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EFFECTS OF TERNARY METAL ADDITIONS ON CORROSION OF SPARK PLASMA SINTERED Ni-Fe ALLOYS IN H_2SO_4 AND NaCl

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ABSTRACT. Effect of ternary additions on electrochemical behavior of Ni-Fe binary alloys developed by Spark Plasma Sintering (SPS) were investigated and reported. All specimens revealed common passivity behavior in H₂SO₄. Both binary and ternary alloys did not reveal passivity region under saline conditions. The alloys have lower corrosion rates in NaCl than in H₂SO₄. Binary Ni₅₀Fe₅₀ alloy is characterized by large pits in H₂SO₄ and homogeneous corrosion on surface in NaCl. Ni₅₀Fe₄₀Ta₁₀ shows better passivity in 1 M H₂SO₄ and Ni₅₀Fe₄₀Al₁₀ has the lowest corrosion rate in 3.5 wt.% NaCl.

KEY WORDS: Spark plasma sintering, Corrosion, Ni-Fe based alloy, Ternary alloys, Powder metallurgy

INTRODUCTION

Ni-Fe base alloys have wide applications as parts of devices in pumps, pipeline, turbines, ultrasupercritical (USC) boiler applications, and in active chemical reactors. Unlike previous research studies in Ni-based alloys which were mainly fabricated via vacuum and induction melting [1] with limitations to eliminate sulfur and phosphorus, powdered metallurgical processes of Ni-Fe based alloys have more flexibility for compositions not possible by other techniques, high dimensional accuracy, short production time, good chemical homogeneity, and capability to fabricate magnetic components and control of magnetic properties to large extent.

As at the time of reporting this study, not many studies have documented, works on the Ni-base alloys produced by spark plasma sintering (SPS). Similarly, the effects of ternary additions as anti-oxidants on the Ni-Fe-base austenitic matrix fabricated by SPS and the corrosion behaviors of sintered Ni-Fe-based alloys in selected media remains unclear. Although the work forms part of a large study to develop Ni-base alloys, it is specially focused on the SPS fabricated Ni-Fe base material with investigation of ternary additions on the binary system and concurrently studying their benefits on the corrosion characteristics of the ternary alloys.

This work used an advanced method of sintering that produces highly dense materials at much lower temperature. It is in particular good for those substances that are not easy to sinter by usual sintering processes in which there could be the contamination by the adsorptive gases on the faces of powder particles as a result of localized high-temperature spark impact. The high-speed ion movement between contacting particles probably caused by the applied electric field is mostly held accountable for helping sintering, supporting diffusion and in this manner materials are transferred in both nano and micro scales [2]. Recently researchers have shown that it is possible to sinter a system of Fe-Ni alloys with high densities successfully [3, 4].

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The present work considers the advantages of the SPS system, the gap in the Ni–Fe based alloy manufacture by SPS process being rarely reported, and in particular researches centered on the corrosion of the Ni–Fe-base sintered alloys. Diverse reports are available on imparts of alloying on the chemical, microstructural and phases characterization/behavior of different type of engineering materials. These include the super-alloys, steels and aluminum alloys [5-7], nickel nano-particles [8] and titanium base alloys, etc., and their corrosion behaviors [9] and control [10-15] in different media [16-18]. The current research development in the Ni-Fe-based austenitic matrix which falls under the category of Ni-based alloys is beginning to receive attention from researchers. The present work aimed at investigating the influence of ternary metal additives (as anti-oxidants on the Ni-Fe-base austenitic matrix) on the corrosion of sintered Ni-Fe based alloy using potentiodynamic polarization technique. The electrochemical methods of corrosion testing were conducted in 3.5 wt. % NaCl and 1 M H₂SO₄. The corrosion results were correlated with the different ternary additions and also against the microstructure for all the alloys and their performance discussed.

EXPERIMENTAL

Material preparation

Elemental ternary metal powder additions (Al, Co, Cr, Mo, Ta and Ti) and commercial elemental powders (Ni and Fe) were used for this study. Table 1 shows the compositions of the raw powders. The binary ($Ni_{50}Fe_{50}$) system was weighed accurately and mixed together with the individual ternary additions powders to make up a desired stoichiometric ratio of $Ni_{50}Fe_{40}X_{10}$ (wt.%) (where X = Al, Co, Cr, Mo, Ta and Ti). Clearly, each $Ni_{50}Fe_{40}X_{10}$ alloy contains 50% of $Ni_{50}Fe_{40}X_{10}$ and $Ni_{50}Fe_{40}X_{10}$ and $Ni_{50}Fe_{40}X_{10}$ alloy contains 50% of $Ni_{50}Fe_{40}X_{10}$ and $Ni_{50}Fe_{40}X_{10}$

The various blends of Ni-Fe and ternary powder additives were made using a Tubular Shaker/Mixer T2F at 49 rpm for 10 h as the optimum speed and time respectively. The powder was fed into the mixing compartment using a 250 mL plastic cylinder being 10% powder filled and agitated under the translational and rotational motions in a dry condition.

Elemental powder	Purity (%)	Particle size range (μm)	Suppliers
1	_ \ /		
Ni	99.5	$\geq 0.5 \text{ to } \leq 3.0$	Wear Tech (PTY) LTD
Fe	99.9	<44	Wear Tech (PTY) LTD
Al	99.7	<25	TLS Technik GMBH & Co., Germany
Co	99.9	<44	Wear Tech (PTY) LTD
Cr	99.2	<10	Wear Tech (PTY) LTD
Mo	99.9	≥2 to ≤4	CERAC Incorporated, USA
Ta	99.9	<44	TLS Technik GMBH & Co., Germany

Table 1. Characteristics of the raw powders.

Grade 1

Sintering of powder mixtures

The powder blends were sintered using a 30 mm-inner-diameter by 10 mm high graphite die in the SPS equipment, (model - HHPD-25 from FCT Systeme GmbH Germany) capable of producing 80-100 mm diameter by 5-8 mm thickness product size ranges.

TLS Technik GMBH & Co., Germany

To remove the sinter easily and reduce the temperature inhomogeneity considerably, graphite foils (0.2 mm thick) were lagged in-between the die, powders and the punches. Also, the die surface was enclosed in a permeable graphite felt (about 10 mm approx. thick) used as a

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heat resistor, thus minimizing heat loss due to temperature gradient and radiation [11, 12]. Sintering operation was done in vacuum at constant pressure (50 MPa) and temperature (1000 °C) and was followed by soaking for 10 minutes at 1000 °C maximum temperature. All cases involved heating from room temperature (25 °C) to the desired temperature varied at a 150 °C/min heating rate. After each soak time regime, equipment was powered off and the specimens were rapidly cooled to the room temperature. An implanted optical pyrometer was placed 3 mm from the top of the sample surface to measure the sintering temperature in SPS apparatus, and sintered discs of 30 mm diameter by 5 mm thick were produced.

Microstructural characterization

Specimen surfaces were prepared by grinding and polished to standard thus removing acquired graphite contamination on the surface. The microstructures of polished/ground specimens surface were examined using SEM (FESEM, JSM-7600F, Jeol, Japan) incorporated with EDX facilities (Oxford X-Max), INCA X-Stream2 pulse analyzer software, and secondary electron detectors. The INCA analyzer software was operated at acquisition time (70 s) and process time (2 s). The phases in the sintered specimen were analyzed by XRD (PANalytical Empyrean model) with Cu K α radiation and analyzer (High score plus software).

Electrochemical behavior studies

A cell with three electrodes [13] having different electrolytes was used for the electrochemical investigations. To study the corrosion behavior of alloys, acidic and chloride-containing solutions were used to simulate different types of exposure conditions. Thus, the electrolytes used in this study was 3.5 wt. % NaCl and 1 M $\rm H_2SO_4$. The specimens were made by joining an insulated copper wire by aluminum tape to a side of the specimen cold mounted in an epoxy resin. The specimen surface was smoothened by grinding using silicon carbide (SiC) grit paper (120 - 1400 μ m) and thereafter by polishing with diamond jelly (3 μ m), distilled water rinsing, degreasing in acetone and warm air drying. The open circuit potential (OCP) and potentiodynamic polarization (PDP) tests were performed to get information on the corrosion behavior of the test materials (Ni-50Fe-10X (X = Al, Co, Cr, Mo, Ta or Ti) alloy system).

Dissolution rate becomes equivalent to hydrogen evolution (reduction of H^+ in the test solution) rate at the corrosion potential ($E_{\rm corr}$). Therefore, it is within 10 mV more noble or more active than the corrosion potential, where the applied current density is linear function of the electrode potential. Tafel plots are thus, constructed from Tafel constants which are calculated from the anodic and cathodic portions. These plots are then extrapolated to meet at the $E_{\rm corr}$. At this point, the value of current density is recorded, and it corresponds to the corrosion rate of the system. The unit of the Tafel constants is V/decade. A decade of current is one order of magnitude.

An Autolab PGSTAT30 potentiostat with counter electrode (graphite rod), reference electrode (Ag/AgCl) saturated with 3 M KCl, and working electrode (specimen) were utilized. Experiment was set up at room temperature (22 \pm 2 $^{\rm o}$ C). Meanwhile, OCP stabilization was run for 2 hours and PDP curves obtained at 2 mV s $^{-1}$ scan rate initiated at -1.0 to +2.5 Volts. At equilibrium, the rate of cathodic reaction (hydrogen evolution) is balanced by an alloy dissolution rate. The state is termed corrosion potential (Ecorr). Thus, with the scanning rate used, on average the observed $E_{\rm corr}$ is reported after approximately \pm 6 min.

RESULTS AND DISCUSSION

Pre-materials and powder mixtures

Prior to mixing, the structures of the powders were studied with a field emission scanning electron microscopy (FESEM, JSM-7600F, Jeol, Japan) equipped with EDS features. The examination of morphological features of the eight as-received powders reveals irregular shapes and agglomeration in Ni, Co, Cr, Mo and Ta powders; while Fe, Al and Ti powders are spherical shape and non-porous [3]; typical of powders made by gas atomization. The SEM images of (a) Ni₅₀Fe₅₀, (b) Ni₅₀Fe₄₀Al₁₀, (c) Ni₅₀Fe₄₀Co₁₀, (d) Ni₅₀Fe₄₀Cr₁₀, (e) Ni₅₀Fe₄₀Mo₁₀, (f) Ni₅₀Fe₄₀Ta₁₀ and (g) Ni₅₀Fe₄₀Ti₁₀ (wt.%) show that mixed powders are distributed uniformly.

Sintering of powder metal blends

Powder sintering is regarded as a thermal treatment usually performed below the fusion temperature of the major components of powder compact in order to enhance the mechanical strength and the thermo-chemical integrity [1-4, 11-12]. After compaction, bordering metal powder particles are bonded by cold welds, which offer the compressed particles sufficient mechanical strength to be handled while diffusion processes form and grow necks at these contacting surfaces at the working sintering temperature [14]. The overall microstructure and properties are dependent of the nature and status of the stating raw materials. The metallochemistry is complex as it involves the multidimensional interactions among all the elemental constituents [15]. As parts of the large work, the production procedures of the NiFe alloy; and the effect of sintering parameters (starting powder morphology and particle sizes, heating rate, temperatures, pressure, sintering time, and holding time) [3] on the densification, microstructure and mechanical properties [4] and behavior (wear) have been explained [16]. Thus, in the present research, the study of the corrosion behavior of the spark plasma sintering of NiFe alloy enhanced with ternary additions is being considered.

Electrochemical behavior in 1 M H₂SO₄ and NaCl

Figure 1 shows the results obtained from the potentiometric (open circuit potential) tests of $Ni_{50}Fe_{50}$ and $Ni_{50}Fe_{40}X_{10}$ (X = Al, Co, Cr, Mo, Ta and Ti) in 1 M H_2SO_4 . The results show that Ni40%Fe-10%Co alloy exhibits the more noble potential throughout the test, while $Ni_{50}Fe_{40}Al_{10}$ showed the lowest potential. The potentials of Ni50Fe and $Ni_{50}Fe_{40}X_{10}$ (X = Co and Ta) initially increased with time and gradually increased toward a steady state value with exposure time of 2 h. This behavior shows an indication of good passivity in 1 M H_2SO_4 .

However, it was observed that the potentials values for $Ni_{50}Fe_{40}X_{10}$ (X=Mo, Ti and Cr) initially decreased and reached minimum values and then gradually increased, while $Ni_{50}Fe_{40}Mo_{10}$ stabilized throughout the 2 h test. However, the potential of $Ni_{50}Fe_{40}Ti_{10}$ was observed to decrease after about 2300 s. The initial decrease in potential signifies initial corrosion on the surfaces of the alloys and later formation of passive films, which caused the increase in the potentials. It is known that when an oxide of chromium formed a continuous layer on the surface of an alloy, it will hinder or reduce the oxidation process thus, protecting the metal. However, in order to obtain a continuous protective chromia layer, chromium contents has to be more than 18% in the alloy [17]. The potentials of the alloys after 2 h of exposure in 1 M H_2SO_4 are of the order: $Ni_{50}Fe_{40}Co_{10} > Ni_{50}Fe_{40}Mo_{10} > Ni_{50}Fe_{50} > Ni_{50}Fe_{40}Ta_{10} > Ni_{50}Fe_{40}Ti_{10} > Ni_{50}Fe_{40}Ti_{10} > Ni_{50}Fe_{40}Al_{10}$.

The $Ni_{50}Fe_{40}Al_{10}$ showed different behaviors in 1 M H_2SO_4 as its potential decreased gradually from the beginning of the test (possibly due to the dissolution of oxide film formed on the surface of the alloy) up to about 20 min and thereby stabilized throughout the test in a

similar trend as explained by [18]. Open-circuit potential (OCP) measurements are carried out to study the reactivity of the alloys, that is, their tendencies to corrode in different media with time. Therefore, the more negative is the samples potential, the less thermodynamic stable is the sample in the test solution and thus, corrodes faster. The alloys of $Ni_{50}Fe_{40}X_{10}$ (X = Al and Ti) exhibit more negative potentials as compared to others.

For their high reactivity, it is expected that $Ni_{50}Fe_{40}X_{10}$ (X = Al and Ti) will be less resistant in the test solution. Although, it is renowned that Al is much more effective than Cr in improving oxidation [19], it also exhibits corrosion and scaling resistance by producing Al_2O_3 layer on the alloys at high working temperatures. The Al_2O_3 layer formed on the surface is more thermodynamically stable and thus, caused no poisoning effect as common with the evaporation of the chromia (Cr_2O_3) scale [20-22].

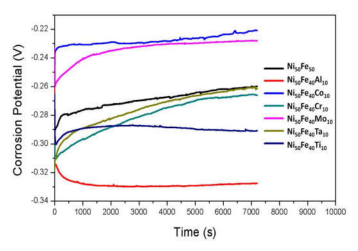


Figure 1. Open-circuit potential of $Ni_{50}Fe_{40}X_{10}$ (wt. %, X = Al, Co, Cr, Mo, Ta and Ti) in 1 M H_2SO_4 .

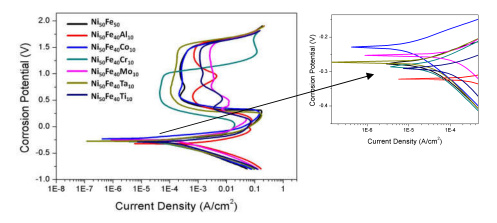


Figure 2. Potentiodynamic polarization curves of $Ni_{50}Fe_{40}X_{10}$ (wt.%), X = Al, Co, Cr, Mo, Ta and Ti) in 1 M H_2SO_4 .

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The polarization behavior of the alloys studied in 1 M H_2SO_4 is shown in Figure 2 and the electrochemical data from the electrochemical test are presented in Table 2. The results obtained confirmed the behavior observed in the OCP test. All the alloys show typical passivity behavior in 1 M H_2SO_4 and exhibit defined active to passive region as a result of metal oxidation [23] with some irregularity seen in the passivity of $Ni_{50}Fe_{40}X_{10}$ (X = Al, Ti, and Mo). This behavior indicates passivation, corrosion and re-passivation of the alloys in the solution. The $Ni_{50}Fe_{40}Ta_{10}$ alloy shows better passivity. This also was seen in the OCP results in Figure 1 where the alloys displayed more noble potential and passivated with time throughout the 2 h of test. Contrary to a report that the presence of molybdenum oxyhydroxide or molybdates within the passive films prevents corrosion, Kozhevnikov *et al.* [24] had shown that that no molybdenum oxides could be formed at the passive potential regions in HCl and H_2SO_4 solutions.

Table 2. Corrosion data obtained from potentiodynamic curves in 1 M H₂SO₄ solution.

Sample	Ecorr (V)	Icorr (A/cm ²)	Rp (ohm)	Ba (V/dec)	Bc (V/dec)	CR (mm/y)
$Ni_{50}Fe_{50}$	-0.221	1.66 x 10 ⁻⁵	2.26×10^2	0.039	0.177	1.86 x 10 ¹
Ni ₅₀ Fe ₄₀ Al ₁₀	-0.326	1.21 x 10 ⁻³	8.41×10^{0}	0.114	0.204	1.34×10^{1}
Ni ₅₀ Fe ₄₀ Co ₁₀	-0.228	1.03 x 10 ⁻⁴	7.59×10^{1}	0.075	0.192	1.15×10^{0}
Ni ₅₀ Fe ₄₀ Cr ₁₀	-0.284	1.18 x 10 ⁻⁴	5.01×10^{1}	0.088	0.17	1.25×10^{0}
Ni ₅₀ Fe ₄₀ Mo ₁₀	-0.252	4.53 x 10 ⁻⁴	2.42×10^{1}	0.114	0.2	4.60×10^{0}
Ni ₅₀ Fe ₄₀ Ta ₁₀	-0.272	8.82 x 10 ⁻⁵	5.45 x 10 ¹	0.068	0.163	9.55 x 10 ⁻¹
Ni ₅₀ Fe ₄₀ Ti ₁₀	-0.289	1.79 x 10 ⁻⁴	2.60×10^{1}	0.063	0.17	2.00×10^{0}

In Figure 2, broad bands are observed in the region 0.2-1.5 V which suggests the intense passive regions for the alloys at diverse current densities. The results in Table 2 show that $Ni_{50}Fe_{50}$ exhibits more positive potential and lowest corrosion current density (1.66 x 10^{-5} A cm⁻²), while $Ni_{50}Fe_{40}Al_{10}$ shows the lowest corrosion potential and highest corrosion current density. These results also reflect in the corrosion rates of the alloys in 1 M H_2SO_4 solution. The $Ni_{50}Fe_{50}$ alloy shows the lowest corrosion rate, followed by $Ni_{50}Fe_{40}Ta_{10}$. However, $Ni_{50}Fe_{40}Al_{10}$ alloy shows the highest corrosion rate. This further correlate with the results obtained from the OCP tests. The order of corrosion rates of the alloys in 1 M H_2SO_4 solution is: $Ni_{50}Fe_{50} < Ni_{50}Fe_{40}X_{10}$ (X= Ta < Co < Cr < Ti < Mo < Al).

Electrochemical behavior in 3.5% NaCl

A study on the tendencies of the alloys to be corroded in the 3.5% NaCl environment with time was carried out using OCP measurements. The OCP results for the alloys tested in 3.5% NaCl are presented in Figure 3. The potentials of alloys $Ni_{50}Fe_{50}$, $Ni_{50}Fe_{40}X_{10}$ (X=Cr, Ta and Co) initially increased, but later decreased with time. An irregular behavior (increase and decrease in potentials) was observed in all the alloys in 3.5% NaCl. This indicates the formation of unstable passive films on the surface of the alloys. Halogen ions such as chloride ion have the tendency to reduce the potential range of the passive region by lowering the breakdown potential resulting from the penetration and destruction of the passive film by the halogen ions. According to Leffler [25], chloride ion can break down the passive film formed and prevent reformation of new films.

The potentials of the alloys containing $Ni_{50}Fe_{40}X_{10}$ (X=Al, Mo and Ti) initially decreased. The potential of $Ni_{50}Fe_{40}X_{10}$ (X=Al and Ti) alloys increased after about 5 min. and thereafter decreased till the end of the test. At the end of the test, the order of the potentials were $Ni_{50}Fe_{40}Co_{10} > Ni_{50}Fe_{40}Ta_{10} > Ni_{50}Fe_{40}Cr_{10} > Ni_{50}Fe_{40}Ti_{10} > Ni_{50}Fe_{40}Mo_{10} > Ni_{50}Fe_{40}Al_{10}$.

1.02 x 10⁻²

βc (V/dec) CR (mm/y) Sample Ecorr (V) Icorr (A/cm²) $Rp(\Omega)$ βa (V/dec) Ni₅₀Fe₅₀ -0.408 3.45 x 10⁻⁶ 1.95 x 10⁴ 0.383 0.327 3.87 x 10⁻² 0.074 1.47 x 10 $Ni_{50}Fe_{40}Al_{10}$ -0.455 1.33×10^{-6} 6.53×10^3 0.268 4.09 x 10 0.373 4.55 x 10 -0.415 1.62 x 10⁴ 0.331 $Ni_{50}Fe_{40}Co_{10}$ -0.402 2.61 x 10 1.34 x 10° 0.248 0.325 2.78 x 10 $Ni_{50}Fe_{40}Cr_{10}$ 2.11 x 10 $Ni_{50}Fe_{40}Mo_{10}$ -0.4482.08 x 10⁻⁶ 1.17×10^4 0.157 0.286 -0.389 2.98 x 10⁻⁶ 0.205 0.376 3.23 x 10⁻² $Ni_{50}Fe_{40}Ta_{10}$ 1.14 x 10⁴

1.69 x 10⁴

0.498

0.136

Table 3. Corrosion data obtained from potentiodynamic curves in 3.5% NaCl solution.

1.75 x 10⁻⁶

Ni₅₀Fe₄₀Ti₁₀

-0.627

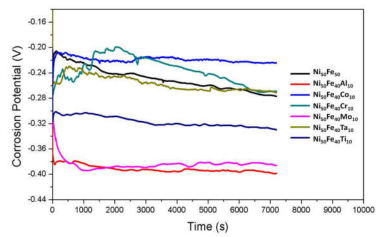


Figure 3. Open-circuit potential of $Ni_{50}Fe_{40}X_{10}$ (X = Al, Co, Cr, Mo, Ta and Ti (wt.%)) in 3.5% NaCl.

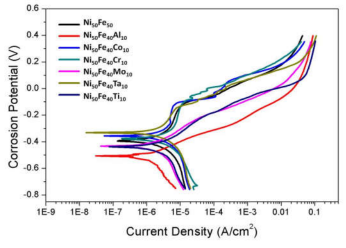
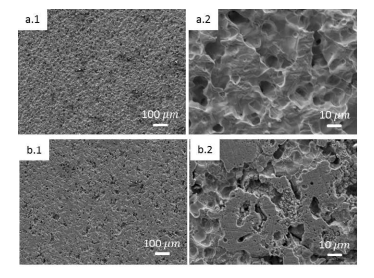


Figure 4. Potentiodynamic polarization curves of $Ni_{50}Fe_{40}X_{10}$ (X = Al, Co, Cr, Mo, Ta and Ti (wt. %)) in 3.5% NaCl.

The behavior of the alloys in 3.5% NaCl solution was further studied using polarization technique. The results are presented in Figure 4 with obtained data in Table 3. The Ni₅₀Fe₄₀Co₁₀ shows a little of active-to passive-region, but other alloys generally did not show any noticeable active-to-passive behavior in the solution, but passivate spontaneously and directly go into a pseudo-passive state. It was observed that as other elements were alloyed with Ni-Fe, a decrease in anodic and cathodic current densities was observed. This decrease indicates anodic dissolution of the alloys and cathodic hydrogen evolution reaction processes. Unlike the behavior of the alloys in 1 M H₂SO₄ solution, addition of other elements to Ni-Fe improved its corrosion resistance in 3.5% NaCl solution with Ni₅₀Fe₄₀Al₁₀ showing the lowest corrosion rate. The corrosion rates of the alloys in 3.5% NaCl solution were lower compared to their corrosion rates in 1 M H₂SO₄. The corrosion rates of the alloys in 3.5% NaCl are as follows: Ni₅₀Fe₄₀X₁₀ $(X=Ti < Al < Mo < Cr < Ta) < Ni_{50}Fe_{50} < Ni_{50}Fe_{40}Co_{10}. \ It \ was \ observed \ that \ all \ the \ alloys$ generally exhibited lower corrosion rates in chloride solution compared to sulfuric acid solution. Similar observation was made by Ameer et al. [26] and Betova et al. [27] in their report, it was seen that the corrosion rate of the alloy tested was higher in the sulfate containing solution than in the chloride solution.

Micrographs of samples after corrosion in 1 M H₂SO₄ and 3.5% NaCl

Figure 5 shows SEM images after corrosion in 1 M H₂SO₄ at different magnifications. The pits are a result of corroded region after they were exposed in the corrosive environment for the different samples showing an indication of how aggressive was the corrosive solution per sample respectively. For Ni₅₀Fe₅₀ (Figure 5 (a2)) in 1 M H₂SO₄ environment, the alloy is characterized by much larger pits as an indication that the corrosive liquid penetrated much deeper into the sample than just its surface Figure 5 (a.2 and c.2). Ni₅₀Fe₄₀Al₁₀ (Figure 5 (b2)) and Ni₅₀Fe₄₀Ti₁₀ (Figure 5 (g.2)) at magnifications of 1000x show that corrosion took place around the grain boundaries.



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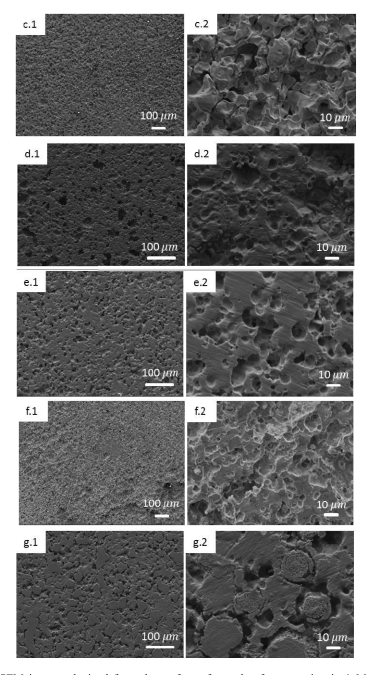
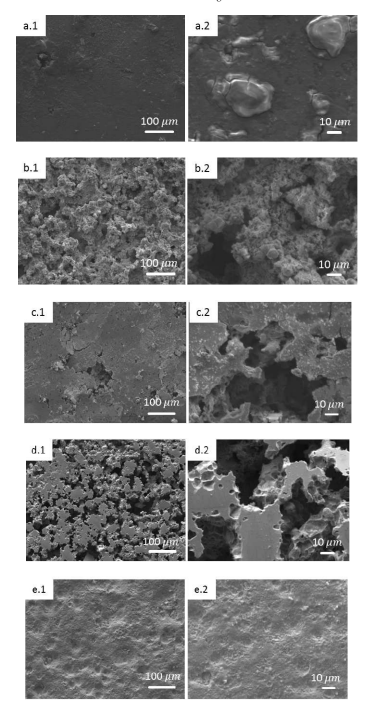


Figure 5. SEM images obtained from the surface of sample after corrosion in 1 M H_2SO_4 at different magnifications ((1) 200x and (2)1000x), (a) $Ni_{50}Fe_{50}$, (b) $Ni_{50}Fe_{40}Al_{10}$, (c) $Ni_{50}Fe_{40}Co_{10}$, (d) $Ni_{50}Fe_{40}Cr_{10}$, (e) $Ni_{50}Fe_{40}Mo_{10}$, (f) $Ni_{50}Fe_{40}Ta_{10}$, and (g) $Ni_{50}Fe_{40}Ti_{10}$.

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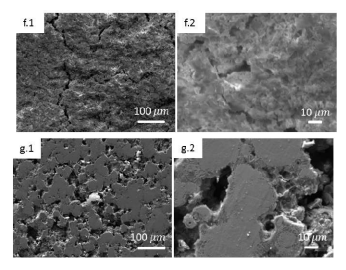


Figure 6. SEM scan obtained from the surface of sample after corrosion in 3.5% NaCl at different magnifications ((1) 200x and (2)1000x), (a) Ni₅₀Fe₅₀, (b) Ni₅₀Fe₄₀Al₁₀, (c) Ni₅₀Fe₄₀Co₁₀, (d) Ni₅₀Fe₄₀Cr₁₀, (e) Ni₅₀Fe₄₀Mo₁₀, (f) Ni₅₀Fe₄₀Ta₁₀, and (g) Ni₅₀Fe₄₀Ti₁₀.

Figure 6 shows SEM images after corrosion in 3.5% NaCl at different magnifications. As observed, $Ni_{50}Fe_{40}Mo_{10}$ sample (Figure 6 (e.2)) has the least susceptibility to corrosion. It is corrosion resistant. Molybdenum (Mo) element reacts slowly when exposed to acids because it is a very hard transition metal but is softer and more ductile than Tungsten. $Ni_{50}Fe_{40}Al_{10}$ (Figure 6 (b.2)) also did not suffer much attack since Al produces a protective layer in corrosive environments. $Ni_{50}Fe_{50}$ (Figure 6 (a.2)) shows some corrosion oxides that are on top of the surface as shown on 1000x higher magnification. The surface revealed a homogeneously corroded surface. However, weak localized corrosion attacks were observed and were related to the increased porosity due to the detaching of the Ni-Fe particles [28]. In Figure 6 (f.1) $Ni_{50}Fe_{40}Ta_{10}$ sample has visible cracks which were no visible before exposure in the corrosive environment, similarly $Ni_{50}Fe_{40}Co_{10}$ (Figure 6 (c.1 and 2)) has some cracks which is associated with huge material loss and the corrosive environment penetrating much deeper into the sample. $Ni_{50}Fe_{40}Cr_{10}$ (Figure 6 (g.1)) and $Ni_{50}Fe_{40}Ti_{10}$ (Figure 6 (d.1)) corrosion attack took place mainly along the grain boundaries. In most of the alloys corrosion is believed to have progressed or initiated on pre-existing micropores.

CONCLUSION

Ternary additions of Al, Co, Cr, Mo, Ta and Ti were successfully incorporated into the binary $Ni_{50}Fe_{50}$ base alloy by SPS. The Co and Mo additions render an alloy nobler (\sim -0.23V) in H_2SO_4 solution. However, Al addition drastically shifted the potential to more negative regions (\sim -0.33V). For all the alloys, intense passive region (\sim 0.25 – 1.5 V) is seen in the acid medium. $Ni_{50}Fe_{40}Cr_{10}$ showed the lowest passive current density (1E-4 A/cm²) though its potential passive range is low. The addition of Co and Cr gave the nobler potentials. In contrast, Al and Mo showed the lowest potentials. Only $Ni_{50}Fe_{40}Co_{10}$ and $Ni_{50}Fe_{40}Cr_{10}$ alloys showed a short region of active to passive. Nevertheless, $Ni_{50}Fe_{40}Al_{10}$ alloy showed the lowest corrosion rate (\sim 1.02 x 10^{-2} mm/y) in 3.5% NaCl. It was found that the corrosion rate of the alloys in 3.5% NaCl (\sim 1.02 x 10^{-2} – 4.55 x 10^{-2} mm/y) is lower than in 1 M H_2SO_4 (\sim 1.86 x 10^{-1} – 1.34 x 10^1 mm/y).

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