Effect of Noble Element Alloying on Passivity and Passivity Breakdown of Ni

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Abstract

Welding of stainless steels can generate welding fumes containing carcinogenic hexavalent chromium (Cr⁶⁺). To mitigate this problem, a new Ni-Cu-Pd welding consumable alloy has been developed. The addition of a small amount of Cu and Pd in Ni enhances the galvanic compatibility of Ni with stainless steels and improves its localized corrosion behavior. In this paper, the artificial pit electrode technique and X-ray photoelectron spectroscopy were used to study the benefits of Cu and Pd alloying. The passive film on Ni-10Cu-1Pd alloy mainly consisted of outer Ni-hydroxide and inner oxide, and the noble elements Pd and Cu apparently did not contribute to the formation of the passive film. However, Pd catalyzed the reduction of Cu at the bottom of artificial pits, which enhanced the cathodic reaction and thus ennobled the protection potential, making stable pit growth more difficult. This catalytic effect of Pd also prevented the propagation of deep pits in a test using a thin foil crevice sample.

During welding of stainless steels (SS), the evaporation and oxidation of Cr from the molten weld pool can result in the generation of carcinogenic hexavalent chromium (Cr⁶⁺) in the welding fume.¹² The U.S. Occupational Safety and Health Administration recently issued a new standard for occupational exposure to hexavalent chromium containing a substantial reduction in the permissible exposure limit, which will make it difficult to weld SS in locations that are not extremely well-ventilated.³⁴ To reduce the emission of Cr⁶⁺ from weld fumes, a Cr-free consumable has been developed for welding austenitic stainless steel⁵,⁶ The resulting welds in type 304L SS exhibit mechanical properties and corrosion resistance comparable to welds made with commonly used Cr-bearing consumables.

Because the new consumable has a different chemical composition than the stainless steel base plate to be welded, it is important to consider the effects of galvanic interaction on the localized corrosion behavior of the new consumable. Therefore, two major criteria were established to develop a new weld metal for stainless steel:⁵

1. The breakdown and repassivation potentials of the weld metal should be higher than the corrosion potential of the stainless steel base metal in environments of interest to prevent localized attack of the weld metal.
2. The corrosion potential of the weld metal should be slightly higher than that of the stainless steel base metal so that the weld metal is cathodically protected.

Ni-Cu alloys initially were selected based on their galvanic compatibility with types 304...
and 316 SS in chloride environments.\textsuperscript{5} 304L SS was welded with Monel filler metal, which contains 28-34 wt% Cu, to create high-quality welds with no cracks.\textsuperscript{5} However, segregated regions rich in Cu were weak spots for corrosion susceptibility in the welds.\textsuperscript{5} Subsequent work on alloys with lower Cu content led to the selection of Ni-10 wt%Cu-1 wt%Pd, which had higher breakdown potential than Monel (though not as high as 304 SS), higher repassivation potential, and slightly higher OCP than 304 SS, and no localized corrosion observed after long term exposure in 0.1 M NaCl. The corrosion properties of the newly developed alloy satisfy the design criteria for a Cr-free filler metal for welding of 304 SS.\textsuperscript{6} However, the role of Pd and Cu on the improvement of the corrosion properties has not been clearly understood.

Cu has been known to have both beneficial and harmful effects on the corrosion resistance of Ni alloys. The corrosion resistance of Ni-Cu alloys was found to be superior to pure Ni only in the active region,\textsuperscript{7} and the optimum content of Cu in Ni-Cu alloys to reduce the corrosion rate was about 30\% in 10\% HCl, 3\% NaCl, and 10\% HNO\textsubscript{3}.\textsuperscript{8} Verink generated experimental Pourbaix diagrams for Cu(0-75\%)Ni systems in Cl\textsuperscript{-}-containing solutions. With decreasing Cu content, the passivity of Ni-Cu alloys was improved in the acidic region. The higher Cu-containing Ni-Cu alloys showed higher weight loss in quiescent seawater after 2 years.\textsuperscript{10} A low Cu content in Ni-Cu alloys promoted the formation of a protective Ni(OH)\textsubscript{2} film, and the dissolution of Cu was inhibited in alkaline solution.\textsuperscript{11}

Although increasing Cu content from 30 through 53 wt\% increased the weight loss, it also decreased the pitting depth of Ni-Cu alloys in the same environment.\textsuperscript{12} After long term immersion, Ni-Cu alloys including Monel 400 showed shallow pits in Cl\textsuperscript{-}-containing environments, whereas Cr-containing alloys such as the stainless steels were deeply attacked at local sites.\textsuperscript{13-17} Isaacs and Newman reported that dissolved Cu was plated out in an artificial pit electrode of type 304 stainless steel in 1 M NaCl solution.\textsuperscript{18,19} This reaction occurred around -0.27 V\textsubscript{SCE}, indicating the reduction of Cu from CuCl\textsubscript{2}. For Ni-Cu alloys with Cu content from 30-50 wt\% in natural seawater, Cu selectively dissolved as Cu\textsuperscript{+} and Cu\textsuperscript{2+}, and redeposited as a metallic Cu.\textsuperscript{20} The kinetics of the dissolution and reduction of Cu can play an important role in breakdown and repassivation behavior of Ni-Cu alloys.

Pd is not reactive in many environments and is very effective as a hydrogeneration catalyst.\textsuperscript{21} The doping effect of the noble Pd on the oxide film of Ni-base alloys was reported to improve the life of the oxide in a deaerated aqueous environment at 288\degree C.\textsuperscript{22-24} Hettiarachchi\textsuperscript{25} also found that Pd incorporation into the growing oxide on the surface of austenitic stainless steel improved the resistance to stress corrosion cracking (SCC).

The addition of a noble metal such as Au, Pt, or Pd ennobled the open circuit potential (OCP) of type 316 stainless steel in 1 N H\textsubscript{2}SO\textsubscript{4} solution for 150 h, and enrichment of these noble metals was detected on the surface.\textsuperscript{26} The ennoblement of the OCP and low passive/active current densities of Fe-25Cr alloy and Fe-18Cr-10Ni alloy were also reported with the enrichment of Pd on the surface by electrolytic deposition.\textsuperscript{27} Potgieter\textsuperscript{28} studied the cathodic modification of alloys with noble metals. The noble metals provided a high exchange current density for the reduction reaction, and then shifted the OCP to the passive range resulting in easy passivation. The effectiveness of this modification was best for platinum group metals (PGMs) and was ordered in terms of high effectiveness from Pt to Pd, Ir, Ru, and Os. Potgieter also found that the noble metal was enriched on the surface after active dissolution and that Mo had a synergetic effect with PGMs. From a study of pure Cr, the addition of 0.1\% Pd lowered the corrosion rate by 10\textsuperscript{5} times, as the PGMs blocked the kink sites where active dissolution preferentially occurred.
In this paper, the influence of Cu and Pd on the passivity and passivity breakdown of Ni is addressed. Passive film composition determined by X-ray photoelectron spectroscopy (XPS), and localized corrosion kinetics determined by artificial pit electrode experiments and proved by a thin foil crevice test, are presented.

Experimental

Five Ni alloys were studied: pure Ni, Ni-10Cu, Ni-1Pd, Ni-30Cu, and Ni-10Cu-1Pd (in wt%). The alloys were melted from pure materials and cast in the shape of small disks. By a series of processes, they were worked to wires with 0.17 or 0.34 mm diam. The wires were degreased and annealed at 1100°C for 1 h in vacuum-sealed quartz tubes. Finally, they were mounted in epoxy. The details of the artificial pit electrode are shown schematically shown in Fig. 1. The samples were ground to no. 600 grit and then inserted facing upward into a cell that was filled with 0.1 M NaCl. A Pt wire counter electrode and saturated calomel electrode (SCE) reference electrode were used.

Three electrochemical steps in series were used to evaluate the polarization behavior of the artificial pits. First, to make an artificial pit of a specific depth, a constant potential was applied, 0.8 V_{SCE} for Ni-30Cu and 0.5 V_{SCE} for the rest of the samples. The time to generate a certain pit depth during the potentiostatic test was calculated using Faraday’s law on preliminary
experimental runs. Four different depths of pits on each sample were tested, where the ratio of the pit depth/diameter was in the range of 0.5 to 4.0. The potential was then stepped down to -0.12 \( V_{\text{SCE}} \) for 30 s to dissolve the salt film in the pit.\(^{29} \) Finally, the potential was scanned from 0.3 \( V_{\text{SCE}} \) to -0.2 \( V_{\text{SCE}} \) at 10 mV/min. The depths of the artificial pits were confirmed after the experiments by observation in an optical microscope and by integration of the current. Some of the wires were extracted by breaking the epoxy mold after the test and then analyzed using SEM-EDS. Upward scanned potentiodynamic polarization experiments were performed on some samples with no prior dissolution, so that the surface of the electrode was coplanar with the epoxy mount.

Samples of each alloy 10 \( \times \) 10 mm in size were polarized in 0.1 M NaCl for 10 min at 0 \( V_{\text{SCE}} \) where all of the samples were in the passive state. After passivation, the samples were rinsed with distilled water and dried. After sitting in the vacuum chamber for about 16 h, the passive films were analyzed using XPS at 13 kV with monochromated X-rays from an Al target. The sample surface was intermittently sputter-etched using Ar ions for 15 or 30 s. The sputtering rate was about 20 \( \text{Å/min} \).

Two alloys, pure Ni and Ni-10Cu-1Pd, were prepared as foils for crevice corrosion testing. Homogenized 1 mm thick plates were polished down to no. 600 grit abrasive paper until reaching a final thickness of 0.17 mm and electrically connected with a Cu wire. This foil was embedded in epoxy with one edge of the foil exposed at the top surface of the mold. A side of the epoxy mold parallel to the foil electrode was thinned to about 0.17 mm by grinding and polished to 1 \( \mu \text{m} \) to facilitate visual observation of the side of the foil (Fig. 2). The foil edge was exposed vertically in a cell containing 0.1 M NaCl and a potential of 0.5 \( V_{\text{SCE}} \) was applied for 6 h. The configuration of the electrodes and applied potential were the same as those used as in the artificial pit electrode experiments. Furthermore, the foil thickness was the same as the diameter of the wires in the artificial pit electrode. The advantage of the foil crevice electrode was that it allowed observation of the change in topography of the surface during polarization. The side view of the foil sample during propagation of the localized attack downward from the foil edge was recorded using a video camera and VCR.

**Results**
**XPS analysis** - XPS analysis was used to evaluate the composition of the prepassivated alloy surfaces. The O 1 s spectra (not shown) indicated that the outermost surface of the passive film for all the tested Ni alloys consisted primarily of hydroxide, and that the inner part of the passive film was mixed oxide and hydroxide. The Pd 3d_{5/2}, Cu 2p_{3/2} and Cu LMM Auger peaks were analyzed to assess the oxidation states of Pd and Cu. These elements mainly existed in the unoxidized metallic state, but small amounts of PdO and CuO were detected in the outermost region, Fig. 3.

The height of the O 1 s, Cu 2p_{3/2}, and Pd 3d_{5/2} peaks of Ni-10Cu-1Pd alloy were analyzed as a function of depth during sputter profiling, Fig. 4. It is possible that some Cu^{+1} is included in the analysis for CuO, but the total amount of oxidized Cu was small. Figure 3 shows that both metallic Cu and Pd were present in the passive film, but in depleted concentrations relative to the content in the bulk alloy.

![XPS results](image)

Figure 3. XPS results of (a) Cu 2p and (b) Pd 3d peaks of Ni-10Cu-1Pd alloy, showing some involvement of Cu and Pd in the outer layer.
Artificial pit electrode studies - Figure 5 shows the polarization curves for artificial pit electrodes of 0.17 mm diam and similar depths in the range of 0.3–0.4 mm, obtained by scanning downward from 0.3 V_{SCE} as described above. The pit current decreased during the reverse scan as the potential decreased. After reaching a zero-current potential, the current changed polarity becoming negative. The cessation of artificial pit growth can be accomplished by two different processes—repassivation or deactivation. Repassivation occurs when the current density decreases during downward potential scanning and the conditions at the pit bottom change to the point at which the active state can no longer be sustained so the surface repassivates.\textsuperscript{30–32} This repassivation process is accompanied by a sharp drop in current density to a lower passive current density. The cessation of pit growth during downward scanning can also result from a deactivation process in which the potential is scanned through the corrosion potential of the electrode surface in the pit environment such that the net current becomes cathodic.\textsuperscript{30,31} The term “protection potential” is generic and encompasses both repassivation and deactivation.

Cessation of the artificial pit electrodes in this study seemed to be a deactivation process occurring as the result of an increase in the cathodic reaction. The addition of 10% Cu decreased the anodic current relative to pure Ni and increased the cathodic current in the

\begin{figure}[h]
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\includegraphics[width=0.5\textwidth]{figure4.png}
\caption{XPS depth-profile results of Ni–10Cu–1Pd alloy after the passivation at 0 V_{SCE} for 10 min. (a) Intensity of Cu 2p_{3/2}, CuO 2p_{3/2} (referred to left axis), O 2p_{3/2} and OH 2p_{3/2} (referred to right axis). (b) Intensity of Pd 3d_{5/2}, PdO 3d_{5/2}, and PdO\textsubscript{2} 3d_{5/2}.}
\end{figure}
pit, resulting in deactivation and an increase in the protection potential. The addition of more Cu, to 30 wt %, provided no further benefit. However, the addition of even 1 wt % Pd to Ni-10Cu resulted in a large increase in protection potential as a result of a large increase in the cathodic reaction rate inside the pit. Noble metal alloys have previously been found to cause such increases in cathodic reaction rates. 21,33

The cathodic reaction in the artificial pit might come from reduction of dissolved oxygen molecules, protons, or metallic ions. Figure 6 shows pit polarization curves for Ni-10Cu-1Pd artificial pits of depth 0.34 mm in deaerated and aerated solutions. The curves are identical, indicating that the ca-thodic reaction was not oxygen. This was expected, as oxygen should be depleted in the constricted crevice. Furthermore, the protection potential of Ni-10Cu-1Pd alloy was 0.15 V_{SCE}, which is much higher than the reversible potential of H^+/H_2, indicating that hydrogen evolution is probably not the
reduction process. However, this potential is very close to the standard equilibrium potential of \( \text{Cu}^{2+}/\text{Cu} \), 0.337 V\text{NHE} or about 0.1 V\text{SCE}.

In an effort to assess the possibility of cupric ion reduction in the artificial pit electrodes, measurements were made in environments with purposeful additions of cupric ions at a concentration of 0.5 M

Figure 7. Artificial pit polarization curves for various alloys in 0.1 M NaCl with or without the addition of 0.5 M cupric ion.

Figure 8. Effect of the pit depth on the polarization curves for Ni-10Cu-1Pd artificial pit electrodes of diameter 0.34 mm. The numbers refer to pit depth in mm.

using samples with no Cu content, Fig. 7. The protection potential for the alloy Ni-1Pd was 0.1 V\text{SCE} in 0.5 M CuCl\(_2\) + 0.1 M NaCl solution, but only -0.4 V\text{SCE} in 0.1 M NaCl solution without cupric addition. The protection potential for Ni-1Pd in the cupric-containing solution was close to that for Ni-10Cu-1Pd alloy in 0.1 M NaCl solution with no cupric addition. Pure Ni also
showed a very high protection potential in cupric-containing solution, but it was lower than that of Ni-1Pd. The protection potentials for both pure Ni and Ni-1Pd in 0.1 M NaCl with no cupric ion were much lower. These observations suggest that the higher rate of the cathodic reaction for Ni-10Cu-1Pd artificial pits in chloride solution was associated with the reduction of the copper ions that were previously dissolved from the alloy into the pit environment. Further evidence for this is given by Fig. 8, which shows the effect of pit depth on the backward-scanned polarization curves for Ni-10Cu-1Pd artificial electrodes of diameter 0.34 mm. For shallow artificial pit electrodes, a sharp decrease in cathodic current of about two orders of magnitude was observed during the back scan. The potential at which this decrease occurred decreased with increasing pit depth. The deepest pit, 1.14 mm deep, was not scanned down far enough to exhibit the current decrease. The sharp decrease is apparently associated with depletion of copper ion from the pit environment. Deeper pits contain a larger reservoir of the dissolved species, and the depletion occurs at a lower potential, after more cathodic charge has passed and more deposition has occurred.

After the first potentiostatic step at 0.5 V_{SCE} in 0.1 M NaCl solution, the epoxy mold around a Ni-10Cu-1Pd sample was mechanically broken and the alloy wire was extracted. The top surface of the wire, which was actually the bottom of the artificial pit electrode, was coated with carbon and analyzed using scanning electron microscopy-energy-dispersive spectroscopy (SEM-EDS), Fig. 9. For Ni-10Cu-1Pd with a pit depth of 0.34 mm, the surface was flat with some scattered crystallographic pits. Local enrichment of Pd and Cu was detected at the same sites across the pit surface. A small amount Cl was also detected at those sites. It has been reported that Cu and Cl can form compounds in pits.\(^{18,19}\)

To monitor the localized attack morphology, a dissolving thin foil crevice sample was viewed from the side face as shown in Fig. 2. A constant potential was applied to pure Ni and Ni-10Cu-1Pd alloy for 6 h in 0.1 M NaCl solution while recording the morphology of the foil from the side. For pure Ni at 0.5 V_{SCE}, breakdown occurred at the two edges of the foil after 1 h and then propagated into the depth. The middle of the foil was unattacked and the surface remained at the position of the original solution/metal (S/M) interface, Fig. 10. For Ni-10Cu-1Pd alloy tested at 0.5 V_{SCE}, the whole

![Figure 9. Pit bottom of Ni-10Cu-1Pd after the first step of potentiostatic test at 0.5 V_{SCE} showing the flat surface with the scattered crystallographic pitting.](image)
surface of the foil was attacked after 1 h and then the S/M interface gradually shifted down with time, Fig. 11. The S/M interface still remained almost as flat as the attack proceeded by repeated breakdown and fast deactivation of many sites. The Pd addition changed the mode of pit propagation from localized growth of few sites to wide spreading of many sites.

**Discussion**

Previous work showed that the alloy Ni-10Cu-1Pd exhibited superior resistance to localized corrosion in potentiodynamic testing

![Figure 10. (Color online) Propagation with time of the localized attack on pure Ni foil at 0.5 V_{SCE} in 0.1 M NaCl solution.](image)

The reasons for this corrosion resistance were explored in this work. The beneficial effects of noble Pd additions on the passivity of other Ni-alloys has been reported previously. However, the results of this study suggest that the addition of Cu and Pd to Ni has little effect on the passive film and that improvement in localized behavior obtained by
alloying is more related to the effects on the anodic and cathodic kinetics in pits than to improvements in the protectiveness of the passive film.

It was shown that Ni artificial pit electrodes scanned downward from high potentials exhibit high zero-current potentials when Cu and Pd are alloyed at low levels or when cupric ions are introduced into the electrolyte. The cessation of artificial pit growth is considered to be a deactivation process resulting from a dramatic increase in the cathodic kinetics in the pit. The cathodic reaction was shown not to be oxygen reduction or hydrogen evolution and it can be concluded that the primary cathodic reaction in the artificial pit electrodes was reduction of Cu ions. The great enhancement in the rate of this reaction for the Ni-10Cu-1Pd alloy relative to the Ni-Cu binary alloys and the co-location of Pd and Cu on the surface of an artificial pit electrode suggest that the Cu reduction was catalyzed at sites where Pd was enriched at the surface, perhaps by surface enrichment from the dissolution of Ni and Cu. The synergism between cupric ions in solution and Pd enrichment on the surface resulted in a strong ennoblement of the deactivation process, which decreased the stability of the pits.

Pd also affects the anodic pit kinetics. Comparison of the artificial pit polarization curves for pure Ni and Ni-1Pd in the cupric free 0.1 M NaCl solution (Fig. 7) indicates that in the absence of Cu, Pd

![Figure 11](image_url)

Figure 11. Propagation with time of the localized attack on Ni-10Cu-1Pd foil at 0.5 V_{SCE} in 0.1 M NaCl solution.
results in a large increase in the Ni active dissolution kinetics. In this cupric-free solution, the primary cathodic reaction is probably hydrogen evolution, and there is no obvious effect of the Pd addition on the rate of this reaction. However, the rate of anodic dissolution at a given potential is increased by many orders of magnitude for the Ni-1Pd alloy. The curves in Fig. 5 indicate that this enhancement of Ni dissolution by Pd was not observed for the Ni-10Cu alloy. However, Fig. 8 shows that the anodic portion of the back-scanned polarization curves for Ni-10Cu-1Pd was dependent on the depth of the pit. The deeper pits with depth of 0.6 and 1.14 mm exhibited much higher dissolution rates at a given potential than the shallow pits with depth of 0.12 and 0.27 mm. It is likely that the extent of Pd enrichment increased for the deeper pits, so the effect of Pd on the anodic kinetics in the pits was more evident for the deeper pits. Pd accelerates the rate of anodic dissolution, but this is more than compensated by the strong effect of Pd on Cu deposition, which results in a very high pit corrosion potential. The mechanism for Pd enhancement of Ni dissolution is not understood.

The creation of a deep artificial pit by forced dissolution at high potential results in the formation of a pit environment that could be different than for a pit that grows at a lower open-circuit potential, so the relevance of the artificial pit electrode measurements to pitting at open circuit must be considered. The breakdown and repassivation potentials for Ni-10Cu-1Pd in 0.1 M NaCl were found to be +220 and +150 mV SCE, respectively. A study of the behavior of Monel-400 (with ~30% Cu) in deaerated chloride solution found that both Cu(I) and Cu(II) were generated during anodic dissolution in a wide range of potential, from -350 to +400 mV Ag/AgCl. At potentials above +150 mV Ag/AgCl (+130 mV SCE), dissolution was dominated by the formation of Cu(II). Therefore, the environment created during the formation of the recess in the artificial pit electrode should be similar to that formed in pits at the breakdown potential measured in cyclic polarization experiments.

Figure 12 shows polarization curves measured from the cross sections of 0.17 mm wires that were not pre-dissolved to form deep artificial pit electrodes, i.e., the surfaces of the wires were coplanar with the epoxy mount. The potential was scanned at 10 mV/min in 0.1 M NaCl solution, with a vertex current density of 0.5 mA/cm².

![Figure 12. Polarization curves for flat cross section of wire electrodes with 0.17 mm diameter without pre-dissolution to form deep artificial pits in 0.1 M NaCl.](image-url)
The average depth of attack during the cyclic scan was much less than the depth of the artificial pit electrodes and the potentials were much lower. The breakdown potentials for pure Ni, Ni-10Cu, and Ni-10Cu-1Pd were all relatively close. Note that the breakdown potentials of the small wire electrodes were lower than for samples with large areas. However, the backscanned portions of the curves for the flat wire cross sections were vastly different from each other. The pure Ni sample exhibited a large hysteresis and a zero-current potential on the backscan equal to the value on the forward scan. The Ni-10Cu-1Pd exhibited a very small hysteresis and much higher zero-current potential on the backscan. The behavior of Ni-10Cu was intermediate. Clearly, the Cu and Pd contents exerted a strong influence on these curves, even with relatively small extents of attack.

The general understanding of critical potentials is that localized corrosion can initiate at potentials above the breakdown potential, and it can continue to grow at potentials above the protection potential. In real applications under open-circuit conditions, localized corrosion initiates when the potential is high, and then the potential drops as a result of the activation at the pit bottom. Therefore, the environment generated in a pit growing at high open-circuit potentials is relevant to the possible cessation of pit growth at lower open circuit potentials.

Newman and Isaacs performed cyclic voltammetry measurements on small Pt wires inserted into an artificial pit electrode made from a commercial 304L SS wire of larger diameter. They observed oxidation and reduction peaks at -200 and -350 mV SCE, respectively, which they assigned to the Cu/CuCl$_2$ redox reaction associated with the 0.33% Cu in the alloy. As mentioned above, the protection potential for the Ni-10Cu-1Pd artificial pit electrode is much higher than these values and close to the standard potential for Cu/Cu$^{2+}$. The lower protection potential found for the flat Ni-10Cu-1Pd wire electrode is still higher than the peaks observed by Newman and Isaacs and might be connected to lower cupric ion concentrations. Our interpretation is that the cathodic reaction for the Cu-containing alloys is the reduction of Cu$^{2+}$ and that it is relevant to the behavior of these alloys at open circuit conditions.

There have been many studies reporting the superior resistance of Cu-containing Ni alloys to localized corrosion in chloride environments for more than 30 years. These alloys usually showed much shallower pits than stainless steel even if the first formation of the pits was earlier for Ni-Cu alloys than for stainless steels. Ni-10Cu-1Pd alloy in this study also showed very shallow pits in the long term crevice corrosion test, compared to type 304L stainless. This behavior can be explained by the stifling of pit growth by the plating of Cu from the pit environment.

**Conclusions**

The effects of Cu and Pd on the passivation and localized corrosion behavior of Ni alloys were investigated using XPS and the artificial pit electrode technique. The following are the important conclusions of this study.

The passive film on Ni-10Cu-1Pd alloy consisted of a mixture of oxide and hydroxide, with the outermost layer being mainly hydroxide. Very little Cu and Pd were found in the passive film and they were mostly in the metallic state. This small change in the composition of passive film cannot explain the improved resistance to the localized corrosion of Ni-Cu-Pd alloy.

The addition of 1 wt % Pd increased the artificial pit electrode protection potential considerably over the whole range of the pit depth studied. This protection potential for the Ni-10Cu-1Pd artificial pit electrode is close to the standard potential for Cu/Cu$^{2+}$. 
The improved resistance to the localized corrosion of Ni-10Cu-1Pd alloy is attributed primarily to the cessation of pit growth by enhanced reduction of copper ions from the pit environment in the presence of Pd.

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