Abstract: The austeno-ferritic Stainless Steels are commonly employed in various applications requiring structural performances with enhanced corrosion resistance. Their characteristics can be worsened if the material is exposed to thermal cycles, since the high-temperature decomposition of ferrite causes the formation of detrimental secondary phases. The Submerged Arc Welding (SAW) process is currently adopted for joining DSS owing to its relatively simple execution, cost savings, and using molten slag and granular flux from protecting the seam of atmospheric gases. However, since it produces high contents of δ-ferrite in the heat affected zone and low content of γ-austenite in the weld, high-Ni filler materials must be employed, to avoid excessive ferritization of the joint. The present work is aimed to study the effect of 3 and 6 hours isothermal heat treatments at 850°C and 900°C in a SAF 2205 DSS welded joint in terms of phases precipitation. The results showed the presence of σ-phase at any time-temperature combination, precipitating at the δ/γ interphases and often accompanied by the presence of χ-phase. However, certain differences in secondary phases amounts were revealed among the different zones constituting the joint, ascribable both to peculiar elements partitioning and to the different morphology pertaining to each microstructure.

Key-words: Duplex Stainless Steels; SAW; Heat treatment; Sigma; Chi.

1. Introduction

Duplex Stainless Steels (DSS) are characterized by an interesting combination of corrosion-resistance and mechanical properties, making them very suitable for structural applications in petrochemical, pharmaceutics and nuclear industries, as well as in off-shore and marine environments. DSS are biphasic steels possessing a 50/50 balanced microstructure composed by almost equal volume fraction of ferrite (δ) and austenite (γ); for this reason, DSS exhibit a partially ferromagnetic behavior together with a good thermal...
conductivity and have a lower thermal expansion coefficient than austenitic steels. Further, the duplex structure assures considerably high strength levels at elevated temperature, good toughness and ductility [1-10].

Nevertheless, the ferritic matrix suffers from a decomposition process during isothermal aging within a critical temperature range, located at 650-950°C, which can cause the precipitation of harmful secondary phases, such as α-phase, χ-phase, carbides and nitrides. These phases are known to be deleterious for the corrosion resistance of DSS, since all of them are considerably enriched in chromium [7-10]. Moreover, these phases represent a solution of continuity in the biphasic structure, acting as preferential sites for crack initiation and, therefore, causing a worsening of the mechanical properties. For this reason, after the hot-working processes, DSS are subjected to a solution-annealing treatment (solubilization) at 1050-1100°C, aimed to restore the balanced (duplex) microstructure and to re-dissolve the eventually formed secondary phases.

In the fabrication of components or equipment, manufacturers frequently employ welding as the principal joining method, which may be followed by an isothermal heat treatment. Such a manufacturing process produces changes in the microstructure that modifies the desired properties. Therefore, since thermal cycles performed within the DSS critical temperature range can cause the precipitation of secondary phases, especial care must be taken during the welding operations. DSS can be joined by all arc welding processes, in which Shielded Metal Arc Welding, Gas Tungsten Arc Welding, Gas Metal Arc Welding and Submerged Arc Welding (SAW) are the most employed. SAW process was developed nearly the 1940 and is well-adapted since then. Their benefits include a molten slag which protects the seam weld from atmospheric contamination, high productivity and high welding quality [11,12]. Is commonly used for joining DSS and previous studies have evidenced that a proper setting of processing parameters in welding the 2205 DSS grade has resulted in a satisfactory δ-γ phase balance in the heat-affected-zone [7-9,13].

However, the microstructure developed after welding is different from that resulting after hot-working in terms of phases morphology and elements partitioning, affecting both the properties of the joint and the precipitation behavior during heat treatments. J. Lou et al. applied in 2205 DSS welds a post weld heat treatment in which their observations conducted to evidence some changes of second phases, segregation and element diffusion improving the microstructure of the heat affected zone (HAZ) of the SAW joints. However, these segregations produced a reduction of the mechanical properties and performance in the DSS joints [14].

Several studies has been performed on phases precipitation in DSS [1-10], most of them concerning the solution heat-treated materials, which have permitted to define the critical temperature range for phases formation and the related loss in properties from the analysis of the Time-Temperature-Precipitation (TTP) curves. In the 2205 grade, the most common secondary phases are the intermetallics α- and χ-phase, whereas carbides and nitrides are less frequent and only visible at very high magnification. In the present paper, the effects of isothermal aging at 850°C and 900°C on the microstructure of a welded SAF 2205 DSS are discussed, focusing on the different behavior of the welded metal respect to the base material after long time treatments (3 and 6 hours).

2. Experimental

The base material was a SAF 2205 DSS, received as plate of 30 mm in thickness and having the composition reported in Table 1. Plates were welded together to form butt joints along the longitudinal direction by adopting the SAW process and employing the ER 2209 as filler metal (composition in Table 1), which contains a higher percentage of Ni respect to the base material in order to counteract an excessive ferritization of the joint.

The isothermal ageing treatments of specimens – previously solubilized at 1050°C – were carried at 850°C and 900°C for 3 and 6 hours and were followed by water quenching. The metallographic sections were prepared for optical (OM) and electron (SEM) microscopy by using standard polishing techniques; for OM observations, the

| Table 1. Chemical compositions of base and filler materials, Fe bal. (wt.%). |
|------------------|--------|--------|--------|--------|--------|--------|--------|
|                  | C      | Cr     | Ni     | Mo     | Mn     | Si     | N      |
| SAF 2205         | 0.019  | 21.9   | 5.7    | 3.0    | 1.5    | 0.5    | 0.16   |
| EN 2209          | 0.020  | 22.5   | 9.0    | 3.0    | 1.2    | 0.7    | 0.15   |
specimens were etched using Murakami solution (30 g KOH, 30 g K$_3$Fe(CN)$_6$ and 60 ml H$_2$O), whereas the SEM investigation was performed on unetched samples.

The volume fraction of the phases was measured by image analysis on the OM and SEM micrographs and the chemical composition of the phases was determined using a Falcon FEI energy dispersive X-ray spectrometer (EDS) attached on SEM; the collected EDS data were quantified using a ZAF correction method on a Quantax software.

### 3. Results and Discussion

The as-received Base Material (BM) consisted of a ferrite matrix containing elongated $\gamma$-grains that were approximately estimated to be the 51% of the total volume fraction (Figure 1a), thus denoting a well-balanced structure. Conversely, the solidification microstructure in the Fusion Zone (FZ) was composed by a lower amount of $\gamma$-phase respect to BM, either in form of dendrites or restructured as Widmanstätten plates, constituting about the 35% of the duplex structure (Figure 1b). In FZ, the observed unbalanced phase ratio was due to the rapid solidification process of the welding joint, which primarily solidified as fully ferritic, whereas austenite was subsequently formed during cooling to room temperature.

![Figure 1. Optical Microscope images of (a) the base material SAF 2205 as-received and (b) FZ/HAZ/BM interfaces.](image)

On the other hand, the Heat Affected Zone (HAZ) adjacent to the fusion line (Figure 1b) presented a further lower amount of $\gamma$-phase if compared to the other zones, where the austenite morphology was altered by the thermal supply due to the SAW process. As expected, the average grain size in FZ and HAZ were increased respect to BM, as a consequence of the high heat input.

Beside the morphological differences, BM- and FZ-phases exhibited different compositions (Table 2), owing to the different thermal histories at which were subjected; indeed, BM was solution-annealed at about 1050°C and then water quenched, while FZ directly derived from cooling the melt. It is well known that solubilization is aimed to obtain partitioning coefficients pointing toward unity, but the usually adopted temperature range (1050–1100°C) determines a greater elements partitioning if compared to melted DSS [1], by following the affinity of each element toward $\delta$- or $\gamma$-phase. In DSS, the solution-annealing temperature is mainly chosen as a function of the alloying elements content, but must be limited in order to maintain the balanced structure, and the higher the temperature the higher the risk to obtain excessive ferrite.

### Table 2. EDS compositions of $\delta$-ferrite and $\gamma$-austenite measured in the base material (wt.%).

<table>
<thead>
<tr>
<th>Zone</th>
<th>Phase</th>
<th>Si</th>
<th>Cr</th>
<th>Mo</th>
<th>Ni</th>
<th>Mn</th>
<th>Fe</th>
</tr>
</thead>
<tbody>
<tr>
<td>BM</td>
<td>$\delta$</td>
<td>0.5</td>
<td>24.5</td>
<td>4.6</td>
<td>3.7</td>
<td>1.2</td>
<td>bal.</td>
</tr>
<tr>
<td></td>
<td>$\gamma$</td>
<td>0.4</td>
<td>19.0</td>
<td>2.2</td>
<td>7.2</td>
<td>1.5</td>
<td>bal.</td>
</tr>
<tr>
<td>FZ</td>
<td>$\delta$</td>
<td>0.5</td>
<td>22.2</td>
<td>2.9</td>
<td>7.7</td>
<td>1.3</td>
<td>bal.</td>
</tr>
<tr>
<td></td>
<td>$\gamma$</td>
<td>0.5</td>
<td>21.7</td>
<td>3.3</td>
<td>8.5</td>
<td>1.4</td>
<td>bal.</td>
</tr>
</tbody>
</table>

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In the material under study, the BM-ferrite was found to be more enriched in Cr and Mo, while Ni and Mn preferentially diffused in austenite; conversely, the elements were more equally distributed within the FZ-phases, due to the reaching of the melting temperature and to the rapid solidification process that characterized the welded material (Table 2).

In 2205 DSS, the nose of the TTP curve is located at about 850°C and, at this temperature, the formation of intermetallics usually requires about 20 minutes of soaking time [3,9]. Therefore, treating the material for more than three hours is expected to cause a copious precipitation of secondary phases and the formation of an increased volume fraction of austenite respect the untreated joints, owing to the eutectoid decomposition of ferrite ($\delta \rightarrow \sigma + \gamma_2$), the presence of secondary austenite can be due to two mechanisms, from the existing austenite or within the ferrite associated with nitrides precipitation [15]. Neither of both mechanisms was presented in this study since $\gamma_2$ needs rapid cooling kinetic formation. BM, HAZ and FZ possessed different microstructures in terms of phase morphologies and elements partitioning and, moreover, FZ can be further split in two distinct areas – an upper (FZ-up) and a lower (FZ-down) zone – owing to heating effects caused by the backhand welding operations.

In Figure 2, the welded microstructures in FZ-up and FZ-down of the sample heat treated at 850°C for 3 hours are reported. In FZ-up, austenite and $\sigma$-phase were found to be the only microstructural constituents, denoting the total decomposition of ferrite in the upper part of the joint. On the contrary, the observations in FZ-down revealed that ferrite was not totally replaced by $\sigma$ and that a not negligible amount of $\chi$-phase was present. On the other hand, HAZ and BM were affected by lower amounts of secondary phases, owing to their different microstructural features respect to FZ (Figure 3).

![Figure 2. SEM images of the sample treated at 850°C for 3 hours: (a) Upper Fusion Zone and (b) Lower Fusion Zone.](image)

![Figure 3. Scanning Electron micrographs of the sample treated at 850°C for 3 hours: (a) Heat Affected Zone and (b) Base Metal.](image)
Considering the presence of σ-phase, similar results were obtained after all isothermal treatments and, even if the estimated volume fractions were different, the same trend was revealed by passing from one zone to another (Figure 4). As expected, the performed treatments caused ferrite decomposition in all the parts of the joints, but the precipitated amounts were conditioned by the characteristic microstructures pertaining to each part.

Figure 4. σ-phase amounts estimated after the heat treatments in the different joint positions.

In FZ, the development of partitioning coefficients almost equal to unity determined a greater metastability of ferrite respect to HAZ and BM, making the phase more prone to secondary phases precipitation. In this zone, ferrite was in any case almost completely replaced by σ and, further, the backhand welding caused a differentiate heating between FZ-down and FZ-up, determining a different metastable situation and resulting in different amount of phase fractions after the treatments.

On the contrary, BM and HAZ were conditioned by other thermal cycles if compared to FZ – solubilization and heating on welding, respectively – which caused a reduced precipitation of σ-phase, owing to a greater elements partitioning between the two duplex phases that lowered the driving force for σ-phase formation.

In BM and HAZ, the σ-phase morphology was found to be different at the two considered temperatures, assuming a coral-like character at 850°C and a bulk shape at 900°C, as also noted by Pohl et al. [10]. In any case, σ-phase nucleated at grain boundaries and grown toward ferrite, and the observed differences in particles shape were due to diffusional/nucleation effects, which conditioned the phases morphology deriving from the eutectoid reaction, causing a greater interconnection between ferrite and σ at lower temperatures (Figure 5). In FZ, owing to the total decomposition of ferrite and to microstructural features, these differences in shape were less evident.

Figure 5. SEM images of the sample heat treated for 6 hours at (a) 850°C and (b) 900°C.
Even though certain regularity among the joint zones was found for σ-phase precipitation, the trends were different if the other intermetallic – the χ-phase – is considered (Figure 6). This secondary phase was mainly observed after the 850°C treatments, since the 900°C temperature is toward the upper limit of the χ-phase stability-field; further, this phase tend to disappear in favor of σ after long thermal exposure, especially at higher temperatures [16]. Treating the joints at 850°C for three hours always caused the formation of greater amounts of χ-phase, which can be considered almost constant in the different welding zones. Conversely, after six hours at 850°C the trend seemed to be inverted respect to σ-phase precipitation and greater volume fractions of χ-phase were observed in BM rather than in FZ. This was due to the tendency of χ to be replaced by σ after long treatment times. The differences among the joint zones can be addressed to the more proneness of FZ-ferrite to σ-phase decomposition, thus causing an anticipation of the precipitation sequence and the formation of lower amount of χ-phase in FZ respect to BM owing to the anticipate reaching of equilibrium conditions. In any cases, χ-phase can be observed as small bright globular particles located along the δ/γ interfaces (Figures 2, 3 and 5).

![Figure 6. χ-phase amounts estimated after the heat treatments in the different joint positions.](image)

### 4. Conclusions

In the present work, the effects of 3 and 6 hours isothermal heat treatments at 850°C and 900°C on a 2205 DSS welding by SAW process were examined. As expected, the employed time-temperature combinations always caused the formation of considerable amounts of σ- and χ-phase, whose volume fractions depended on the considered part of the joint (i.e. base material, BM; heat-affected zone, HAZ; and fusion zone, FZ).

Indeed, all the parts constituting the joints where affected by ferrite decomposition, but the precipitated amounts were conditioned by the characteristic microstructures pertaining to each part. In FZ, owing to the rapid cooling from the melt, the alloying elements were almost equally partitioned within the two duplex phases, allowing for a greater driving force for σ-phase precipitation. On the contrary, HAZ and BM were less affected by σ-phase formation, since they were only subjected to thermal cycles. In these latter parts of the joint, the elements were more partitioned within the phases by following their δ- or γ-character, thus determining a greater stability against ferrite decomposition.

The presence of χ-phase was mainly revealed at 850°C, since at 900°C this phase is not even stable; at this latter temperature, either σ-phase precipitation is more favored or χ is quickly replaced by σ owing to higher diffusional contributions. Both the secondary phases nucleated along the δ/γ interfaces and, while σ-phase grown toward the ferritic matrix and possessed peculiar morphologies depending on the ageing temperature, the χ-phase always precipitated at the boundaries in globular form.
Detrimental Cr-rich Phases Precipitation on SAF 2205 Duplex Stainless Steels Welds After Heat Treatment

References


