In-situ Raman Spectroscopic Study of Oxide Films on Alloy 600 in Simulated PWR Water

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Abstract

Although there has been no general agreement on the mechanism of intergranular stress corrosion cracking (IGSCC) as one of major degradation modes of Ni-base alloys in PWR’s, common belief derived from previous studies is that the damage to the alloy substrate is related to mass transport characteristics and/or repair properties of overlaying oxide film. Recently, it was shown that the oxide film structure and IGSCC initiation time as well as crack growth rate were systematically varied as a function of hydrogen partial pressure in high temperature water, providing supporting evidences. This study is aimed at demonstrating a capacity to characterize the oxide film by an in-situ Raman spectroscopy as a function of PWR primary water conditions in order to understand how the oxide film chemistry can vary with water chemistry and participate in the cracking process as a key variable. This is achieved by analyzing the oxide film properties obtained for alloy 600 in various conditions leading to different IGSCC susceptibility. The in-situ Raman spectroscopic information obtained for various oxide films on alloy 600 in primary water of PWR as a function of hydrogen partial pressure is compared with thermodynamic predictions.

1. Introduction

Primary water stress corrosion cracking (PWSCC) is one of major degradation modes that have been observed to occur pre-dominantly along grain boundaries in Ni-base structural materials for steam generators (SG’s) or nozzles in the primary circuit of nuclear power plants. Such an intergranular stress corrosion cracking (IGSCC) has been observed under specific combinations of materials and environmental conditions. The PWSCC of alloy 600 is directly related with safety concerns of nuclear power plant from the viewpoint that the SG tubings or nozzles may rupture as result of the cracking although alloy 600 components are protected against rupture by relatively large safety margins. Also, it has been intensified in recent years as Ni-base alloys display the increasing number of IGSCC events in components.
that are critical to the life extension of nuclear power plants.

The PWSCC phenomenon of alloy 600 has been extensively studied in order to formulate remedies and to establish theoretical and/or empirical models for predicting the crack initiation time and crack growth rates for field components. Several theoretical and semi-empirical models, based on slip dissolution/oxidation, corrosion-enhanced plasticity, internal oxidation, and creep have been proposed. Even though there is no general agreement on the mechanism of IGSCC, one common postulation from these studies is that the damage to the alloy substrate can be related to mass transport characteristics, rupture and/or repair properties of overlaying oxide film.

Earlier studies have shown that dissolved hydrogen concentration in high temperature water affects the PWSCC susceptibility of nickel base alloys. Test results have demonstrated that the maximum of crack growth rate, with respect to dissolved hydrogen concentration, is observed in proximity to a key phase transition between the nickel metal (Ni) and nickel oxide (NiO). Fig. 1 summarizes Influence of hydrogen concentration on the crack growth rate of PWSCC in alloy 600 at 360°C. As shown in Fig. 1, the region of degradation by PWSCC is located at transition of nickel metal (Ni) to nickel oxide (NiO) phase. Recently, it was also shown that the oxide film structure, PWSCC initiation time and crack growth rate were systematically varied as a function of hydrogen partial pressure in high temperature water. These are evidences that the oxide film on the surface of alloy 600 in the primary water plays a key role in the process of PWSCC.

This study is aimed at characterizing the oxide film by in-situ Raman spectroscopy under PWR primary water conditions in order to understand how the oxide film behaves in the cracking process as a key element. This can be done by analyzing oxide film properties obtained in various conditions exhibiting different PWSCC susceptibility. The in-situ Raman spectroscopic information is to be acquired in this study from the oxide film of alloy 600 metal surfaces in simulated PWR primary water conditions through a hermetically sealed window. The system developed in this study is capable of examining the effect of temperature, dissolved hydrogen concentration, and impurity concentration on chemistry and kinetics of film formation at temperature of up to 350°C and pressure of up to 18 Mpa. On the theoretical side, thermo-chemical calculation has been made to identify the thermodynamically stable oxide phases and to compare with observed results in this work.

2. Experimental

2.1. Materials

The alloy 600 used in this work for analysis of the oxide films have been produced by Sumitomo Metals, Inc., for International Cooperative Group of Environmentally Assisted Cracking (ICG-EAC) round robin test program following the specification of a reactor pressure vessel (RPV) head penetration materials for control rod drive mechanisms (CRDMs) in PWRs.

Chemical composition of alloy 600 used in this study is given in Table 1. The material was solution-annealed at 1050°C for 2 hours and then water-cooled.

The microstructure showed particularly solutionized carbides with an average grain size of
about 50 μm.

2.2. Specimens for Raman Spectroscopic Measurement

The in-situ Raman specimen for aqueous corrosion was machined of alloy 600 into a round disk with 7 mm diameter and 1 mm thickness. All the specimens were mechanically polished down to 1μm Al₂O₃ powder, then rinsed with ethanol, and finally with deionized water prior to an installation for exposure.

For measurements of ex-situ spectra in the air, the commercial high purity powders (Johnson Matthey) of NiO (99.998%), NiFe₂O₄ (99%), Cr₂O₃ (99.998%), and NiCr₂O₄ (90%) were used to obtain reference spectra for aqueous corrosion of alloy 600. Average particle size was 44μm for NiO, for 200 μm NiFe₂O₄, 800μm for Cr₂O₃, and 150μm for NiCr₂O₄. Powder color was dark green for NiO, brown for NiFe₂O₄, and green for Cr₂O₃ and NiCr₂O₄. Powder samples were placed in the recess of sample holder, mixed with deionized water, and then dried prior to deposition on the sample holder.

2.3. Test Conditions

For the surface oxide formation, the alloy 600 specimens were placed inside a high-pressure optical cell and exposed to deionized and deaerated water environments. The same chemical condition as typical PWR primary water was used. Water with a resistivity of 18 MΩ-cm was mixed with additional chemicals to produce 1,000 ppm boron in the form of boric acid(H₃BO₃) and 2 ppm lithium in the form of lithium hydroxide (LiOH).

Two different levels of dissolved hydrogen gas (30 cm³(STP)/kg or 1 cm³(STP)/kg) concentration were prepared with corresponding dissolved hydrogen concentrations of 2.68 ppm and 0.89 ppm, respectively, according to the solubility data. The operating pressure was about 18 Mpa and the temperature was varied from room temperature to 350°C.

2.4. Apparatus for In-situ Raman Measurement

The optical cell for the in-situ observation at high temperature and pressure condition was constructed with a custom-designed 1 liter-capacity autoclave. In order to reduce corrosion products from the surfaces of construction materials at high temperature, the cell was made of alloy 690. For the same reason, the head of a water-charging pump was made of titanium and balls of check valves were made of sapphire. All other components exposed to high temperature water including compression fittings and tubings were made of alloy 600. The nominal flow rate to the cell was maintained at 4 liters per hour during experiments. The cell was machined with two penetrations in the wall of cylinder to place optical windows for the access of Raman spectroscopy. The initial window assembly, composed of 4.5 mm thick sapphire with gaskets to seal high-pressure water, was designed after Hurst et al.’s work. In this work, gaskets were made of Ni-base alloy 718 electroplated with gold to 25 μm thickness. Because the optical surface of the sapphire was degraded by pitting corrosion in high
temperature water during the study, a 0.25 mm thick CVD diamond disk was placed on the water side of the sapphire window and this has greatly reduced surface attack on the sapphire window. The alloy 600 specimen for aqueous corrosion was held in the recess of an oxidized zirconium metal by a washer made of alloy 718 with the gasket of an oxidized zirconium sheet so as to electrically insulate the specimen for the cell.

The Raman spectroscopy system is consisted of an excitation laser source, a spectrometer and optical components including mirrors and filters. Fig. 2 shows the layout drawing of optical system with near backscattering geometry including optical cell and water chemistry loop used in this work. A more detailed description for the Raman system is given elsewhere.29

3. Results and Discussions

3.1. Morphology of Oxide Film on Alloy 600

Fig. 3 shows the morphology of the surface oxide film observed by a field emission scanning electron microscope (FE-SEM) for alloy 600 exposed to 350°C PWR water for 110 hours. The surface oxide film is composed of a compact layer by small crystallites of few tens of nm in diameter. Some larger crystallites of about 200 nm in diameter are present on the compact layer.

Fig. 4 shows the transmission electron micrographs of a cross-section of alloy 600 specimen oxidized in simulated PWR water with 30 cm³(STP)/kg of dissolved hydrogen concentration for 71 hours. The thickness of oxide film on alloy 600 was observed to be in the range of 12 ~ 180 nm. Compact layers and other crystalline phases on alloy 600 observed by TEM have the same morphology as those in the FE-SEM micrographs.

3.2. Reference Raman Spectra for High Purity Oxide Powders

The first series of experiments were conducted to obtain reference Raman spectra on oxides using powder samples. Results were compared with literature data in order to verify the developed system. The results of experiments with nickel and chromium oxides and spinels in room temperature air environment are summarized in Table 2. As shown in Table 2, the Raman features of the reference powder spectra for NiCr₂O₄ powder show good agreements between the measurement in this work and literature data. Further details of results are given elsewhere29.

3.3. In-situ characterization

3.3.1. Aqueous Corrosion in PWR Water with 30 cm³(STP)/kg of Dissolved Hydrogen Concentration

To maintain the dissolved hydrogen concentration (DH₂) of 30 cm³(STP)/kg in the
simulated PWR water, pure hydrogen gas was injected into the water storage tank with the
cover gas pressure of 170 kpa. Based on Henry’s law, a dissolved hydrogen concentration of
2.68 ppm is expected. The operating pressure was about 18 Mpa and the temperature was
gradually increased from room temperature to 350°C at an average rate of 40°C/hour.
Temperatures at which Raman spectra were collected with the remained time length are
summarized in Table 3.

Fig. 5 shows the in-situ Raman spectra of alloy 600 specimen obtained by holding
temperatures at 250, 290, 320, and 350°C, respectively, while temperature increased. Peaks
observed at ca. 417 cm⁻¹, 743 cm⁻¹ in the in-situ Raman spectra were originated from the
sapphire window. A feature in the range of ca. 546-587 cm⁻¹ in the spectrum acquired at
250°C was known to be attributed to α-CrOOH based on Maslar et al.’s work. According
to the explanation described in Ref. 19, [CrIII06] octahedral are common structure to both Cr2O3
and α-CrOOH. Therefore, one might reasonably expect the Raman wavenumbers of α-
CrOOH to be observed in approximately the same wavenumber range as those of Cr2O3, i.e.,
ca. 300-613 cm⁻¹.

The crystallographic structure of the Cr2O3·nH2O powder is yet unknown. The Cr2O3·nH2O XRD pattern does not match any chromium hydrate/hydroxide reference patterns. The broad XRD peaks are typical of a hydrate consisting of a mixture of different metastable hydrates/hydroxides. This result is consistent with a report on the Cr2O3·nH2O possessing variable water content. While the observed Raman peaks for Cr2O3·nH2O cannot be attributed to a particular structural characteristic, the Cr2O3·nH2O Raman spectrum illustrates that the Raman peaks of some hydrated CrIII oxide are observed in approximately the same wavenumber range as those of Cr2O3. Additionally, the infrared (IR) spectrum of α-
CrOOH exhibits features in this wave number range, i.e., peaks at 525 and 610 cm⁻¹. Therefore, the shared structural features of Cr2O3 and α-CrOOH and the wavenumber range of the Raman features in the Cr2O3 and Cr2O3·nH2O reference spectra are consistent with
the assignment of the measured Raman features to α-CrOOH.

Several intense peaks are also observed in the 840-880 cm⁻¹ range of spectra in Fig. 5. According to the extensive discussion by Maslar et al., this feature would be attributed to CrVI or crystalline CrIII/CrVI species assuming a chromium species is responsible for this feature. This explanation can account for the weak signature observed in the 340-350 cm⁻¹ range in the spectra of the specimen at 250°C. Chromium oxide features in this wavenumber range are generally attributed to CrVI –oxygen terminal stretching modes or mixed CrIII/CrVI oxide vibrational modes. Hydrated surface chromate species have been reported to exhibit a vibrational mode at ca. 865 cm⁻¹. The most intense peak in the aqueous HCrO4− Raman spectrum is reported from ca. 880 to 899 cm⁻¹. A number of mixed CrIII/CrVI oxides have been identified during the thermal decomposition of various chromium-containing materials. Crystalline compounds such as Cr3O6, Cr2O5, and XCr3O8 (X = Na, K, Rb) exhibit their most intense Raman spectral features in the ca. 820-904 cm⁻¹ range at room temperature. The most intense Raman spectral feature of an amorphous CrIII/CrVI compound has been observed at 859 cm⁻¹. In the spectra obtained at 250 and 290 °C, weak features at 704 cm⁻¹ appeared and these were attributed to nickel ferrite spinel (NiFe2O4) features.

In the spectra obtained at 320 °C, features of pure chromium oxide including α-CrOOH,
CrVI and crystalline CrIII/CrVI compounds become weaker, while nickel oxide features are firstly observed and become more apparent in subsequent spectra up to 350°C. The features at 550 cm⁻¹ and 910 cm⁻¹ were attributed to NiO phase. Also, features of nickel chromium spinel (NiCr₂O₄) were detected in this temperature range. The features at ca. 682 cm⁻¹ and 430 cm⁻¹ were attributed to NiCr₂O₄ phase. Also, nickel oxide and nickel chromite phase were still detected on the specimen surface as exposure time increases at 350°C. The NiCr₂O₄ is one of thermodynamically stable phases in a reducing aqueous environment with the range of hydrogen overpressure at about 300°C. A more detail study that was performed to identify thermodynamically stable phase of nickel base alloy in high temperature water was described in ref. 29.

3.3.2. Aqueous Corrosion in PWR Water with 1 cm³(STP)/kg of Dissolved Hydrogen Concentration

To maintain the dissolved hydrogen concentration of 1 cm³(STP)/kg in water, 5% hydrogen gas balanced with 95% helium gas was injected into the water with the overpressure of 0.7 atm and it made a dissolved hydrogen concentration of 0.89 ppm in this study. Temperatures at which Raman spectra were collected with the remained time length are summarized in Table 3.

Fig. 6 shows the in-situ Raman spectra of alloy 600 specimen obtained while temperature increased. A feature in ca. 546-587 cm⁻¹ range in the spectrum acquired at 250°C was attributed to α-CrOOH from the dame rationale as that described earlier in the result with DH₂=30 cc/kg. Also, the intense peaks feature that would be attributed to CrVI or crystalline CrIII/CrVI species was observed in the 840-880 cm⁻¹ range.

As temperature increases, clear features in 550 and 910 cm⁻¹ which were attributed to nickel oxide (NiO) were firstly appeared and became weaker in subsequent spectra up to 350°C. After 23 hours at 350°C in this condition, the features of nickel oxide phase were still detected on the alloy 600 specimen surface at 350°C.

3.3.3. Effect of Dissolved Hydrogen Concentration Change in PWR Water

To investigate the effect of dissolved hydrogen concentration on the formation of oxide film of alloy 600 in high temperature water, dissolved hydrogen concentration was increased from 1 to 30 cc/kg, immediately following the measurement at 1 cc/kg. In-situ Raman spectroscopic measurement of alloy 600 surfaces was made both before and after the change of hydrogen concentration.

Fig. 7 shows the change of Raman spectra with the increase of dissolved hydrogen concentration in water at 350°C. In high temperature water with 1 cc/kg of hydrogen concentration, features of pure chromium oxide including α-CrOOH, CrVI and crystalline CrIII/CrVI compounds and nickel oxide (NiO) features were observed at the oxide film of alloy 600 specimen. With the increase of dissolved hydrogen concentration, nickel chromium spinel (NiCr₂O₄) features were appeared in the observed Raman spectrum.
4. Discussion

Table 4 summarizes the comparison between experimental observations and prediction of oxide phases of alloy 600 at each condition of temperature and hydrogen concentration. It can be known that the consideration of formation of chromium oxide hydroxide (CrOOH) and nickel spinels can make differences in the thermochemical prediction results, regardless of dissolved hydrogen concentration in 350°C water. At both conditions of dissolved hydrogen concentration, nickel ferrite (NiFe₂O₄) phases which were predicted to be thermodynamically stable assuming that nickel spinels can be formed in this system were not experimentally detected. When the observed oxide phases are compared with the predicted stable phases of alloy 600 at 350°C, all the observed oxide phases but Cr-oxides are included in predicted stable phases. But the thermochemical predictions with CrOOH and Ni spinels do not agree with the observed results at both hydrogen concentrations in 350°C water. In contrast, the predicted Ni/NiO equilibrium line is in good agreement with the observed results. Therefore, it can be said that the oxide phases identified by in-situ Raman spectroscopic investigation can be used to verify thermodynamically stable phases.

Table 5 summarizes the experimental observations of surface oxide phases of alloy 600 in 350°C water in comparison with literature data⁹,¹²,³⁰ at each dissolved hydrogen concentration. At higher dissolved concentration, NiFe₂O₄ precipitates reported in ref. 10 and 30 could be made by Fe ion dissolution from high temperature flowing system including autoclave, tubes and fitting for longer exposure time than this thesis work. At lower dissolved concentration, the observed oxide phases agree well with those reported in literature.

From the viewpoint that electrochemical potential-pH diagrams can be useful in predicting thermodynamically stable phases, thermochemical calculations can provide very much meaningful information to explain the corrosion behavior of alloy 600 in high temperature water. Although the pH was not measured at high temperature and pressure during this investigation, this value can be estimated using aqueous thermochemical calculation softwares. In this work, pH values of primary water in PWR were calculated as a function of temperature using MULTEQ®.²⁸ Electrochemical potential value was calculated by using the Nernst equation assuming that the surface of alloy 600 behaves as a hydrogen electrode in the hydrogen-containing water environment. The potential of hydrogen electrode is given as below:

\[
E_{\text{H}^+/\text{H}_2}(T) = E_{\text{H}^+/\text{H}_2}^\circ (T) - 2.303 \frac{RT}{2F} \left[ \log f_{\text{H}_2} + 2 \text{pH} \right]
\]

\[
= -2.303 \frac{RT}{2F} \left[ \log f_{\text{H}_2} + 2 \text{pH} \right] \quad \text{vs. SHE (T)}
\]

From the results of in-situ Raman spectra of alloy 600 with temperature increase, dissolved hydrogen concentration is one of the most important experimental parameters in this study. While nickel oxide was observed at 320°C or higher at the higher hydrogen concentration, nickel oxide was observed at 290°C or higher at the lower hydrogen concentration. To compare experimental results with calculations, thermochemical calculations for alloy 600
were made as a function of temperature, as described in ref. 29. In this calculation, for simplicity, it was assumed that nickel oxide could be produced whereas spinel oxide could not form. In Fig. 8, dashed line represents a previous experimental result of nickel/nickel oxide transition using a contact electric resistance (CER) instrument. The deviation between present and previous study can arise from inaccuracy of thermochemical data at high temperature for calculation in this work or experimental uncertainty in previous work.

Fig. 8 summarizes the estimated electrochemical potential of alloy 600 for two different hydrogen concentrations in this experiment as a function of temperature. It can be seen that nickel oxide was in a thermodynamically stable state at 270°C or higher at the lower hydrogen concentration. NiO phase was observed to be stable at 320°C or higher at the higher hydrogen concentration.

In-situ Raman spectroscopy is shown to be a very useful experimental tool that can provide valuable information about oxide films and the SCC behavior of alloy 600 in high temperature aqueous environments. In this work, dissolved hydrogen concentration was experimentally known to play an important role on the formation of oxide films on the alloy 600 in high temperature water. Different compositions of oxides were produced at different condition of dissolved hydrogen concentration in high temperature water.

Previous studies showed that the dissolved hydrogen concentration of 30 cm³(STP)/kg at 360°C corresponded to the maximum susceptibility of alloy 600 to crack initiation. The present in-situ Raman spectroscopic study showed that the condition corresponds to Ni/NiO equilibrium line in agreement with thermochemical predictions. Based on the results, it can be concluded that the mixture of nickel and chromium spinel oxide on the surface of alloy 600 is more sensitive to the SCC initiation than nickel oxide on that in 350°C PWR water condition. Mechanistic understanding on why the condition leads to the higher susceptibility to SCC of alloy 600 in PWR water may require a more detailed study on the nanostructure of oxide films that is beyond the scope of this study.

5. Summary and Conclusion

In-situ Raman spectroscopic investigation has been conducted for a nickel-base alloy 600 exposed to deaerated water with 1,000 ppm boron and 2 ppm lithium at a pressure of 18 Mpa, temperatures ranging up to 350°C and dissolved hydrogen concentrations ranging from 1 to 30 cc/kg in an optically accessible cell. From this study, following conclusions are made:

1. An in-situ Raman spectroscopic system was developed successfully to provide valuable information for the oxide film of structural alloys in water reactor environments, as a function of temperature, water chemistry and stress.

2. Potential-pH diagrams for Ni-Cr-Fe alloy at high temperature water were predicted by thermochemical calculations considering both ternary alloying effect and spinel oxide formation.

3. At the lower dissolved hydrogen concentration (1 cm³(STP)/kg), nickel oxide(NiO) phase was detected on the specimen surface at temperature ranges between 290 and 350°C. Nickel oxide phase appeared to be stable over a long period of time (21 hours) at 350°C.

4. At the higher dissolved hydrogen concentration (30 cm³(STP)/kg), chromium oxide
hydroxide (CrOOH) phase was detected on the specimen surface at relatively low temperature ranging from 250 to 290. Also, features attributed to the phase of NiO and NiCr2O4 were observed at 320 or higher.

5. Thermochemical calculations on the formation of NiO agreed very well with the in-situ Raman observation results.

6. Irrespective of dissolved hydrogen concentration, CrOOH, Ni(Fe)Cr2O4, and NiFe2O4 were predicted to be stable in 350°C water, assuming the formation of nickel spinels. CrOOH, NiO and FeCr2O4 were predicted to be stable in 350°C water, assuming the absence of nickel spinels.

7. Based on the results of in-situ Raman spectroscopic measurement and thermochemical calculation of oxide film on alloy 600, the mixture of nickel and chromium spinel oxide on the surface of alloy 600 is more sensitive to the SCC initiation than nickel oxide on that in 350°C PWR water condition.

Acknowledgements

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References


Table 1. Chemical Composition of Alloy 600 Used in This Study (wt.%)  

<table>
<thead>
<tr>
<th>Element</th>
<th>C</th>
<th>Mn</th>
<th>Fe</th>
<th>S</th>
<th>Si</th>
<th>Cu</th>
<th>Ni</th>
<th>Cr</th>
<th>Al</th>
<th>Ti</th>
<th>Nb</th>
<th>P</th>
<th>B</th>
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<td>Compo.</td>
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<td>8.31</td>
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<td>0.16</td>
<td>0.36</td>
<td>0.04</td>
<td>0.09</td>
<td>0.02</td>
<td>0.01</td>
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</table>

Table 2. Nominal Raman Peak Wavenumbers for Oxide Powders\textsuperscript{a} Measured in Room Temperature Air Environment.

<table>
<thead>
<tr>
<th>Oxide</th>
<th>This work\textsuperscript{b}</th>
<th>Ref. 1\textsuperscript{c}</th>
<th>Ref. 2\textsuperscript{c}</th>
</tr>
</thead>
<tbody>
<tr>
<td>NiO</td>
<td>1,074 (^{15}) 910 (^{15}) 725 (^{15}) 532 (^{15}) 400 (^{15})</td>
<td>(1,074 (^{15}))</td>
<td>(490 (^{15}))</td>
</tr>
<tr>
<td>NiFe(_2)O(_4)</td>
<td>702 (^{17}) 654 (^{17}) 595</td>
<td>702 (^{17}) 655 (^{17}) 592 (^{17})</td>
<td>700 (^{18}) 655 (^{18})</td>
</tr>
<tr>
<td>Cr(_2)O(_3)</td>
<td>610 (^{19}) 550 (^{19}) 528 (^{19}) 352 (^{19}) 302 (^{19})</td>
<td>(613 (^{19}))</td>
<td>585 (^{19})</td>
</tr>
<tr>
<td>NiCr(_2)O(_4)</td>
<td>687 (^{17}) 550-560</td>
<td>(685 (^{17}))</td>
<td>(686 (^{18}))</td>
</tr>
</tbody>
</table>

\textsuperscript{a}The most intense peak(s) in each spectrum is underlined.  
\textsuperscript{b}The wavenumber of a well-resolved peak has an associated uncertainty of \(\pm 2\) cm\(^{-1}\).  
\textsuperscript{c}The excitation laser with 647.1nm wavelength was used.
Table 3. Temperature and Exposure Time of Alloy 600 Specimen Prior to In-Situ Raman Spectra Measurements in High Temperature Water.

<table>
<thead>
<tr>
<th>DH2 (cc/kg)</th>
<th>Temperature (°C)</th>
<th>Total exposure time prior to first measurement at temperature (h)</th>
<th>Hold time at each temperature prior to first measurement (h)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>30</td>
<td>250</td>
<td>28</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>290</td>
<td>33.5</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>320</td>
<td>42</td>
<td>7</td>
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<tr>
<td></td>
<td>350-I</td>
<td>47.5</td>
<td>3.5</td>
</tr>
<tr>
<td></td>
<td>350-II</td>
<td>69.5</td>
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<td></td>
<td>350-I</td>
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<td>2</td>
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<tr>
<td></td>
<td>350-II</td>
<td>74</td>
<td>23</td>
</tr>
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Table 4. Summary of Experimental Observations and Thermochemical Predictions of Surface Oxide Phases of Alloy 600 in 350°C Water at Each Dissolved Hydrogen Concentration.

<table>
<thead>
<tr>
<th>DH2 = 30cc/kg</th>
<th>Predicted at 350°C</th>
<th>Observed at 350°C</th>
</tr>
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<tbody>
<tr>
<td>With CrOOH</td>
<td></td>
<td>NiO</td>
</tr>
<tr>
<td>NiFe2O4</td>
<td>NiO</td>
<td>NiCr2O4</td>
</tr>
<tr>
<td>NiCr2O4</td>
<td>NiO</td>
<td>NiCr2O4</td>
</tr>
<tr>
<td>FeCr2O4</td>
<td>FeCr2O4</td>
<td>NiO</td>
</tr>
<tr>
<td></td>
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<td>NiO</td>
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</table>

<table>
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<th>DH2 = 1cc/kg</th>
<th>Predicted at 350°C</th>
<th>Observed at 350°C</th>
</tr>
</thead>
<tbody>
<tr>
<td>With CrOOH</td>
<td></td>
<td>NiO</td>
</tr>
<tr>
<td>NiFe2O4</td>
<td>NiO</td>
<td>NiCr2O4</td>
</tr>
<tr>
<td>NiCr2O4</td>
<td>NiO</td>
<td>NiCr2O4</td>
</tr>
<tr>
<td>FeCr2O4</td>
<td>FeCr2O4</td>
<td>NiO</td>
</tr>
<tr>
<td></td>
<td></td>
<td>NiO</td>
</tr>
</tbody>
</table>

Table 5. Summary of Experimental Observations of Surface Oxide Phases Of Alloy 600 in 350°C Water in Comparison with Literature Data at Each Dissolved Hydrogen Concentration.

<table>
<thead>
<tr>
<th>Method</th>
<th>This work</th>
<th>Soustelle11</th>
<th>Caron12</th>
<th>Terachi30</th>
</tr>
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<tbody>
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<td>Temperature</td>
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<td>GDOS + EDS</td>
<td>XPS</td>
<td>Synchrotron XRD + TEM</td>
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<tr>
<td>(°C)</td>
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<td>360</td>
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<td>Exposure</td>
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<td>1869</td>
<td>1000</td>
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<tr>
<td>time (hrs)</td>
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<tr>
<td>Oxide(s) at</td>
<td>CrOOH, Cr-Oxide</td>
<td>Compact layer (NiCr2O4) + Precipitates (NiFe2O4)</td>
<td>NiCr2O4</td>
<td>(Ni+Cr-rich) oxide + Precipitates (NiFe2O4)</td>
</tr>
<tr>
<td>high DH2</td>
<td>NiO + NiCr2O4</td>
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<tr>
<td>Oxide(s) at</td>
<td>CrOOH, Cr-Oxide</td>
<td>Compact layer (NiCr2O4)</td>
<td>NiO</td>
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<td>low DH2</td>
<td>NiO</td>
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Fig. 1 Influence of hydrogen concentration on the crack growth rate of PWSCC in alloy 600 at 360°C.

Fig. 2 Layout of in-situ Raman spectroscopic measurement system in high temperature aqueous environment.

Fig. 3 Field emission scanning electron micrographs of oxide film on alloy 600 after 110 hours exposure in simulated PWR water.
Fig. 4. TEM micrographs of oxide film on alloy 600 after 71 hours exposure in simulated PWR water.

Fig. 5. In-situ Raman spectra obtained for alloy 600 by holding at 250, 290, 320 and 350°C, respectively, during heating up to 350 °C in PWR water condition with DH₂=30cc(STP)/kg.

Fig. 6. In-situ Raman spectra obtained for alloy 600 by holding at 250, 290, 320 and 350°C, respectively, during heating up to 350°C in PWR water condition with DH₂=1cc(STP)/kg.

Fig. 7. In-situ Raman spectra obtained for alloy 600 at 350°C in PWR water as the DH₂ in water was increased from 1 to 30 cc/kg.

Fig. 8. Summary plot on electrochemical potential for nickel in the alloy 600 as a function of temperature.