# Effect of Nitrogen on the Corrosion Behavior of Austenitic Stainless Steel in Chloride Solutions

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# Abstract

The effect of partial replacement of nickel with nitrogen on the corrosion resistance of newly designed austenitic stainless steel samples without and with heat treated was investigated in 3.5wt% and 5wt% NaCl solution using open-circuit, potentiodynamic, cyclic anodic polarization and electrochemical impedance spectroscopy techniques. The results showed that, passivation in sample 1 where the highest addition of nickel and low addition of nitrogen is different from that for sample 4 where the nitrogen is greatest and the nickel is reduced almost to the third comparing sample 1. The difference in responses of heat treated samples to localized and general corrosion could be attributed to the difference in their phase compositions. The appearance of ferrite phase for samples (2, 4, 5 and 6) after heat treatment resulted in lowering the general and localized corrosion resistance than as forged samples in contrast with samples 1 and 3, where they still pure austenite. The obtained results are confirmed by surface examination.

Keywords: localized corrosion, pitting corrosion, anodic polarization, nitrogen stainless steel

# 1. Introduction

Nitrogen is considered as an important alloying addition to austenitic stainless steel in terms of corrosion resistance. It promotes passivity, widens the passive range in which pitting is less probable, improves stress corrosion cracking resistance in some media, and enhances the resistance to general and intergranular corrosion (Osozawa & Okato, 1976, p. 135)(Newman, Lu, Bandy & Clayton, 1984, p.394)(Mozhi, Nishimoto, Wilde & Clark, 1986, p.197)(Speidel, 1989, p. 251)(Kamachi, 1993). Moreover, nitrogen dissolved in austenitic stainless steel was found to increase its strength(Osozawa, Okato, Fukase, Yokota & Boshoku-Gijyutsu, 1975. p.1)(Qian, Zhang, Qiu & Guo, 2011, p.4065). The following mechanisms have been suggested to explain how nitrogen operates: (1) nitrogen in solid solution is dissolved and produces  $NH_4^+$ , depressing oxidation inside a pit (Osozawa & Okato, 1976, p. 135) (Osozawa, Okato, Fukase, Yokota & Boshoku-Gijvutsu, 1975, p.1) (Fermenia, Levgraf & Lookkonen, 2001, p.1939)(Bandy, Lu, Newman & Clayton, 1984, p.471); (2) concentrated nitrogen at the passive film/alloy surface stabilizes the film, and prevents attack of anions (Cl<sup>-</sup>)(Clayton, Rosenzweig, Oversluizen & Lu, 1986, p.323)(Halada, Claton, Kim & Kearns, 1995, p.531)(Osozawa & Okato, 1976, p.135)(Newman, Lu, Bandy & Clayton, 1984, p.394); (3) produced nitrate ions improve the resistance to pitting corrosion(Baba, Kodama & Katada, 2002, p.2393); (4) nitrogen addition stabilizes the austenitic phase(Jargelivs, 1986); and (5) nitrogen blocks the kink, and controls the increase of electric current for pit production(Olesson, 1995, p.467). In order to develop a resources saving stainless steel with excellent general and localized corrosion resistance, 2 key technologies were adopted. The first key is high nitrogen alloying. Nitrogen enrichment improves pitting and crevice corrosion resistance without increasing chromium or molybdenum content. Therefore, adding nitrogen may contribute to reduce necessary chromium and molybdenum content. Moreover, as nitrogen is austenite former, adding nitrogen lowers the nickel content required for forming a single austenitic phase. Lowering the nickel in stainless steel lowers the costs of production. The aim of this paper is to study the effect of partial replacement of nickel with nitrogen on the corrosion behavior of newly designed austenitic stainless steel samples without and with heat treated.

# 2. Experimental

The chemical composition of the samples is described in Table 1. The newly designed samples were prepared, cold worked and normalized by steel laboratory in Central Metallurgical Research and Development Institute (CMRDI). The newly austenitic stainless steel samples forged as square rode at 1100  $^{\circ}$ C. Heat treatment of six samples alloys was carried out at temperature 1050  $^{\circ}$ C, and then isothermally held for 30 minutes followed by water quenching. Then stress relief for 2 hours at 750-770  $^{\circ}$ C followed finally by air cooling. Heat treatment cycle were shown in Figure(1) and the condition after heat treated are listed in Table (2). Austenitic stainless steel (sample 1) was used as the blank material for all the samples. In samples 2- 6 the nitrogen content was varied while the nickel content was gradually decreased.

Experiments were designed to investigate the role of partial replacement of nickel by nitrogen on the corrosion behavior of austenitic stainless steel. The solutions used in this investigation were 3.5 and 5wt% NaCl solution. Open-circuit, potentiodynamic, cyclic anodic polarization and electrochemical impedance spectroscopy were used to test corrosion behavior of the samples in different solutions at 25  $^{\circ}$ C . Finally scanning electron microscope (SEM) and energy dispersive X-ray analysis (EDAX) was used to detect the morphology of some samples surface. Volta Lab 40(PGZ301)-Radiometer computerized potentiostat was used for testing.

# 3. Results and Discussion

## 3.1 Open–Circuit Potential Measurements

## a. Forged samples

Figures(2,3) show the time dependence open circuit potentials of stainless steel samples in 3.5 and 5 wt % NaCl solutions. The curves showed general tendency for the initial open circuit potential values to shift towards more negatives.

From the values of immersion potential  $(E_{im})$  and steady state potential  $(E_{s.s})$  which presented collectively in Tables 3, it is concluded that:

- Sample 4 which contains the highest amount of nitrogen was having more positive E<sub>im</sub> and E<sub>s.s</sub> values indicating passive film formation on the sample surface.
- In contrast, curves indicated that, sample 2, due to the lowest percentage of nitrogen and the highest percentage of carbon, showed the most negative E<sub>im</sub> and E<sub>s.s</sub> values.
- Sample 1 with the highest percentage of nickel showed the most positive E<sub>im</sub> and E<sub>s.s</sub> values which facilitate passivation and producing more stable passive film.
- For samples 3, 5 and 6, E<sub>im</sub> and E<sub>s.s</sub> values shifted to more positive with increasing in nitrogen content in tested samples.

The passive film acts as barrier, separating the sample surface from the corrosive ions in the environment. Thus the understanding of the corrosion resistance of the stainless steel lies in comprehending the properties of the passive film formed on the stainless steel. The passive film formed on high nitrogen stainless steel surface in NaCl solutions was n-type semiconductor (Qian, Zhang, Qiu & Guo, 2011, p.4065).

#### b. Heat Treated Samples:

Figure(4) shows typical comparison between open–circuit potential measurements for as forged and heat treated test alloys in 3.5 wt% NaCl solution.

Analysis of the curves Figure(4) show that, the open-circuit potential of sample 1 after heat treatment had a tendency to shift in positive direction, but as forged was almost stable at low negative value, this results indicating that, the heat treatment improved the corrosion resistance of sample 1. While other samples showed potential drop to the negative direction for both the as–forged and heat treated samples indicating dissolution of the oxide film formed on the surface of these samples. Open-circuit potential for heat treated samples increases than the as forged for samples, 2 and 3, decrease for sample 4 and 6 or almost the same in samples 5.

# 3.2 Potentiodynamic Polarization Measurements

# a. Forged samples:

Figures(5,6) represent the potentiodynamic polarization measurements of stainless steel samples in aerated 3.5 and 5wt% NaCl solution using scanning rate 2mV/sec in potential range from -500 to 2000 mV(SCE) at 25  $^{\circ}$ C. According to Figs(5,6), for six samples in 3.5 and 5wt% NaCl solutions, the polarization curves can be put into two groups. A first group with samples 2, 3 and 5 show anodic dissolution, the Cl<sup>-</sup> ions prevent passivation to occur. A second group with samples 1, 4 and 6 there is a general tendency for passivation. The break in current

as the potential increases explains this passivation. The passivation current was taken as the average current of the passive region.

The superior behavior of samples 1 and 4 over the rest samples was confirmed from the corrosion kinetic parameter which listed in Table (4), where:

For sample 1, where the highest Ni content  $\sim 9$  wt% and low nitrogen content  $\sim 0.028$  wt%, the zero current potential, Ecorr, shifted to more positive value 2.3mV than the other samples and passivation current is estimated to be about 0.06mA. The passivation region extended to higher potential which is shifted to more positive. The break in this passive region occurred at almost 0.8 V.

- For sample 4, where the highest N content ~0.24 wt% and Ni content reducing to the third comparing sample 1, the passivation occurred by a slight decrease in current followed by passivation region started at 0.01mA and extended to more positive higher potential. The break down potential is observed at ~ 1.25V.
- 2) For samples from 2 to 6 the corrosion current density, I<sub>corr</sub>, and the corrosion rate are decreased with increasing in the nitrogen contents without respect to the percentage of Ni.

Increasing in NaCl concentration to 5wt%, Figure(6), lead to decrease in break down potential and shrinkage in passive region for samples 1 and 4 in comparison with that in 3.5wt% NaCl solution. As clear from Table(5), for six samples, the corrosion current density, Icorr, was shifted to higher values which suggested worse behavior of all samples by increasing in NaCl concentration. In addition the results reveal that the value of I<sub>corr</sub> depends also on the composition of the alloys at a given NaCl concentration.

A general view to the potentiodynamic curves of Figure(5,6) and corrosion parameters of Tables (4,5), reveals that nitrogen promote passivation of nitrogen stainless steel and passivation requires a certain proportion of nickel to be present.

It is worthy to observe that the mode of passivation in sample 1 where there is a high addition of nickel and low addition of nitrogen is different from that for sample 4 where the nitrogen is greatest and the nickel is reduced almost to the third comparing sample 1. In sample1, anodic current is almost the lowest. In sample 4, the current decreases greatly with increasing the potential and passivation takes place over a wide range of potential.

It could be possible that the corrosion property was improved in sample1, mainly due to the high nickel contents which led to an increase in the coarseness of the grain size and a corresponding decrease in grain boundaries. Since local corrosion events usually start at electrochemical active site such as grain boundaries (Fermenia, Leygraf & Lookkonen, 2001, p.1939), it is therefore believed that lower corrosion rate which observed in sample 1 was due to lower amounts of electrochemical active site occurring at these grain boundaries. The highest nickel content 9.05% would have been expected to make sample 1 the most resistance to uniform corrosion.

In terms of passive film properties for sample 4, N has been observed to decrease the passive current density. It is also widely admitted that N accelerates repassivation, both for Fe-N alloys (Bandy, Lu, Newman & Clayton, 1984, p.471) (Clayton, Rosenzweig, Oversluizen & Lu, 1986, p.323) (Halada, Claton, Kim & Kearns, 1995, p.531) and for stainless steels (Osozawa & Okato, 1976, p.135) (Newman, Lu, Bandy & Clayton, 1984, p.394), particular in presence of chloride. Scratch repassivation experiments have shown that N dissolving in steel has a strong capacity to promote repassivation (Baba, Kodama & Katada, 2002, p.2393). Upon dissolution of the steels, mainly  $NH_4^+$  is formed from the dissolved nitrogen. Most probably nitrogen from the metal surface, segregated in a negatively charged state, is directly converted according to the chemical reaction:

$$N^{3-} + 4H^+ \longrightarrow NH_4^+$$

The formation of ammonium ions consumes protons and there by increases the pH which buffering the solution in the passive layer region (Jargelivs, 1986) (Olesson, 1995, p.467).

#### b. Heat Treated Samples:

Figure(7) show typical comparison between potentidynamic polarization measurements for as forged and heat treated test samples in 3.5wt% NaCl solution.

The difference in responses of theses alloys to general corrosion could be attributed to the difference in their phase compositions. The appearance of ferrite phase for samples (2, 4, 5 and 6) after heat treatment result in lowering the general corrosion resistance than as forged samples in contrast with samples 1 and 3, where they still pure austenite. In heat treated samples (2, 4, 5 and 6), the ferrite phase would be anodic towards the austenite one which might possibly result in selective dissolution of the ferrite phase leading to lowering the general corrosion resistance of these samples. So while the ferrite phase increases, there is a gradual decrease in corrosion resistance of these samples. This was also confirmed by previous work of Yau and Stericher(Yau &

Streicher, 1988, p. 220), Cortie and Potgieter(Cortie & Potgieter, 1991, p. 2173)and from the electrochemical parameters, Table 6. Samples 1 and 3 have lower corrosion current densities, Icorr hence lower corrosion rate than as forged but the heat treated samples 2, 4, 5 and 6, have higher I<sub>corr</sub> values than as forged ones.

3.3 Potentiodynamic Cyclic Anodic Polarization Measurements

# a. Forged Samples:

Figures (8,9) represent the potentiodynamic cyclic anodic polarization measurements for six samples in 3.5wt% and 5wt% sodium chloride solutions using scan rate 2mV/sec in potential range from -500 to 2000 mV/sec at 25 °C.

As is clear from Figures(8, 9), after the corrosion potential, the anodic current density starts to increase to form the active region. The increase of the potential in the positive directions lead to increasing of the anodic current, which corresponds to oxidation of iron to iron ions, with increasing the potential a passive film  $[Fe(OH)_2, Fe_3O_4]$  and or  $Fe_2O_3$  conform(M.Poubaix, 1974). When the potential move towards more anodic, the current density starts to increase again, forming the transpassive region before oxygen evolution. The current rises suddenly without any sign of oxygen evolution for samples 2, 3, 5 and 6, denoting break down of the passive layer, when the polarization potential reaches a certain critical potential ( $E_{pit}$ ).

Hysteresis loop area is a rough measure of the intensity of pitting corrosion. Larger size means greater intensity of pitting corrosion (Bikic, Rizvanovic and Avdusinovic, 1973, p. 471).

For samples 1 and 4 in 3.5wt% NaCl solution, the cyclic polarization curves don't have a hysteresis loop where the forwards scan current density are higher than back words at the same potential, indicating the good stability and self repairing ability of the passive film. For the rest samples, the hysteresis loops were observed.

As shown from Figure (8), the hysteresis loop area is increased with decreasing in the nitrogen content of the sample reaching its maximum area in sample 2 where the lowest nitrogen content sample.

As noted, it is the formation of the chromium – rich oxide layer that accounts chiefly for the corrosion resistance of stainless steel. However, this layer is susceptible to damage, particularly in the presence of chlorides, and such damage can lead to the onset of localized corrosion such as pitting and crevice corrosion. Nickel does not influence the initiation phase but is important in reducing the rate at which both pitting and crevice corrosion propagates. This is critical in determining how serious corrosion will be and this explain the highly pitting resistance of sample 1(Okatmot, 1973, p. 471).

For sample 4, the improvement in the pitting resistance is due to the highest nitrogen content. Nitrogen in the stainless steel improves pitting and crevice corrosion resistance by the ways consumption of the acid in the pit nuclei leading to  $NH_4^+$  formation (Palit, Kain & Gadyiyar, 1993, p. 979), enrichment of nitrogen on the passivated surface and enrichment of nitrogen on the active surface (Byrives, Grujcic and Owen, 1987, p. 1853)( Newman & Shahrabi, 1987, p. 827). Thus the oxide rupture strain should increase due to better stability and improved characteristics of the passive film resulting in a reduction in the passive film breakage frequency.

Regarding the manganese effect the addition of this element didn't modify significant corrosion behavior of the stainless steels, due to the low ability of Mn(II) to form in soluble compound(Bautista, Blanco, Velasco, Gutierrez, Soriano, Palomares & Takenouti, 2009, p.785). Moreover, the increase in the manganese content lead to increasing in manganese oxide inclusions which creates more pit initiation sites in the alloy (Pardo, Merino, Coy, Viejo, Arrabal & Matykina, 2008, p. 780). Thus, for sample 2, the largest hysteresis loop area may be due to the high manganese content, 8.38%, in addition to the lowest nitrogen content, 0.026%.

As the concentration of sodium chloride increased to 5wt% Figure(9),the potentiodynamiccyclic anodic polarization curves indicated the presence of pitting or crevice corrosion break down potential which sustained the increase of anodic current density. Samples 1 and 4 failed facing rising in sodium chloride concentration and according to Fig (9), for all stainless steel samples the occurrence of the hysteresis loop is related to the presence of aggressive solution formed inside the occluded area which maintain when the potential was reversed.

As clear from the corrosion parameters, Tables (7,8), for samples (2 to 6) the corrosion current,  $I_{corr.}$ , decreased and pitting potential,  $E_{pit}$ , increased with increasing nitrogen contents. Moreover, an increase in NaCl concentration shift  $E_{pit}$  of all samples towards more negative (active) direction, indicating a decrease of pitting corrosion resistance. A linear dependence coexists between  $E_{pit}$  and log concentration of Cl- ions, satisfying the following equation:

$$E_{pit} = a - b \log C_{Cl}$$

Where (a) and (b) are constants depend upon the composition of the alloys. The values of Epit at any given  $Cl^-$  concentration depend upon chemical composition of the samples.

The data reveal that, the value of Epit for the samples (at constant NaCl concentration) follow the sequence Sample:

$$1 \approx 4 > 6 > 3 > 5 > 2$$

This sequence reflects the beneficial effects of nitrogen in the alloy. Element has beneficial effects on pitting resistance shift Epit to more positive potentials. It is clear that higher content of N (sample 4) not only reduce susceptibility to pit nucleation, but also diminish rate of pit growth.

#### **B.** Heat Treated Samples:

Potentiodynamic cyclic anodic polarization measurements for the comparison between as forged and heat treated samples in 3.5 wt % NaCl solution are shown in Figure 10.

From Figure 10, the following facts can be summarizing:

• The heat treatment for sample 1 and 4 did not affect the cyclic polarization curves where still no hysteresis loop as the forward scan current densities are higher than back words at the same potential which indicating good stability of the passive film after heat treatment.

Sample 1 is still pure austenite after heat treatment with the highest Ni content leading to considerable improvement of the pitting properties. Since nickel is less readily oxidized than iron and chromium, there could be enrichment of nickel in its metallic state in the metal closet to the oxide, metal interface, there by prolonging the passive region (Potgieter, Olubambi, Cornish, Machio & El- Sayed, 2008, p. 2572).

For sample 4 although ferrite phase where appeared after heat treatment, but, the highest nitrogen content of this sample act as strong austenite ( $\chi$ ) stabilizer, makes it possible to retain austenite phase with reduction in ferrite phase. And, due to the moderate nickel content in this sample, 3.38 wt%, it could be possible that the pitting resistance after heat treatment mainly due to the effect of Ni on the volume fraction of ferrite and austenite. The reduction of ferrite phase urged chromium atoms moved into austenite phase and there by increased the relative amount of Cr in austenite phase, which meant that the pitting resistance of austenite phase (Da-Wei, Change- Sheng, Xiang-juan, Jun, Lu-lu & Xue-Shan, 2012, p. 50).

The heat treatment for the rest samples (2, 3, 5, 6,) has deteriorated effect on their pitting resistance comparing as forged samples.
Ha et al (Ha, Thand & Sj, 2012, p.80), investigated the pitting corrosion resistance of nitrogen stainless steel

and showed that, the combination of N and C addition enhances pitting corrosion resistance of multigen stanless steer protection passive film, however, the pitting corrosion resistance deteriorated by lamellar  $Cr_2N$  precipitate formed during aging at 900 °C. Table 9, represent the electrochemical parameters  $E_{pit}$ ,  $E_{corr}$ ,  $I_{corr}$  and corrosion rate for the heat treated six samples. Data from Table 9 shows the superior behavior of heat treated of samples 1 and 4 comparing as forged one and vies versa the rest samples.

3.4 Electrochemical Impedance Spectroscopy (EIS)

# **Forged Samples:**

The analysis of Nyquist plots from experimental data is done by using circuit in Figure (11) in which ( $R_s$ ) represents the electrolyte resistance; ( $R_{ct}$ ) represents the charge transfer resistance and the constant phase element ( $C_{PE}$ ).

The impedance spectra of six stainless steel samples in 3.5% and 5wt% of sodium chloride solutions were investigated at open circuit potentials at room temperature. The impedance data are presented as Nyquist plot for six stainless steel alloys in Figures (12,13).

In Figure(12), in case of sample 1 and 4 which contain the highest percentage of nickel and nitrogen respectively. The capacitive loops for two samples were broad, indicating better film stability. In general, the samples surface is covered with continuous oxide film (Qiao, Zheng, Ke & Okafor, 2009, p.979).

The Nyquist plots of stainless steel samples 2, 3, 5 and 6 in Figure(12) show a depressed semicircular shape. This observation indicates that the corrosion of stainless steel in 3.5% NaCl solution is mainly controlled by a charge transfer process. The Nyquist plots are not perfect semicircles and generally attributed to the frequency dispersion and mass transport resistant as well as electrode surface heterogeneity resulting from surface roughness, impurities, dislocations, grain boundaries and formation of porous layer (Machnikova, Pazderova, Bazzaoui & Hackerman, 2008, p.1543).

Figure (13) represent the Nyquist plots for all tested stainless steel samples in 5% NaCl solution. Inspection of

these curves shows that, all spectra exhibited a similar behavior except sample 1.

Sample 1 shows a straight line with a high magnitude of impedance, which indicated the presence of a highly resistant passive film. This confirm that the high percentage of nickel facilitate passivation and produce more stable passive film.

Clearly all the impedance spectra for other samples 2, 3, 4, 5 and 6 exhibit one single depressed semicircle, which indicates that the corrosion of stainless steel is mainly controlled by a charge transfer process, and usually related to the charge transfer of the corrosion and double layer behavior. These capacitive loops are not perfect semicircles which can be attributed to the frequency dispersion effect as a result of the roughness and in homogeneousness of electrode surface. The diameter of the capacitive loop depends on the composition of the test samples.

The main parameters deduced from the analysis of Nyquist diagram which reported in Tables (10,11) are:

- The resistance of charge transfer R<sub>ct</sub> (diameter of high frequency semicircle)
- The capacity of double layer C<sub>dl</sub> which is defined as:

$$C_{dl} = 1 / 2\pi f_{max} R_{ct}$$

Where  $f_{max}$  is the maximum frequency. c- The degree of surface coverage ( $\theta$ ) is calculated from the EIS data. From the impedance data given in Table (10,11), we conclude that:

• The value of Rct increases in the order sample

1>4>6>3>5>2

this can be attributed to the formation of protective film on alloy / solution interface.

On the other hand, the high Rct values are generally associated with lower corroding sample (Machnikova,

Pazderova, Bazzaoui & Hackerman, 2008, p.1543).

• The value of double layer capacitance Cdl decreases in the order:

sample 2>5>3>6>4>1. This attributed to increase in thickness of electric double layer, suggested that samples 1 and 4 give a better protection against the corrosion. This agreement with the results obtained from the other electrochemical techniques.

#### a. Heat Treated Samples:

Nyquist impedance plots obtained for the comparison between as forged and heat treatment alloys of stainless steel in 3.5 wt% NaCl, Figure 14, showed single semicircle shifted along the real impedance ( $Z_{real}$ ) axis. For both as forged and heat treatment the shape of figure almost the same with the small increasing or decreasing the values of real impedance( $Z_{real}$ ) and frequency. The result were interpreted as mentioned before.

## 3.5 Spectroscopic Surface Examination

#### a. Scanning Electron Microscope (SEM):

Figures (15,16) illustrates the scanning electron microscope images which describe the morphology of samples no. 1, 2 and 4 after potentiodynamic polarization in 3.5 wt% NaCl solutions. Figures (15(a,b)) show no evidence of corrosion attack and formation of passive film on alloy surfaces. This observation indicates that samples no. 1 and 4 gave excellent corrosion resistance due to the protective passive film with the thickness of a few nanometers formed on the surface of the samples. While Figure(15,c) which belong to the alloy no.2 in 3.5wt% NaCl solution showed that, the surface of the samples was highly corroded. The corroded areas are shown as grooves in the sample with gray and white products.

Figures (16,a-c) showed the scanning electron microscope images which describe the morphology of the surface of the tested samples no. 1,2 and 4 after potentiodynamic polarization measurements in 3.5 wt% NaCl solution after heat treated. As can be seen the samples no 1 and 4 were free of any corrosion attack, and possesses a single austenite phase and two phases austenite and ferrite respectively. But sample no .2, exhibit decreasing in the level of corrosion than before heat treatment.

The (SEM) results are in good agreement with the electrochemical measurements results.

# b. Energy Dispersive X-Ray Analysis (EDAX):

Figure(17) show the energy dispersive X-ray analysis (EDAX) for samples no. 1, 2 and 4 in 3.5wt% NaCl solutions at room temperature.

The observed spectra indicated the presence of Fe, Cr, Ni, O, C, S and Cl ions. According to the peak strength Fe and Cr dominates the component of the film, though the peak of the nitrogen is not appear so nitrogen peak

intensity is independent of the nitrogen content (Qiao, Zheng, Ke & Okafor, 2009, p.979). The presence of chloride ion on the passive film is known to create a concentration gradient that facilitates its diffusion into the film, where it undergoes hydrolysis and reduces the local pH, causing dissolution of the film (YQiao, YZheng, Ke & Okafor, 2009, p.979).

#### 4. Conclusion

- The general and localized corrosion attack was found to be retarded by nitrogen addition and the samples were able to passivate as the nitrogen content is increased.
- Addition of nitrogen to stainless steel promotes the passivation but passivation requires a certain proportion of nickel to be present.
- The difference in responses of heat treated samples to general and localized corrosion could be attributed to the difference in their phase compositions. So while the ferrite phase increases, there is a gradual decrease in corrosion resistance of these samples.

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#### Appendix

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Table 1. The chemical composition of the samples used, mass %

Sample #	Cr	Ni	Ν	Mn	Si	V	С	Р	S	Fe
1	18.01	9.05	0.028	0.92	1.75	.0115	0.15	0.016	0.020	bal.
2	19.1	5.51	0.026	8.38	1.16	0.106	0.24	0.022	0.028	bal.
3	20.18	5.36	0.222	6.37	2.0	0.103	0.22	0.015	0.025	bal.
4	20.1	3.38	0.240	7.03	2.6	0.103	0.19	0.022	0.016	bal.
5	20.2	2.76	0.030	9.89	1.08	0.106	0.15	0.022	0.011	bal.
6	20.7	0.16	0.235	4.04	2.7	0.060	0.13	0.030	0.015	bal.

Table 2. Phases of the heat treated stainless steel samples

Sample No.	Phases
1	Pure austenite
2	Austenite +ferrite
3	Pure austenite
4	Austenite +ferrite
5	Austenite+ferrite
6	Austenite+ferrite

Table 3. Values of Eim and Ess for samples in different concentrations of NaCl solutions

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Conc.	Sample No.	NaCl	
		$E_{im}(V)$	$E_{s.s}(V)$
	1	-0.136	-0.180
%	2	-0.446	-0.577
₹	3	-0.337	-0.425
3.5	4	-0.138	-0.380
	5	-0.419	-0.465
	6	-0.402	-0.423
	1	-0.353	-0.393
	2	-0.415	-0.590
t%	3	-0.425	-0.481
ă In	4	-0.133	-0.431
u /	5	-0.375	-0.486
	6	-0.401	-0.433

Table 4. The corrosion parameters of six samples in 3.5wt% NaCl, solution

Sample	E <sub>corr</sub>	I <sub>corr</sub>	Tafel	Slopes	<b>Corrosion rate</b>
No.	(mV)	μA/cm <sup>2</sup>	β <sub>a</sub> n	nVβ <sub>c</sub>	μm/y
1	2.30	9.06	119.60	-154.60	106.00
2	-393.30	47.50	158.60	-620.30	555.20
3	-209.80	35.72	201.70	-314.90	417.80
4	-168.70	10.54	120.00	-148.70	123.30
5	-310.50	36.21	232.10	-166.80	423.60

6	201 50	12.00	126 10	44.00	151.00
0	-201.30	12.98	120.10	-44.00	151.90

Sample No.	E <sub>corr</sub> (mV)	$I_{corr}$ $\mu A/cm^2$	Tafel β <sub>a</sub> n	Slopes nVβ <sub>c</sub>	Corrosion rate µm/y
1	-6.10	23.08	195.10	-351.80	270.00
2	-400.50	2494.30	346.70	-192.80	29170.00
3	-215.50	98.74	110.40	304.70	1155.00
4	-190.70	26.70	188.50	-163.00	312.30
5	-350.80	1236.20	384.60	-317.20	14450.00
6	-205.50	34.90	283.00	-207.40	408.20

Table 5. The corrosion parameters of six samples in 5wt%NaCl solution

Table 6. The corrosion parameters of six samples after heat treatment in 3.5wt%NaCl, solution.

Sample No	E <sub>corr</sub> mV	$I_{corr}$ $\mu A/cm^2$	Tafel Slopes β <sub>a</sub> β <sub>c</sub> mV		Corrosion rate µm/y
1	371.70	6.90	81.60	-316.80	80.79
2	-558.60	331.50	137.20	-341.70	3877.00
3	-207.00	27.15	224.20	-49.60	317.60
4	-180.80	22.34	100.00	-287.40	261.30
5	-353.30	39.36	493.60	-917.70	460.40
6	-205.50	23.66	26.40	-42.40	276.70

Table 7. The corrosion parameters of six stainless steel samples in 3.5wt% NaCl solution

Sample No.	E <sub>corr</sub> (mV)	E <sub>pit</sub> (mV)	I <sub>corr</sub> μA/cm <sup>2</sup>	Corrosion rate µm/y
1	-283.80		2.39	27.97
2	-536.10	0.29	29.36	343.40
3	-430.40	0.37	9.13	106.80
4	-390.00		6.17	72.23
5	-439.00	0.36	12.45	145.70
6	-428.50	0.66	8.06	94.27

Table 8. The corrosion parameters of six stainless steel samples in 5wt% NaCl solution

Sample	E <sub>corr</sub>	E <sub>pit</sub>	$I_{\rm corr}$	Corrosion
INO.	(mv)	(mv)	µА/сш	rate µm/y
1	-370.00	0.54	33.71	394.20
2	-594.80	0.22	9964.90	116500.00
3	-440.50	0.35	1925.10	22510.00
4	-397.20	0.50	158.79	1857.00
5	-455.10	0.32	3653.90	42730.00
6	-431.20	0.43	161.40	1887.00

Table 9. The corrosion parameters of six stainless steel samples after heat treatment in 3.5wt%NaCl solution

Samples No.	E <sub>corr</sub> mv	E <sub>pit</sub> mv	I <sub>corr</sub> μA/cm <sup>2</sup>	Corrosion rate µm/y
1	-70.60		1.80	21.11
2	-540.20	0.25	56.19	762.50
3	-440.20	0.33	9.46	110.60
4	-250.80		4.80	56.38
5	-450.10	0.32	13.36	159.70

6	-435.20	0.43	8.20	96.00

Table 10	FIS noromotore	of austanitic stainly	a ataal Allava in (	2 Swt% NoCl Solution
	LIS Darameters	of austennue stanne	55- SICCI AIIUVS III .	J.JWL/0 INACI SUIULIULI

Conc. of NaCl	Sample	R <sub>ct</sub>	C <sub>dl</sub>	Angel
	No.	ohm.cm <sup>2</sup>	μF/cm²	(θ)
	1	$146.4 \times 10^{6}$	1.934X10 <sup>-3</sup>	$25.3^{\circ}$ -
	2	1.96	202.60	$29^{0}$
%	3	13.00	137.00	-10.5
3.5	4	$1.126 \times 10^{6}$	5.031X10 <sup>-3</sup>	$-18.7^{0}$
	5	9.32	140.00	23.6
	6	34.21	41.67	-5.24

Table 11. EIS parameters of austenitic stainless- steel Alloys in 5wt% NaCl Solution



Figure 1. Heat treatment cycle





Figure 2. Potential – time curves for samples (1-6) in 3.5wt% NaCl solution.

Figure 3. Potential – time curves for samples (1-6) in 5wt% NaCl solution.



Sample 5 Sample 6 Figure 4. Variation of open – circuit potential for as forged (a) and heat treated (b) Samples (1-6) in 3.5wt% NaCl Solution









-1



Sample 3





Sample 5 Sample 6 Figure 7. Variation ofPotentiodynamic polarization for as forged (a) and heat treated (b) of Samples (1-6) in 3.5wt%NaClSolution





Figure 8. Cyclic anodic polarization curves for samples(1-6) in 3.5wt% NaCl solution











Sample 5 Sample 6 Figure 10. Variation of cyclic anodic polarization for as forged (a) and heat treated (b) of samples (1-6) in3. 5wt% NaCl solution



Figure 11. Electric equivalent circuit model used to fit the results of impedance





Figure 12. Nyquist plots for samples in 3.5wt% NaCl solution

Figure 13. Nyquist plots for samples in 5wt% NaCl solution



Sample 5 Sample 6 Figure 14. Variation of Electrochemical impedance for as forged (a) and heat treated (b) of Samples (1-6) in 3.5wt% NaCl solution



(a). Sample1

(c). Sample 2

Figure 15. Surface morphology of samples in 3.5wt% NaCl solution



(a). Sample 1(b). Sample 4(c). Sample 2Figure 16. Surface morphology of samples in 3.5wt% NaCl solution after heat treatment



Figure 17. EDAX spectra Surface morphology of samples in 3.5wt% NaCl solution

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