Primary Water SCC Understanding and Characterization Through Fundamental Testing in the Vicinity of the Nickel/Nickel Oxide Phase Transition

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Primary Water SCC Understanding and Characterization Through Fundamental Testing in the Vicinity of the Nickel/Nickel Oxide Phase Transition

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ABSTRACT

This paper quantifies the nickel alloy stress corrosion crack growth rate (SCCGR) dissolved hydrogen level functionality. SCCGR has been observed to exhibit a maximum in proximity to the nickel/nickel oxide phase transition. The dissolved hydrogen level SCCGR dependency has been quantified in a phenomenological model in terms of the stability of nickel oxide not the dissolved hydrogen level. The observed SCCGR dependency has been extended to lower temperatures through the developed model and Contact Electrical Resistance (CER) measurements of the nickel/nickel oxide phase transition. Understanding obtained from this hydrogen level SCC functionality and complementary SCC subprocesses test results is discussed. Specifically, the possible SCC fundamental subprocesses of corrosion kinetics, hydrogen permeation and pickup have also been measured for nickel alloys. Secondary Ion Mass Spectroscopy (SIMS) analysis has been performed on SCCGR specimens tested in heavy water (D2O).

INTRODUCTION

Hydrogen is often added to high temperature water to maintain low levels of dissolved oxygen and thereby minimize corrosion of structural metals. Numerous studies have shown that hydrogen dissolved in high temperature water (288 to 360°C) or hydrogen gas in steam (400°C) affects the stress corrosion cracking (SCC) performance of nickel alloys (e.g., alloys 600 and X-750). Test results have demonstrated that a crack growth rate maximum, with respect to coolant hydrogen variation, is observed in proximity to a key phase transition, the nickel (Ni) to nickel oxide (NiO) phase transition, see Figure 1.

Reference (1) identified that the SCC hydrogen dependency is fundamentally described by the extent that the alloy's corrosion potential deviates from the potential of the Ni/NiO phase transition. This potential difference represents the relative stability of the SCC controlling oxide films (e.g., crack tip oxides are often of a NiO structure).

This paper reports results from numerous tightly controlled SCC crack growth rate (SCCGR) tests as a function of dissolved hydrogen level for three nickel alloys (X-750 AH, X-750 HTH and alloy 600). Except for new tests of alloy 600 at a stress intensity factor of 65.9 MPa√m, these tests are a direct continuation of the Reference (1) effort. A phenomenological SCCGR correlation has been developed to describe the observed nickel alloy SCC dependency on dissolved hydrogen. This hydrogen SCC model, in association with Contact Electrical Resistance (CER) measurements of the Ni/NiO phase transition from 288 to 360°C permits the ability to extrapolate the SCCGR hydrogen functionality to lower temperatures.

EXPERIMENTAL PROCEDURE

The nickel alloys investigated in this study were mill-annealed alloy 600, alloy X-750 condition HTH, and alloy X-750 condition AH. The compositions of these alloys have been provided in Reference (1). Alloy 600 was fabricated into standard 25.4 mm compact tension (CT) specimens with 10% side grooves and alloy X-750 was fabricated into standard 10.2 mm CT specimens. All specimens were fabricated in a longitudinal-transverse (LT) orientation. Test specimens were air fatigue precracked, to a nominal a/W of 0.5 for alloys 600 and X-750 AH and 0.45 for alloy X-750 HTH, using the fatigue precracking procedure from ASTM E399 Annex 2. The alloy X-750 tests were conducted under pure constant load test conditions whereas a 10% daily unload was performed in the alloy 600 tests.
The initial nominal stress intensity factor ($K_I$) in the alloy X-750 AH tests was 27.5 MPa$\sqrt{m}$ and 49.4 MPa$\sqrt{m}$ for the alloy X-750 HTH tests. Alloy 600 tests were conducted at $K_I$ levels of 27.5 and 65.9 MPa$\sqrt{m}$\(^{(1)}\).

Constant load stress corrosion crack growth rate (SCCGR) tests were conducted in active load servoelectric machines with digital control electronics monitored through a computerized data acquisition system. A description of the recirculating high flow rate autoclave system has previously been described.\(^{(2)}\) Crack lengths were monitored in-situ using electrical potential drop (EPD) according to ASTM E647, Annex 3, and were verified by post-test destructive examinations (DE). Alloy 600 EPD measured active voltages (proportional to crack extension) were normalized for alloy 600 resistivity changes measured in an unloaded non-precracked 10.2 mm CT specimen of the same heat. Since alloy X-750 resistivity does not change appreciably with time\(^{(3)}\), X-750 EPD resistivity normalization was not performed in tests with these materials.

**Test Environment**

The SCCGR test chemistries were deaerated high purity water with various levels of dissolved hydrogen. The feed tank solution was buffered to a room temperature pH of 10.1 to maintain a near neutral autoclave pH at the test temperature. Separate tests were performed at multiple hydrogen concentrations for each alloy (between 1 and 149 cc/kg). The desired hydrogen concentrations were obtained by varying the feed tank hydrogen overpressure according to Henry's law. A room temperature Henry's law coefficient of 5.9 kPa/(cc/kg) was employed. Cylinders of mixed gas containing 4% or 14.7% hydrogen, with the balance argon, were used to obtain hydrogen levels less than 20 cc/kg. Pure nitrogen gas was employed to obtain test conditions with very low hydrogen levels\(^{(2)}\). Since the autoclave turnover rate is relatively rapid (once every hour), autoclave effluent dissolved gas levels are analogous to feed tank levels, and the oxygen levels for all tests were <10 ppb.

Alloys 600 and X-750 AH tests were conducted at 338°C and alloy X-750 HTH tests were conducted at 360°C. Prior to the start of the alloys 600 and X-750 AH SCC tests, test specimens were prefilmed in the load train for 10 and 14 days, respectively, at $K_I = 4$ MPa$\sqrt{m}$ and 260°C at the hydrogen level of the SCC test. Alloy X-750 HTH specimens were not prefilmed. The differences in material prefilming procedures were a consequence of historical testing practices.

Trends in pH and contaminant levels were monitored through system in-line conductivity probes. Anion contaminant levels were verified as less than detectable (~25 ppb) by ion chromatography (IC) analysis of autoclave effluent samples. Autoclave cation contaminants were monitored through analysis of autoclave effluent samples by inductively coupled plasma (ICP) spectroscopy. Cation contaminant levels typically coincided with cation solubility limits (e.g., ~2 ppb Fe) at temperature.

Upon completion of SCC testing, alloy 600 and X-750 AH test specimens were heat-tinted at 454°C in air for 8 hours to facilitate fractography inspections. Following heat-tinting, the alloy 600 specimen underwent an air fatigue extension. This process prevents deformation of the SCC surface during post test fracture of the specimen. The higher strength 10.2 mm alloy X-750 CT specimens, as well as the fatigue extended alloy 600 specimen, were ultimately pulled apart in tension to expose the fracture surfaces. The amount of SCC in each specimen was measured by optical examination using the procedure in ASTM E813.

\(^{(1)}\) K validity size requirements are not met for a 25.4 mm CT specimen at the "pseudo" stress intensity factor of 65.9 MPa$\sqrt{m}$ (i.e., the specimen is too small). The fact that K validity is not met under this condition will not be highlighted throughout this document.

\(^{(2)}\) In a "nitrogen sparge" X-750 AH SCC test (338°C) the dissolved hydrogen level was estimated at ~0.08 cc/kg via an Orbisphere hydrogen analyser.
RESULTS

SCCG Analysis

Electrical potential drop (EPD) indicated immediate SCC growth at the start of all alloy 600 and X-750 condition AH tests. In a large percentage of the X-750 condition HTH tests an appreciable crack growth rate incubation period (i.e., dwell time prior to the establishment of a constant crack growth rate) was noted by EPD (up to 100 hours). Figure 2 provides a representative example of EPD indicated crack growth rate incubation. Actual crack extensions were consistently underpredicted by EPD by a factor ranging from 1.5 to 3. This under-prediction occurs consistently for materials with branched crack fronts and results from an uncracked metal ligament conduction path. EPD would have dramatically over predicted the actual alloy 600 crack extension if material resistivity increases were not taken into consideration. Because of these EPD limitations, reported crack growth rates were determined by dividing the actual destructive examination (DE) intergranular SCC extension by the portion of the test in which cracking occurred (i.e., test duration-incubation time).

It was hypothesized that X-750 HTH SCCGR incubation time would correlate inversely with the crack growth rate. That is, if the environment (in this study hydrogen level) was conducive to accelerate crack growth rates then this environment should also tend to minimize the SCC incubation period. Figure 3 provides a plot of measured X-750 HTH incubation times vs. SCC growth rate (SCCGR) for data generated in this study (i.e., ~0 to 150 cc/kg dissolved hydrogen) and some recently conducted tests as a function of pH (3.8 to 8.6 at 360°C) and in heavy water (D₂O) with 50 cc/kg dissolved hydrogen. All of these X-750 HTH tests were conducted at 360°C with a stress intensity factor of ~49 MPa.m. Figure 3 illustrates that the X-750 HTH incubation data was comprised of two populations: one group of specimens which had no discernible incubation period for a wide range of crack growth rates and a second set in which the incubation time was weakly correlated with the crack growth rate. Interestingly, a lack of crack growth rate incubation was associated with 4 out of the 5 slowest X-750 HTH crack growth rates. Note the unusually large incubation times of the X-750 samples tested in D₂O relative to those tested in light water. This observation may be related to the formation of a more perfect passive oxide in D₂O.

Influence of Hydrogen on SCCGR

Dissolved hydrogen crack growth rate test results for each of the three materials investigated in this study are summarized in Figures 4, 5, and 6. These figures illustrate the dramatic (up to 7x) effect of dissolved hydrogen concentration on the crack growth rate of the three nickel alloys investigated in this study. Each of these figures includes plots of the measured crack growth rate as a function of (1) the hydrogen level and (2) the electrochemical corrosion potential (ECP) distance away from the nickel to nickel oxide phase transition (ECPₐlNiO-ECP). Since the corrosion potential of nickel alloys behaves as hydrogen electrodes in hydrogenated water, this ECP difference reduces simply to the difference of two hydrogen electrodes:

\[
ECP_{NiO/Ni} = \Delta ECP = 1000(RT/2F)\ln([H_2]_{test}/[H_2]_{NiO}) \quad \text{in mV} - 1
\]

In equation (1), R is the gas constant [8.314 J/(mol·K)], F is Faraday’s constant (96,500 C/eq), T is the absolute temperature (K), [H₂]_{test} is the test hydrogen concentration, and [H₂]_{NiO} is the hydrogen concentration of the Ni/NiO phase transition at the given temperature in the same units as [H₂]_{test}. In these figures, \( \Delta ECP \) was improved

(1) EPD X-750 HTH crack growth rate incubation often appeared as a region of slow growth prior to steady crack growth rate. A significant portion of this slow growth is believed to be an EPD artifact due to small X-750 resistivity changes that were not accounted for in this study.

(2) In a Pourbaix diagram the Ni/NiO phase transition (single line) and hydrogen electrodes (multiple lines as a function of H₂ level) have the same slope. These lines are common at the hydrogen concentration of the Ni/NiO phase transition.

(3) 13.8 and 25 cc/kg were employed as hydrogen concentrations of the nickel/nickel oxide phase transition at 338°C and 360°C, respectively. The 360°C value is based upon contact electrical resistance (CER) measurements and the
by subtracting the measured specimen to platinum EcP measurement. This modification corrects the specimen EcP for deviations from that of a true hydrogen electrode. This correction becomes appreciable at hydrogen concentrations less than ~5 cc/kg.

For each of these three materials, a maximum in crack growth rate occurred in proximity to the Ni/NiO phase transition. The maximum SCCGR for alloy X-750 condition HTH coincides almost exactly with the CER measurement of the Ni/NiO phase transition, Figure 4. The maximum SCCGR for X-750 AH was ~30 mV into the nickel oxide regime (Figure 5) and maximum for alloy 600 was ~15 mV into the nickel metal regime. Besides the precise location of the maximum susceptible to SCC, with respect to the hydrogen concentration, the magnitude (peak height) and breadth of the effect (peak width) also varied slightly between the three materials. That is, for X-750 HTH, a 5x (1.71 mils/day/0.35 mils/day) maximum SCCGR effect was observed and the region in which variation in hydrogen level accelerated the SCC extended over 80 mV, Figure 4. For alloys 600 and X-750 AH these values were 3x, 95 mV and 7x, 150 mV respectively, Figures 5 and 6.

The environmental SCCGR influence of dissolved hydrogen, as expected, was not appreciably influenced by the stress intensity factor (mechanical effect) in this study. As illustrated in Figure 6, the effect of hydrogen level on alloy 600 crack growth rate is extant, location and breadth) was not appreciably different at the two stress intensity factor levels of 27.5 and 65.9 MPa/m. The lack of an interaction between the hydrogen functionality and the stress intensity factor is significant. This result indicates that testing to describe the influence of hydrogen level on nickel alloy SCC need not be conducted as a function of the stress state.

**Phenomenological Correlations**

A correlation of the observed crack growth rate hydrogen functionality (Figures 4-6) was developed in terms of the fundamental independent parameter \( EcP_{N_{i}O}-EcP (\Delta EcP) \) and a symmetric exponential function as illustrated in equation (2).

\[
\text{SCCGR/SCCGR}_{\text{min}} = \left( \frac{\text{SCCGR}}{\text{SCCGR}_{\text{min}}} \right) \exp \left\{ -\frac{1}{2} (\Delta EcP + \Delta EcP_{a})/\lambda \right\}^2
\]

SCCGR\(_{\text{max}}/\text{SCCGR}_{\text{min}} \) is the ratio of the observed maximum to minimum crack growth rate for a given material. This ratio describes the magnitude of the SCCGR dissolved hydrogen level effect for a specific material (e.g., ~5x for X-750 HTH, see Figure 4). \( \Delta EcP_{a} \) is a fitted parameter which describes the extent that the maximum crack growth rate is offset from \( EcP=0 \) (i.e., the location of the Ni/NiO phase transition). \( \lambda \) is another fitting parameter, which describes the breadth of SCCGR vs. EcP peaks. Values for \( \Delta EcP_{a} \) and \( \lambda \) were obtained by linear regression analysis of a linearized version of equation (2). Resulting values of \( \Delta EcP_{a} \) and \( \lambda \) are given below for each material tested. Figure 7 illustrates that these correlations describe the observed nickel alloy dissolved hydrogen crack growth rate functionality.

<table>
<thead>
<tr>
<th>Test Condition</th>
<th>SCCGR(<em>{\text{max}}/\text{SCCGR}</em>{\text{min}} )</th>
<th>( \Delta EcP_{a} )</th>
<th>( \lambda )</th>
<th>Range of Applicability(^{(4)} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>X-750 HTH 360°C</td>
<td>4.89</td>
<td>3.0</td>
<td>20.7</td>
<td>-40 mV &lt; ( \Delta EcP &lt; 39 ) mV</td>
</tr>
<tr>
<td>X-750 AH 338°C</td>
<td>7.19</td>
<td>29.9</td>
<td>41.8</td>
<td>-100 mV &lt; ( \Delta EcP &lt; 50 ) mV</td>
</tr>
<tr>
<td>A600 338°C</td>
<td>2.81</td>
<td>-8.9</td>
<td>35.0</td>
<td>-40 mV &lt; ( \Delta EcP &lt; 55 ) mV</td>
</tr>
</tbody>
</table>

338°C measurement is based upon CER measurements as well as corrosion coupon data which indicates that 12.5 cc/kg H₂ is within the NiO regime and that 20 cc/kg is within the Ni regime at 338°C.

\(^{(4)}\) If the nitrogen sparge crack growth rate (0.069 mils/day at <0.1 cc/kg) was employed as the minimum observed SCCGR this effect would be 25x. This crack growth rate could not be plotted in Figure 4b since measurement of the specimen to Pt EcP was not available. This EcP measurement is required at low hydrogen levels (< 5 cc/kg) to correct for deviations from the potential of a hydrogen electrode.

\(^{(7)}\) The SCCGR effect of dissolved hydrogen level appears to have an appreciable influence in a narrow range of EcP about the Ni/NiO phase transition.
To evaluate this hypothesis, repassivation tests and Electron Spectroscopy Chemical Analysis (ESCA) corrosion measurements were performed as a function of aqueous dissolved hydrogen. Repassivation results could not discern a metal corrosion hydrogen functionality because of a hydrogen oxidation interference. Figure 9 is a summary of the ESCA corrosion film thickness results. ESCA oxide film thickness determinations were made on external surfaces of three alloy X-750 HTH CT specimens tested as part of this study. As illustrated in Figure 9, ESCA average oxide film thickness did not exhibit the same functionality that was observed with SCC with respect to the dissolved hydrogen level. X-750 HTH 360°C SCC crack growth is ~20 times slower under nitrogen sparge test conditions compared to 14 cc/kg test conditions. However, the nitrogen sparge oxide film thickness was approximately twice as thick as the 14 cc/kg oxide film (6500 compared to 2820 angstroms).

As noted in Reference (15), the lack of correlation between SCC and bulk oxide film thickness suggests that either: 1) bulk surface corrosion is not an accurate measure of the corrosion kinetics that occurs at the crack tip or 2) the functionality of SCC with respect to the coolant hydrogen level is unrelated to a corrosion functionality. The authors If the location of Ni/NiO phase transition is known, equation (2) can be extrapolated to lower temperatures through the assumption that the magnitude of the dissolved hydrogen effect is temperature independent. CER measurements (288 to 360°C) of the Ni/NiO phase transition were extrapolated to lower temperatures through a two parameter thermodynamic model. The Gibbs free energy (ΔG=ΔH298-TΔS298) of the phase transition reaction (Ni+H2O→NiO+H2) is equal to the -RTln(P_H2) where R and T are the quantities defined in equation (1) and P is the hydrogen concentration in atmospheres. ΔH298 and ΔS298 are the enthalpy and entropy of formation for the phase transition reaction. ΔH298 and ΔS298 were empirically determined to be 20.1 kJ and 0.011 kJ/K, respectively, by linear regression analysis of the CER data (i.e., plotting ln(P_H2) vs. 1/T). The resulting thermodynamic correlation for P_H2 is provided below.

P_H2=exp[-(20,100-11T)/RT] (3)

Figure 8 presents model predictions for the coolant hydrogen crack growth rate functionality of alloys 600 and X-750 condition HTH at 288°C with 10 cc/kg dissolved hydrogen based upon the Ni/NiO phase transition of 4.7 cc/kg H2 and the assumptions previously identified. This figure suggests a 2.7x influence of 10 cc/kg dissolved hydrogen at the lower temperature of 288°C. For alloy 600 at 288°C, 10 cc/kg H2 is close to the SCCGR maximum. For alloy X-750 HTH this condition yields a hydrogen SCCGR effect which is approximately midway between the maximum and the nickel metal region minimum.

General Corrosion Measurement

A maximum in nickel alloy SCC susceptibility has been observed in a narrow (~100 mV) region near the Ni/NiO phase transition. Increased SCC near an oxide phase transition is often considered a classic sign of a corrosion-controlled SCC mechanism. Since corrosion is responsible for crack advance in the film rupture oxidation (FRO) mechanism and is believed to be the main source of detrimental hydrogen in a hydrogen assisted crack (HAC) mechanism, both mechanisms as applied to low electrochemical corrosion potential SCC (LPSCC) are essentially corrosion-driven mechanisms. Specifically, if either the FRO or the HAC mechanism is the dominant LPSCC mechanism, then metal corrosion should exhibit a maximum in proximity to the Ni/NiO phase transition (i.e., the same functionality with respect to aqueous hydrogen concentration as exhibited by LPSCC).

To evaluate this hypothesis, repassivation tests and Electron Spectroscopy Chemical Analysis (ESCA) corrosion measurements were performed as a function of aqueous dissolved hydrogen. Repassivation results could not discern a metal corrosion hydrogen functionality because of a hydrogen oxidation interference. Figure 9 is a summary of the ESCA corrosion film thickness results. ESCA oxide film thickness determinations were made on external surfaces of three alloy X-750 HTH CT specimens tested as part of this study. As illustrated in Figure 9, ESCA average oxide film thickness did not exhibit the same functionality that was observed with SCC with respect to the dissolved hydrogen level. X-750 HTH 360°C SCC crack growth is ~20 times slower under nitrogen sparge test conditions compared to 14 cc/kg test conditions. However, the nitrogen sparge oxide film thickness was approximately twice as thick as the 14 cc/kg oxide film (6500 compared to 2820 angstroms).

As noted in Reference (15), the lack of correlation between SCC and bulk oxide film thickness suggests that either: 1) bulk surface corrosion is not an accurate measure of the corrosion kinetics that occurs at the crack tip or 2) the functionality of SCC with respect to the coolant hydrogen level is unrelated to a corrosion functionality. The authors

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(8) P_H2 is converted to the more common hydrogen concentration units of cc/kg through the use of an applicable Henry's law coefficient. With the approach employed in this analysis to extrapolate Ni/NiO phase transition measurements to lower temperatures all thermodynamic corrections were lumped into ΔG.

(9) The assertion that corrosion must be the dominant source of detrimental hydrogen if a HAC mechanism is responsible for SCC is supported by the dissolved hydrogen functionality observed in this study. If hydrogen dissolved in the water (i.e., aqueous hydrogen) was the dominant source of detrimental hydrogen in the metal, one would expect SCC susceptibility to increase monotonically with the aqueous hydrogen concentration, rather than exhibiting a maximum in SCC susceptibility.
view the former argument as being more likely. Steady state bulk surface corrosion kinetics are typically a function of the square root of time due to a process controlled by diffusion through the protective oxide. In contrast, at a crack tip where periodic rupture can occur the kinetics of initial film formation are likely controlling the corrosion.

**Metal Hydrogen Pickup Measurements**

If the HAC mechanism is the dominant LPSCC mechanism, then the buildup of detrimental hydrogen within the material should exhibit the same aqueous hydrogen concentration functionality as observed in LPSCC. In order to investigate this theory, hydrogen concentration measurements have been made on both stressed and unstressed samples of Alloy 600, Alloy X-750 HTH and on high purity nickel. The global hydrogen concentrations were quantified via thermal extraction (i.e., LECO). Results from the LECO measurements are summarized in Figure 10. These results illustrate that the global hydrogen concentration does not follow the same aqueous hydrogen functionality as observed in LPSCC. Consistent with Reference (16), increased absorbed hydrogen levels \(^{(10)}\) were not observed in the non-stressed specimens.

The SCC regions of stressed samples were investigated via nuclear reaction analysis (NRA), secondary ion mass spectroscopy (SIMS), and LECO measurements. No clear evidence of increased hydrogen concentrations near stress corrosion cracks were observed from these tests. While NRA measurements suggested that locally high (~100 wt. ppm) hydrogen concentrations may exist near the SCC surface, the relatively thick oxides (100's of nm), shallow penetration depth of the NRA beam (~0.5 μm), and the likelihood of adsorbed water or hydrated oxides on the surface confounded determination of the true hydrogen concentration in the metal. Analysis of the fatigue precrack, SCC, and post-test fracture regions of compact tension specimens of Alloy 600 and X-750 tested in \(D_2O\) via SIMS were also inconclusive. Evidently, the dissolved hydrogen concentrations in these regions are below the ~100 ppm detection limit of the SIMS system employed.

Lastly, sectioning of the SCC region of X-750 HTH compact tension specimens and analysis via LECO did not show any hydrogen enrichment (2-4 wt. ppm), solid circles in Figure 10. Note that this is in contrast to previous work on X-750 and Alloy 600, in which, locally high hydrogen concentrations (~43 and 83 wt. ppm) have been reported\(^{18,19}\). One possible explanation for this discrepancy is that the research that reported elevated hydrogen concentrations near SCC\(^{18,19}\) were from constant extension rate (CERT)-type samples rather than a compact tension specimen. It is possible that the CERT specimens contained a greater SCC crack area per volume of hydrogen sample and facilitated the measurements. Additional investigations into the concentration of hydrogen near a stress corrosion crack tip are critical to assessing the viability of the HAC mechanism.

**CONCLUSIONS**

1. Testing has identified a dramatic (up to 7x) effect of dissolved hydrogen level on the crack growth rate of nickel alloys. This hydrogen crack growth rate phenomena has been fundamentally described by the extent that the alloy's corrosion potential deviates from the corrosion potential of the Ni/NiO phase transition. Crack tip oxides are often of the nickel oxide structure and this corrosion potential difference represents the relative stability of the SCC controlling oxides. A phenomenological model has been developed to extrapolate the hydrogen functionality to lower temperatures.

2. The determination of an accurate thermal activation energy (Q) requires that all fundamental parameters, other than temperature, are maintained constant. The knowledge that the SCC hydrogen functionality is controlled by proximity to the Ni/NiO phase transition not the specific aqueous hydrogen level suggests that SCC thermal activation tests should actually be conducted within a single oxide stability region at all temperatures.

\(^{(10)}\) Increased absorbed hydrogen levels imply hydrogen values which are greater than would be expected from Sieverts law solubility expectations.
3. The environmental SCCGR influence of dissolved hydrogen was not appreciably influenced by the stress intensity factor (mechanical effect) in this study. The lack of an interaction between the hydrogen functionality and the stress intensity factor indicates that testing to describe the influence of hydrogen level on nickel alloy SCC need not be conducted as a function of the stress state.

4. The buildup of hydrogen within nickel alloy does not exhibit the same functionality as SCC with respect to the aqueous hydrogen concentration. This result implies that if hydrogen assisted cracking (HAC) is the dominant SCC mechanism, then isolated active path regions (i.e., surface initiation sites or cracks) must be actively corroding and must be the dominant source of elevated detrimental hydrogen fugacities. However, increased hydrogen levels were not measured in the SCC region of X-750 specimens.

5. Alloy X-750 HTH ESCA bulk surface corrosion film thickness did not exhibit the same functionality that was observed with SCC with respect to the hydrogen level. This observation suggests that either:
   (1) bulk surface and crack tip corrosion kinetics are different, or
   (2) the functionality of SCC with respect to the hydrogen level is unrelated to a corrosion functionality.

ACKNOWLEDGEMENTS

The following individuals are recognized for their contribution to this effort: Mark Ando for his assistance in test design and diligence in test facility operation and Mike Hanson for his alloy X-750 ESCA analyses.

REFERENCES

Figure 1: Effect of Dissolved Hydrogen on X-750 HTH 360°C Crack Growth Rate

![Figure 1: Effect of Dissolved Hydrogen on X-750 HTH 360°C Crack Growth Rate](image1)

- Nickel Metal Stability
- HTH Stability

Figure 2: EPD indicated Crack Growth Rate Incubation in a 360°C X-750 Condition HTH Test at 10 cc/kg and K=51.5 MPa/m

![Figure 2: EPD indicated Crack Growth Rate](image2)

Figure 3: X-750 HTH SCC Incubation Time vs. Crack Growth Rate (360°C, K~49 MPa/m)

![Figure 3: X-750 HTH SCC Incubation Time vs. Crack Growth Rate](image3)
25 cc/kg was employed as the hydrogen concentration of the nickel/nickel oxide phase transition at 360°C. The 360°C value is based upon contact electrical resistance (CER) measurements. 13.8 cc/kg was employed as the hydrogen concentrations of the nickel/nickel oxide phase transition at 338°C. This 338°C value is based upon CER measurements as well as corrosion coupon data which indicates that 12.5 cc/kg hydrogen is within the NiO regime at 338°C.
Figure 7: Phenomenological Correlation of Alloy X-750 Condition HTH Coolant Hydrogen Crack Growth Rate Functionality at 360°C

![Graph showing SCCGR proportionality to EcP_Ni/NiO - EcP (mV).

SCCGR/SCCGR_{\text{max}} = SCCGR_{\text{max}}/SCCGR_{\text{min}} \exp\left(-\frac{1}{2}(\Delta EcP + 3.0)/20.7\right)

SCCGR_{\text{max}}/SCCGR_{\text{min}} = 1.71 \text{ mils/day}/0.35 \text{ mils/day}

3.0 and 20.7 are fitted parameters

The correlation is valid in the approximate range of -40 mV < \Delta EcP < 39 mV

Figure 8: The Effect of Dissolved Hydrogen on Alloys 600 and X-750 HTH SCCGR Extrapolated Down to 288°C

![Graph showing SCCGR proportionality to EcP_Ni/NiO - EcP (mV).

Figure 9: Nickel Alloy SCC and ESCA Oxide Film Thickness as a Function of Dissolved Hydrogen Concentration

![Graph showing oxide film thickness vs. hydrogen concentration for various alloys.

Figure 10: Nickel Alloy Metal Hydrogen Pickup as a Function of Dissolved Hydrogen Concentration

![Graph showing metal hydrogen pickup vs. aqueous hydrogen level for various alloys.

a) X-750 HTH, 360°C, 49 MPa\sqrt{m}

b) X-750 HTH, 360°C, 28 day ESCA determined oxide film thickness vs. H₂