermostated at 23°C, using an 18 mm O-ring working electrode. The configuration of the system allowed purging the electrochemical media with a 20/80 H₂/N₂ mixture for 1 hour before each experiment without any contact between the media and the sample during purging. For each sample, the electrochemical protocol started with a 5-minute potentiostatic current measurement at -0,6 V/SCE in a 0.07 mol/L H₂SO₄ + 200 g/L NaCl solution. Then, polarization curves were immediately drawn, starting at -0,6 V/SCE towards anodic potentials with a 20 mV/min scanning rate till -0,1 V/SCE.

2.3 Internal stresses measurement

Internal stress measurements were carried out on an X-ray diffractometer using the sin² Ψ method along the axial direction of tensile straining. X-rays were generated by a cobalt tube. The peak at 89.173° was chosen for the study of austenite, and at 99.374° for ferrite, with a scanning range of 6° and a step of 0.06°.
3 RESULTS AND DISCUSSION

3.1 Interrupted tensile testing

Engineering stress-strain curves plotted from the extensometer data (Figure 5) allowed to know the final plastic strain for each pre-strain level (Table 2). It can be noted from the plots that curves are quite stackable, indicating the same strain path. Henceforth, given that the aim of the present study is to correlate plastic strain and corrosion, only the final plastic strain will be used. For practical reasons, numbers will be written under percentage form: 0.4%, 0.9%, 3.3%, 4.3%, 9.4%, 19.3%.

![Figure 5. Engineering stress-strain curves of interrupted tensile testing.](image)

### Table 2. Targeted total elongation and corresponding plastic strain

<table>
<thead>
<tr>
<th>Targeted total elongation</th>
<th>εₚ (extensometer)</th>
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<tbody>
<tr>
<td>0.5%</td>
<td>0.004</td>
</tr>
<tr>
<td>1%</td>
<td>0.009</td>
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<tr>
<td>3%</td>
<td>0.033</td>
</tr>
<tr>
<td>5%</td>
<td>0.043</td>
</tr>
<tr>
<td>10%</td>
<td>0.094</td>
</tr>
<tr>
<td>20%</td>
<td>0.193</td>
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</tbody>
</table>

3.2 Effect of plastic strain on anodic dissolution and hydrogen evolution

Polarization curves show two activity peaks (Figure 6), as it was expected for a DSS. It has already been demonstrated that the peak at more negative potentials corresponds to ferrite, and the more anodic one to austenite [23]. A distinction can be observed between low strained (unstrained to 0.9%), and strongly strained (3.3% to 19.3%) samples, especially in terms of cathodic currents and ferrite's activity peak. The same curves are drawn in Figure 7, but currents are expressed in logarithmic scale. On this graph, the reduction of both cathodic and anodic rates with pre-strain
are highlighted. These current reductions mean that, for a given cathodic potential, hydrogen production decreases with pre-strain. For a given anodic potential, ferrite’s oxidation also decreases with strain. Moreover, strain induces a slight shift of the corrosion potential ($E_{corr}$) towards more positive potentials. Given that at the corrosion potential the oxidation mainly concerns ferrite, this phase is believed to have become nobler with strain. Even if austenite is not the phase being predominantly corroded in this medium, a negative shift of the minima between ferrite’s and austenite’s activity peaks ($E_{min}$) is observed, suggesting that the austenitic phase might have become less noble.

In order to better evaluate the evolution of corrosion rates ($J_{corr}$), they were calculated using the cathodic Tafel rights and plotted against plastic strain on Figure 9a. Corrosion rate first increases with strain for low pre-strain levels, and then decreases and stabilizes for higher strains. That non-linear observation corresponds well to previous studies and could be explained based on the role of dislocations. As raised by H. Krawiec et al and M. Sahal et al [12-16], the increase of the number of dislocations due to plastic strain enhances cathodic reaction to a certain level, and then slows it down due to the formation of specific dislocation structures, which hinders adsorption processes. L. Peguet and Y. Huang et al [3,4] also mentioned an increase of dissolution rates due to an increase of dislocations density.

Figures 9b and 9c show the variations of the corrosion potential and the potential at the beginning of the activity peak of austenite compared to the potentials for the unstrained sample. Although the interpretation of the variation for austenite might be complex, for ferrite (corrosion potential) it appears to correspond to the evolution of the corrosion rates, except for low strains. It is possible that the $E_{corr}$ shift towards more noble potentials is globally a result of a more marked effect of pre-strain on the oxidation reaction of ferrite than on cathodic reduction, as found by X. Ly et al [11].

Residual stress measurements on unstrained and 19.3% pre-strained samples are given on Table 3. Austenite is under tensile stress before straining, and ferrite under compressive stress. After straining, the character of internal stresses in each phase does not change, except for its magnitude that increases. That could also explain the shifts on $E_{corr}$ and $E_{min}$, since compressive stresses are supposed to make materials nobler, and inversely for tensile stresses [20-22]. In Duplex steels, it can be supposed that, given that the two phases do not have the same mechanical behavior, after pre-strain and elastic relaxation the softer phase would retract more than the harder phase, increasing compressive residual stresses in the first one and tensile in the second. Further studies should allow to evaluate the contribution of dislocations and also to better understand the evolution of cathodic and anodic behaviors separately after straining.
Figure 6. Polarization curves of unstrained and pre-strained Lean Duplex UNS 32304 in 0.07M H₂SO₄ + 200 g/L NaCl.

Figure 7. Log-scale polarization curves of unstrained and pre-strained Lean Duplex UNS 32304 in 0.07M H₂SO₄ + 200 g/L NaCl – evolution of currents and potentials.
4 CONCLUSION

A protocol for evaluation of the effect of plastic strain obtained by interrupted tensile testing on the electrochemical behavior of Duplex stainless steels has been set up. Potentiodynamic experiments on seven pre-strain levels allowed to detect a difference on the cathodic and anodic reaction rates as a consequence of plastic deformation. First, corrosion rates increase with strain for the low pre-strain levels. This would be due to the development of new dislocations in the material. For higher deformation, corrosion rates decrease and then stabilize, probably as a consequence of the evolution of specific dislocation structures that hinder adsorption processes.

A slight ennoblement of ferrite has been observed for plastic strain levels going from 3.3% to 19.3%. The opposite behavior was noted for austenite. If that trend is
confirmed, it could be due both to the effect of strain on the anodic dissolution reaction and to the residual stress state in the material.

REFERENCES


