Effect of Aging Heat Treatment H950 and H1000 on Mechanical and Pitting Corrosion Properties of UNS S46500 Stainless Steel

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The effect of aging temperature on mechanical and pitting corrosion properties of UNS S46500 was investigated. Tensile and Hardness tests were carried out and the microstructure was analyzed by optical microscopy, scanning electron microscopy and X-ray diffraction; Thermo-Calc simulations helped in the phase identification. Pitting corrosion properties were investigated in 0.6M NaCl electrolyte with sulfate additions by Potentiodynamic Polarization (PP). Hardness, tensile and yield strength of the UNS S46500 steel after lower aging temperature, 510ºC (H950), are higher than the ones found in the 538ºC (H1000) aged steel. This result is explained by microstructure, X-ray diffraction and Thermo-Calc analysis, which indicated the presence of austenite, chi phase and probably Ni$_3$Ti precipitates finely and uniform distributed throughout the martensite matrix. Pitting corrosion resistance is equivalent in both aging temperatures. The sulfate inhibitor effect on UNS S46500 steel was enhanced for 538ºC condition when the electrolyte reaches 1Cl$^-$:1SO$_4^{2-}$ ratio, which is explained by Ni sulfate adsorption and the amount of interfaces in the microstructure resulting in smaller amount of adsorption sites, such as coarsened Ni$_3$Ti precipitates, smaller fraction of chi phase and recovery of dislocations in martensite structure.

Keywords: martensitic precipitation hardening stainless steel, aging heat treatment, pitting corrosion.

1. Introduction

The mechanical properties of precipitation hardening steels can be attributed to the presence of a fine and uniformly distributed intermetallic particles in a martensitic matrix. The UNS S46500 steel is strengthened by precipitation of Ni$_3$Ti. It is known that longer aging times and/or higher aging temperatures imply in growth and coarsening this precipitates, leading to mechanical properties reduction.

The heat treatment of these materials is accomplished by solution heat treatment followed by water or oil quenching. The presence of retained austenite is possible and a quick cryogenic treatment is employed to further reduce or eliminate the fraction of retained austenite. However, when high toughness and high strength material is desired, the presence of low quantities of this phase is important. Schnitzer et al. proposed that austenite is formed during the aging of the material, and this austenite is called reverted austenite, which is formed at the same time of precipitates nucleation in the martensitic matrix, i.e. the austenite stabilizer element, Ni, is present in the precipitate nucleation Ni-X and simultaneously there is its diffusion into less ordered regions, e.g. grain boundaries or martensite lath boundaries, occurring its microsegregation and leading to austenite nucleation.

The precipitation hardening stainless steels have been widely used in aerospace application due to its combination of strength, toughness and corrosion resistance. In this material, the main form of corrosion is the localized corrosion (pit and crevice), where cavities formed act as stress concentrator, which can lead the material to a catastrophic failure.

Pitting corrosion can be studied by many methods, such as potentiodynamic polarization, salt spray, weight loss, always in the presence of chloride ion. In the potentiodynamic polarization test, the pitting potential (Epit) is measured relative to a reference electrode in different environments depending on the purpose of the study. Some electrolytes, when excessively aggressive, do not permit the Epit determination, since they promote general surface corrosion, or present a lack of sensitivity to distinguish the difference between pitting corrosion resistance where its properties are similar. The addition of inhibitors compounds in these electrolytes can improve the sensitivity and allow differentiation in the materials performance. In this investigation was used 0.6M NaCl electrolyte with sulfate additions to decrease the aggressively and try to differentiate the performance of the steels.
Thus, the combination between mechanical and corrosion properties shall be evaluated when these materials are applied. UNS S46500 has as peak strength aging temperature 482°C (900°F). However, this temperature will provide the highest strength but not necessarily the best corrosion resistance. In fact, this aging cycle promotes the sensitivity to stress corrosion cracking. Thus, there are limitations on aging temperatures used for precipitation hardening stainless steels. For the UNS S46500, the suggested aging temperature for optimum strength and corrosion resistance is known as H950 and H1000 according to AMS 5963. Both are carried out for 4 hours, varying only the aging temperature, where for H950 is 510°C (950°F) and for H1000 is 538°C (1000°F).

The purpose of this study is to clarify the effect of aging condition, H950 or H1000, on the mechanical properties and localized corrosion resistance of martensitic precipitation hardening stainless steel UNS S46500.

2. Experimental Procedures

The chemical composition of investigated martensitic precipitation hardening stainless steel is given in Table 1. This material, trade named as Custom 465®, was provided by Carpenter Technology Inc. in the form of a solution heat treated rolled bar with 25.4 mm of diameter, and was then subjected to different aging treatments, 510°C (H950) and 538°C (H1000), both for 4 hours according to AMS 5963. The tests and microstructure examination was done for UNS S46500 on solution heat treated, H950 or H1000 conditions.

Tensile tests were conducted on Wolpert type 20TUZ 750 equipment. For each aging condition, 6 cylindrical specimens with a diameter of 6 mm were tested in the longitudinal direction and tensile elongation values were measured on a 25 mm Panantec gage length according to ASTM E8/E8M. The tensile test at ambient temperature was registered by Panantec ATMI software. Three measurements of Rockwell C (HRC) hardness test, conducted on Pantec RBS, was performed directly on transversal direction of heat treated bars according to ASTM A370.

The specimens were metallographically polished up to 1 µm and then examined with and without Vilcella (1 g picric acid, 5 mL hydrochloric acid, 100 mL ethanol) and Fry (5 g copper(II) chloride, 33 mL hydrochloric acid, 33 mL ethanol, 33 mL distilled water) etching, at room temperature and with immersion times between 15 and 30s.

Metallographic specimens were observed in an optical microscopy (OM), it was also used a scanning electron microscopy (SEM), Jeol JSM-6490LV, generating secondary electron images and energy-dispersive X-ray analysis.

X-ray diffraction patterns were obtained using Cu-Kα radiation (wavelength of 1.5418 nm), at 30 kV and current of 30mA at Shimadzu XRD-7000 diffraction equipment. Scan rate of 2°/min e scan step of de 0.04° were used.

Potentiodynamic polarization tests were conducted in a plane cell (leaving a working area of 1 cm²) using potentiostat/galvanostat 273A Princeton Applied Research and 352 SoftCorr III software. The working surface of disks with 24 mm diameter and 6 mm height was ground subsequently up to 600 grit SiC paper, cleaned with distilled water and alcohol, and dried in cold air. The test solution was 0.6M NaCl electrolyte with Na₂SO₄ additions (0.06M; 0.3M; 0.6M) to decrease the aggressiveness of the electrolytes, allowing a better differentiation of the heat treatment conditions’ performance. The steel specimen was used as working electrode, platinum wire with 1 mm diameter and 25 mm long as counter electrode and a silver/silver chloride (Ag/AgCl) electrode as reference electrode. The specimen was cathodically polarized at 250 mV, followed by potential scan, with scan rate of 1 mV.s⁻¹, up to current density of 10⁻³ A.cm⁻². Pitting potential (Epit) was read directly on polarization curve at the potential which the anodic current increased abruptly. The electrochemical tests, performed at room temperature (23 ± 2 ºC), were repeated 10 times for each heat treatment condition. Immediately after the polarization, sample surfaces were washed with distilled water and then with ethyl alcohol (C₂H₅OH), and finally dried with hot blown air and observed in optical and scanning electron microscopy in order to analyze the corroded surface morphology.

3. Results

Table 2 shows the average results of mechanical properties. For both aging conditions, H950 and H1000, the tensile strength, yield strength, elongation (EL) and hardness values are above the typical specifications of AMS 5963, which are 47 HRC of hardness, yield strength of 1515 MPa, tensile strength of 1655 MPa and total elongation of 10%. It is possible to observe that increasing the aging temperature leads to the reduction of the tensile and yield strengths, and a slight tendency to decrease of hardness, while elongation increased. This result is in accordance with previous works and Narendra.

| Treatment | Tensile Strength (MPa) | Yield Strength (MPa) | EL (%) | Hardness (HRC) |
|-----------|------------------------|----------------------|--------|----------------|
| H950      | 1881 ± 21              | 1836 ± 22            | 10 ± 0.5 | 52 ± 1        |
| H1000     | 1702 ± 14              | 1676 ± 15            | 12 ± 0.9 | 49 ± 3        |

Table 1. Chemical composition (wt.%) of the investigated UNS S46500.

| C    | Mn | Si | P   | S    | Cr | Ni | Mo | Ti | N   | Fe |
|------|----|----|-----|------|----|----|----|----|-----|----|
| 0.005| 0.02 | 0.04 | 0.003 | 0.0005 | 11.63 | 11.02 | 0.94 | 1.59 | 0.002 | Balance |
et al.\textsuperscript{13}, the first authors aged the steels at 550°C, 580°C and 620°C reporting maximum hardness at 550°C (33.1 HRC) and minimum at 620°C (28 HRC). On the other hand, Narendra et al.\textsuperscript{13} carry out aging heat treatments from 510°C (H950) to 570°C with intervals of 20°C and several time conditions, it was report that specimen aging heat treated at 510°C during 2h had maximum hardness but the values were not informed in that paper. A recent review paper reports hardness value of 49 HRC for the H950 condition\textsuperscript{4}.

The microstructure analysis of polished specimens, solution heat treated, H950 and H1000 revealed the presence of disperse titanium nitride particles in the metallic matrix. Figure 1 and Table 3 show the morphology of titanium nitride (TiN) particles and the EDS results of particle and matrix of steel on H950 aging condition confirming that the particle is enriched with titanium.

Figure 2 shows the optical micrographs of Vilella etched specimens. As observed in the Lee et al.\textsuperscript{14} work, the main microstructural feature consists of martensite lath packets with different orientation within relatively equiaxed prior austenite grains. The titanium nitride precipitates observed in the polish specimens where also observed after etching.

Investigations by SEM (Figures 3-5) show more clearly the microstructural differences of three heat treated conditions (solution heat treated, H950 and H1000). The evaluation of the same heat treated condition, it is possible to observe that the different metallographic etched of solution specimens did not revealed differences in the quantity and distribution of precipitates and phases. However, for the aging specimens, H950 and H1000, the Vilella etch revealed more pronounced the martensite lath, white phases and grain boundaries.

The comparison between the three heat treated conditions (solution heat treated, H950 and H1000) revealed increased quantity of white phase (Figures 3 to 5) for aged conditions than solution heat treated specimens; and the Figure 6 show that H950 present higher fraction of white phases than for H1000 condition. The EDS of solution heat treated specimen revealed that the white phase is enriched by Mo and Ti elements (Figure 7 and Table 4). It was not possible see the white phase using the backscattered electrons detector (by chemical composition), only with the secondary electrons detector which gives information about the morphology of matrix. Complementing the microscopy analysis, Figure 8 shows the X-ray diffraction patterns of three heat treatment condition, solution heat treated, H950 and H1000. According the XRD analysis all heat treatment conditions indicate the presence of body-centered cubic phase which represents the martensite (\(\alpha\)) matrix. Equilibrium phase simulation for the aged conditions (Table 5) predicted the formation of chi (\(\chi\)) phase in both aging conditions, but the small fractions of expected equilibrium chi phase, and its diminue dimensions resulted from the precipitation process, did not allowed its identification by XRD. In the same way, TiN observed in the OM and SEM images were not found in the X-ray diffractions, and this can be atributted to lower fraction of nitrogen (0.002%) on the steel chemical composition, leading to very small volume fractions of TiN, not detectable by X-ray diffraction.

Thermodynamic simulations in Thermo-Calc software using TCFE8 database was performed; whereas the aging treatments H950 and H1000 occurring at 510°C and 538°C respectively, the equilibrium phases and its volume fractions at those temperatures are presented in Table 5, and some considerations about this simulation can be addressed:

- BCC phase, body centered cubic ferrite, is the martensite matrix structure.
- Ni,Ti precipitates are responsible for strength the material. They were not observed on OM, SEM and X-ray diffraction analysis, since their very small size, however the literature\textsuperscript{2,15} indicates their presence. The tensile mechanical properties of the UNS S46500 can be attributed to a precipitation of a fine and disperse distribution of these precipitates in the martensite matrix. The identification of Ni,Ti is possible with specialized research and major microscopy resolution as transmission electron microscopy (TEM).
- Chi phase, according to studies\textsuperscript {16}, is enriched with Fe, Cr and Mo (Fe\(_{0.96}\),Cr\(_{12}\),Mo\(_{10}\))\textsuperscript {16}, and was observed in the microscopy analysis for both aging conditions.

Table 3. Chemical composition of matrix and precipitate of Figure 1.

| Elements (wt.%) | Cr   | Ni   | Ti  | N   | Fe   |
|---------------|------|------|-----|-----|------|
| Matrix        | 11.74| 12.32| 1.81| -   | 74.13|
| Precipitates  | -    | -    | 81.93| 13.79| 4.29 |
FCC phase, face centered cubic austenite, were not observed on OM and SEM analysis, however it is mentioned in the literature works.

The quantity analysis of Thermo-Calc indicated that the main difference between H950 and H1000 aging condition is in the quantity of austenite and chi phase, as observed in the Table 5.

Thus, the Thermo-Calc analysis and OM, SEM, EDS and X-ray diffraction tests results indicate that the UNS S46500 steel in the H950 and H1000 aging present a martensite matrix with Ni,Ti and chi precipitates, and some austenite phase.

Figure 2 show typical polarization curves of UNS S46500 in each heat treatment condition. The polarization tests results of solution heat treated material was used as corrosion reference for the material in the aging conditions.

The polarization curves show a cathodic region followed by passive region under all heat treated conditions, and the Epit appear defined in all polarization curves. It is also observed that when the sulfate addition achieves 1:1 of Cl:SO\textsuperscript{2-}, the Epit marked increased compared with other electrolytes concentrations.

The average and deviation of Epit obtained on polarization curves are indicated at Figure 10. The analysis of Epit for both aging conditions indicated that the Epit increased with SO\textsuperscript{2-} addition, which indicates that the pitting corrosion resistance increased. This behavior is expected since the literature indicates the inhibitor effect of this ion for stainless steels. It is also observed that the inhibitor effect of SO\textsuperscript{2-} for UNS S46500 steel in H1000 condition is enhanced gradually with increased SO\textsuperscript{2-} additions, allowing it to reach the highest Epit for this condition when the electrolyte is in proportion to 1Cl: 1 SO\textsuperscript{2-}.

Analyzing the polarization curves in terms of current density (Figure 9), it is clear that the material in both conditions are in the passive state, showing current densities between 10\textsuperscript{-5} and 10\textsuperscript{-6} A/cm\textsuperscript{2}, nevertheless the evaluated conditions using only 0.6M NaCl always showed slightly higher current density before reach the Ep indicating that this chloride concentration affects the passive layer more pronouncedly.

It is known that pitting corrosion can be evaluated by mean of several parameters, among these are, Ep, pit depth, pits density and pit mouth morphology. In this work, it was used the parameters of Ep and pit mouth morphology.
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Figure 3. SEM images of solution heat treated condition, etched with: (a) Vilella and (b) Fry.

Figure 4. SEM images of H950, etched with: (a) Vilella and (b) Fry.

Figure 5. SEM images of H1000, etched with: (a) Vilella and (b) Fry.

Figure 6. SEM images of (a) H950 and (b) H1000, etched with Vilella, showing the quantity difference of white precipitates.
(Figure 11) to infer that different heat treatments did not alter the pitting mechanism, explaining the equivalent pitting potential for the different heat treatment conditions in the same electrolyte.

4. Discussion

In this study, the effects of aging temperatures, H950 and H1000, on mechanical and corrosion properties of UNS S46500 steel were investigated. The mechanical tests results showed that the tensile strength and elongation of the material is strongly depend to heat treatment applied.

The Thermo-Calc simulation indicated that higher aging temperature produced higher quantity of austenite phase: it was found 0.8% for H950 and 2.9% for H1000 (Table 5). This result is in accordance with Schnitzer et al.\[3,20\] and Nakagawa and Miyazaki\[21\] studies.

Austenite acts reducing the tensile strength and increasing the elongation of the material.

The chi phase precipitation was observed for both aging treatments applied for UNS S46500, and the Thermo-Calc analysis indicated that its presence is slight lower for H1000 condition (2.3% for H950 and 1.7% for H1000), in accordance to the observations of this study (Figure 6). The chi phase appeared in the form of relatively coarse precipitates in intergranular and transgranular grain boundaries. Thus,
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Table 4. Chemical composition of matrix and precipitate of Figure 7.

| Elements (wt.%) | Cr   | Ni   | Ti   | Mo  | Fe   |
|----------------|------|------|------|-----|------|
| Matrix         | 11.81| 11.85| 2.12 | -   | 74.11|
| Precipitate    | 10.63| 4.67 | 16.13| 20.85| 47.74|

Table 5. Phases determined by Thermo-Calc (TCFE8 database).

| Phases                  | 510°C, H950 condition (% vol.) | 538°C, H1000 condition (% vol.) |
|-------------------------|--------------------------------|---------------------------------|
| BCC                     | 86.2                           | 84.3                            |
| Ni₃Ti                   | 7.2                            | 7.1                             |
| CHI                     | 2.3                            | 1.7                             |
| FCC                     | 4.3                            | 6.9                             |

The reduction of 2.3% to 1.7% of this phase can contribute to tensile properties reduction. Here the evidence of chi phase is only reported based on termodinamic simulations, nevertheless it was found in literature a research with low carbon martensitic stainless steel and similar Cr and Mo content reporting the presence of Cr and Mo rich precipitates by TEM. This is an evidence that chi phase can appear in this kind of material when the material is exposed to aging temperatures.

Another factor evaluated is the presence of fine and well distributed Ni₃Ti precipitates in the martensite matrix. As these intermetallic precipitates interact with dislocations, they are the main strengthening mechanism in precipitation hardening steels. Schnitzer et al., in their study for PH 13-8Mo, with nickel aluminum precipitates, calculated that 40% of the loss in yield strength during aging can be attributed to the influence of higher amounts of reverted austenite and the rest of the observed decrease in strength is due to coarsening nickel aluminum precipitates. In fact, the Thermo-Calc (Table 5) results showed that amount of Ni₃Ti precipitates for UNS S46500 did not change significantly with aging temperature (7.2% for H950 and 7.1% for H1000). However, considering that the aging time is identical for the two heat-treatment conditions, in the H1000 condition there is higher temperature for coarsening of precipitates; coarsened Ni₃Ti precipitates and higher austenite content can be the reasons for the lower mechanical resistance of the H1000 condition.

Thus, the results indicated that mechanical properties of H950 and H1000 aging conditions can be attributed to higher
amount of austenite, diminish of chi phase and coarsening of Ni₃Ti precipitates.

On the other hand, to evaluate the effect of aging temperature on corrosion resistance, studies 23-24 suggest that material with higher amount of chi phase, as the H950 condition, has a lower resistance to pitting corrosion, since smaller amounts of chromium and molybdenum would be in solid solution. However, the pitting corrosion resistance was very close for all heat treatment conditions studied: solution heat treated, H950 and H1000. Only the sulfate inhibitory effect was more pronounced for the material in the H1000 condition. The mechanism of inhibition of pit nucleation depends upon adsorption of the sulfate ion in the active sites that would initially be occupied by chloride ion, leading to formation of pits. Whereas the chi phase precipitation consumes chromium and molybdenum, leaving the matrix enriched with other elements, particularly nickel, the H1000 condition differs of H950 by having a matrix richer in this element. Seeking in the literature 25, the values of free energy of formation (ΔGº) for NiCl₂ and NiSO₄ compounds are -259.4 and -760.9 kJ. mol⁻¹, respectively, which show the chemical affinity between these species. The higher thermodynamic potential for the formation of nickel sulfate compared to the nickel chloride suggests that the adsorption of sulfate is more severe as higher nickel contents are found in solid solution. In this case, this is the condition of the aging heat treatment H1000, resulting in a greater inhibitory effect sulfate for this condition. It is possible to think in the enrichment and depletion of other elements such as Fe, Cr, Mo and Ti. However, thermodynamic data are not found to allow comparison of affinity of these elements with Cl⁻ and SO₄²⁻, which is probably owes his lack of chemical affinity.

Another factor that may have contribute to the higher Epit provided with 1Cl⁻: 1SO₄²⁻ for H1000 may be related to the amount of interface area that act as adsorption sites. The condition H1000 has higher aging temperature, which has as a consequence higher recovery of dislocations in a martensitic structure, coarsened Ni₃Ti phase and smaller fraction of chi phase. These microstructural changes ultimately reduce the number of interfaces in the microstructure, resulting in smaller amount of adsorption sites.

5. Conclusions

The lower aging temperature (H950 condition) studied in UNS S46500 steel presented greater tensile and yield strengths and hardness due to the presence of minor amounts of austenite, and greater amount of chi phase and Ni₃Ti precipitates finely and uniform distributed throughout the martensite matrix. The two aging conditions, H950 and H1000, have the same pitting corrosion resistance, and the UNS S46500 has excellent response to pitting nucleation inhibition with increasing additions of sulfate.

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