

# Irradiation-Assisted Stress Corrosion Cracking Behavior of Austenitic Stainless Steels Applicable to LWR Core Internals

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# Irradiation-Assisted Stress Corrosion Cracking Behavior of Austenitic Stainless Steels Applicable to LWR Core Internals

Manuscript Completed: March 2005 Date Published: January 2006

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Prepared for Division of Engineering Technology Office of Nuclear Regulatory Research U.S. Nuclear Regulatory Commission Washington, DC 20555-0001 NRC Job Code Y6388



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### Irradiation-Assisted Stress Corrosion Cracking Behavior of Austenitic Stainless Steels Applicable to LWR Core Internals

by

H. M. Chung and W. J. Shack

#### Abstract

This report summarizes work performed at Argonne National Laboratory on irradiation-assisted stress corrosion cracking (IASCC) of austenitic stainless steels that were irradiated in the Halden reactor in simulation of irradiation-induced degradation of boiling water reactor (BWR) core internal components. Slow-strain-rate tensile tests in BWR-like oxidizing water were conducted on 27 austenitic stainless steel alloys that were irradiated at 288°C in helium to 0.4, 1.3, and 3.0 dpa. Fractographic analysis was conducted to determine the fracture surface morphology. Microchemical analysis by Auger electron spectroscopy was performed on BWR neutron absorber tubes to characterize grain-boundary segregation of important elements under BWR conditions. At 0.4 and 1.4 dpa, transgranular fracture was mixed with intergranular fracture. At 3 dr.a, transgranular cracking was negligible, and fracture surface was either dominantly intergranular, as in field-cracked core internals, or dominantly ductile or mixed. This behavior indicates that percent integranular stress corrosion cracking determined at  $\approx 3$  dpa is a good measure of IASCC susceptibility. At  $\approx$ 1.4 dpa, a beneficial effect of a high concentration of Si (0.8-1.5 wt.%) was observed. At  $\approx$ 3 dpa, however, such effect was obscured by a deleterious effect of S. Excellent resistance to IASCC was observed up to  $\approx 3$ dpa for eight heats of Types 304, 316, and 348 steel that contain very low concentrations of S. Susceptibility of Types 304 and 316 steels that contain >0.003 wt.% S increased drastically. This indicates that a sulfurrelated critical phenomenon plays an important role in IASCC. A sulfur content of <0.002 wt.% is the primary material factor necessary to ensure good resistance to IASCC. However, for Types 304L and 316L stee! and their high-purity counterparts, a sulfur content of < 0.002 wt.% alone is not a sufficient condition to ensure good resistance to IASCC. This is in distinct contrast to the behavior of their high-C counterparts. At S concentrations >0.002 wt.%, the deleterious effect of S is so dominant that a high concentration of C is not an important factor. A two-dimensional map was developed in which susceptibility or resistance to IASCC is shown as a function of bulk concentrations of S and C. Data reported in the literature are consistent with the map. The map is helpful to predict relative IASCC susceptibility of Types 304 and 316 steels. A similar but somewhat different map is helpful to predict IASCC behavior of Type 348 steels. Grain-boundary segregation of S was observed for BWR neutron absorber tubes irradiated to  $\approx 3$  dpa. On the basis of the results of the stress-corrosion-cracking tests and the microstructural characterization, a mechanistic IASCC model has been developed.

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### Foreword

This report examines the effects of simulated light-water reactor coolants, material chemistry, and irradiation damage on the susceptibility to stress corrosion cracking of a series of commercially available and laboratory-melted stainless steels. This report is one of a series dating back about 8 years, describing such results, which are required to support analysis of the structural integrity of reactor internal components, many of which are subject to irradiation-assisted stress corrosion cracking (IASCC). Earlier publications in this series include NUREG/CR-5608, "Irradiation-Assisted Stress Corrosion Cracking of Model Austenitic Stainless Steels Irradiated in the Halden Reactor"; NUREG/CR-6826, "Fracture Toughness and Crack Growth Rates of Irradiated Austenitic Stainless Steels"; and NUREG/CR-6687, "Irradiation-Assisted Stress Corrosion Cracking of Model Austenitic Stainless Steel Stress Corrosion Cracking of Model Austenitic Stainless Steels Stress Corrosion Cracking of Model Austenitic Stainless Steels Irradiated Austenitic Stainless Steels"; and Austenitic Stainless Steels Stress Corrosion Cracking of Model Austenitic Stainless Steel Alloys."

This report and its sister report, which details crack growth rates in heat-affected zones adjacent to stainless steel weldments, comprise the final publications based on specimens irradiated in Phase I (of twc) in the Halden test reactor. Phase I irradiations principally comprised stainless steels of wide-ranging chemistry (including commercial steels of typical chemistry) and conventional heat treatment and product form processing, while Phase II irradiations included a plurality of innovatively fabricated and engineered alloys designed to possibly be more resistant to IASCC. Irradiation levels in both phases ranged up to about 3 dpa, and the high-temperature water environment employed in these tests contained 200 ppb to 8 ppm dissolved oxygen. The materials tested included several commonly used stainless steels, such as Types 304, 316 (and their low-carbon counterparts), and 347, and several laboratory heats that were expected to have an impact on IASCC properties. Taken together, these materials and test conditions make these results most applicable to boiling-water reactor internals. Publications concerning test results from Phase II irradiations are expected in the 2006 – 2008 time frame.

As in earlier reports, the deleterious effect of sulfur, which increases susceptibility to IASCC, was confirmed and further quantified. Reducing sulfur content to less than about 0.002 wt. % provides protection against IASCC. By contrast, susceptibility increased dramatically with heats that contain >0.003 wt. % S. A sufficiently low concentration of sulfur is the primary material factor necessary to ensure good resistance to IASCC. However, a sulfur concentration of less than 0.002 wt. % alone is not a sufficient material condition to ensure resistance to IASCC. Rather, a moderately high (>0.03 wt. %) level of carbon is also required in Type 304 and 316 steels. This is an important observation, given that many stainless steels (Type 304L or 316L) are decarburized to protect against thermal sensitization; too low a carbon level could render such steels susceptible to IASCC. Based on these results and related microstructural characteristics and data from electron microscopy studies, the authors have developed a model of grain boundary processes leading to IASCC. Such mechanistic understanding may shed light on the principles that might become the basis for a calculational model for use in predicting IASCC susceptibility.

Carl J. Paperiell, Director Office of Nuclear Regulatory Research U.S. Nuclear Regulatory Commission

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### Acronyms

- AAEM = advanced analytical electron microscopy
- AES = Auger electron spectroscopy
- AISI = American Iron and Steel Institute

ASTM = American Society for Testing and Materials

- BWR = boiling water reactor
- CGR = crack growth rate

CP == commercial purity

- CT == compact tension
- dpa == displacement per atom
- DO == dissolved oxygen
- ECP = electrochemical potential
- EDS = energy dispersive spectroscopy
- FEG-AAEM = field-emission-gun advanced analytical electron microscopy
- HBV/R = heavy boiling water reactor
- HP ≕ high purity
- IASCC = irradiation-induced stress corrosion cracking

IG = intergranular

- IGC = intergranular cracking
- IGSCC = intergranular stress corrosion cracking
- JRR == Japan Research Reactor
- LWR = light water reactor
- PWR = pressurized water reactor
- SAM = scanning Auger electron microscope
- SEM = scanning electron microscopy
- SS = stainless steel

SSRT = slow-strain-rate tensile

TE = total elongation

TEM = transmission electron microscopy

TG = transgranular

TGSCC = transgranular stress corrosion cracking

UE = uniform elongation

UHP = ultra high purity

UTS = ultimate tensile strength

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YS = yield strength

### **Executive Summary**

This report summarizes experimental work performed at Argonne National Laboratory on irradiationassisted stress corrosion cracking (IASCC) of austenitic stainless steels that were irradiated in the Halden boiling heavy water reactor in order to simulate irradiation-induced degradation of boiling water reactor (BWR) core internal components. Slow-strain-rate tensile tests in BWR-like water were conducted or. 27 austenitic stainless steel alloys that were irradiated at 288°C in helium to three fluence levels, i.e.,  $\approx 0.3 \times 10^{21}$ n·cm<sup>-2</sup>,  $\approx 0.9 \times 10^{21}$  n·cm<sup>-2</sup>, and  $\approx 2.0 \times 10^{21}$  n·cm<sup>-2</sup> (E > 1 MeV) (neutron damage  $\approx 0.4$ ,  $\approx 1.3$ , and  $\approx 3.0$ displacement per atom [dpa]). Fractographic analysis by scanning electron microscopy was conducted to determine the susceptibility to IASCC as manifested by the degree of intergranular fracture surface morphology. Microchemical analysis by Auger electron spectroscopy was performed on BWR neutron absorber tubes to characterize the behavior of grain-boundary segregation of important elements under BWR operating conditions. On the basis of the results from the stress-corrosion-cracking tests and microstructural analyses, a mechanistic model for IASCC was developed.

At  $\approx 0.3 \times 10^{21} \text{ n} \cdot \text{cm}^{-2}$  (E > 1 MeV)( $\approx 0.4 \text{ dpa}$ ), IASCC susceptibility was negligible for all tested heats, with the exception of a 316L-like high-purity model alloy that contained 0.003 wt.% C and 0.004 wt.% S. At  $\approx 0.4$  and  $\approx 1.3$  dpa, significant transgranular fracture was mixed with intergranular fracture. At  $\approx 3$  cpa, susceptibility to transgranular stress corrosion cracking was negligible, and fracture surface morphology was either dominantly intergranular, as in field-cracked core internal components, or dominantly ductile or mixed. This behavior indicates that percent intergranular stress corrosion cracking determined at  $\approx 3$  dpa is a good measure of IASCC susceptibility.

At  $\approx 1.4$  dpa, a beneficial effect of a high concentration of Si (0.8-1.5 wt.%) was observed. At  $\approx 3$  dpa, however, such effect was obscured by a strongly deleterious effect of S. From the standpoint of BWR IASCC, which usually occurs at >2.5 dpa, typical variation in Si concentration in the range of 0.4-1.1 wt.% in Types 304, 316, and 348 steel is, therefore, not an important factor.

Excellent resistance to IASCC was observed up to  $\approx 3$  dpa for seven heats of Types 304 and 316 steel that contain very low S concentrations of  $\leq 0.002$  wt.% and for one heat of Type 348 steel that contains 0.005 wt.% S. Susceptibility of Types 304 and 316 steel that contain >0.003 wt.% S increased drastically. These observations indicate that a sulfur-related critical phenomenon plays an important role in IASCC. A sufficiently low concentration of S (i.e., <0.002 wt.%) is the primary material factor necessary to ensure good resistance to IASCC. For Types 304L and 316L steels and their high-purity counterparts that contain <0.025 wt.% C, a sulfur concentration of <0.002 wt.% alone is not a sufficient material condition to ensure good resistance to IASCC. This is in distinct contrast to the behavior of the high-C counterparts (i.e., Types 304 and 316 steel that contain <0.002 wt.% S and >0.03 wt.% C), which exhibit good resistance to IASCC. The contrasting behavior indicates that, at sufficiently low concentrations of S, a high concentration of C is beneficial. However, at S concentrations >0.003 wt.%, the deleterious effect of S is so dominant that a high concentration of C is not an important factor.

A two-dimensional map was developed in which susceptibility or resistance to IASCC is shown as a function of bulk concentrations of S and C. Data reported in the literature are consistent with the map. The two-dimensional map is helpful to predict relative IASCC susceptibility of Types 304 and 316 steels. A similar map is helpful to predict IASCC behavior of Type 348 steels.

At S concentrations >0.003 wt.%, IASCC susceptibility is thought to be influenced significantly by various material-related factors, such as the degree of S segregation to grain boundaries, annealing and cooling during fabrication, component thickness, presence of a small amount of delta ferrite, Mn content, precipitation of sulfides, solubility of sulfides in the crack-tip water, and the thickness and morphology of the Cr oxide on the crack-tip grain boundary, which controls oxygen permeation and diffusion along the grain boundary. A model alloy that contains  $\approx 3$  vol.% globular-shaped delta ferrite and  $\approx 21$  wt.% Cr exhibited excellent resistance to IASCC, despite the fact that it contained a very high S concentration of 0.028 wt.%. The excellent resistance is attributed to the effect of delta ferrite, which traps S atoms, and the effect of a high concentration of Cr, which forms Cr oxide and helps to block oxygen permeation and diffusion along the austenite grain boundary.

Grain-boundary segregation of S was observed for BWR neutron absorber tubes irradiated to  $\approx 3$  dpa. It is not clear, however, whether the S segregation occurred via a thermal process, an irradiation-induced process, or both. The S segregation behavior is consistent with the results of stress-corrosion-cracking tests that showed a predominantly deleterious effect of S on IASCC at high fluence. Based on the results of stresscorrosion-cracking tests and microstructural characterization, an IASCC model has been developed. The model is based on several key steps postulated to occur at the crack-tip grain boundary, i.e., grain-boundary segregation of Ni and S, preferential oxidation of Cr and Fe atoms over Ni atoms by oxygen that diffuses along the grain boundary, formation of a Cr-Fe spinel oxide ahead of the crack tip, formation of Ni- and Srich thin films and islands at the metal/spinel boundary, and melting or amorphization of the Ni-and Srich thin films and islands. The model also predicts that the thickness and morphology of Cr oxide that covers the crack-tip grain boundary strongly influence the rate of oxygen permeation and diffusion along the grain boundary.

### Acknowledgments

The authors thank T. M. Karlsen, OECD Halden Reactor Project, Halden, Norway for many years of specimen irradiations in the Halden reactor; H. Valseth and J. R. McDade for transportation of the irradiated specimens; D. O. Pushis and D. McGann for specimen retrieval; W. E. Ruther, D. R. Perkins, R. W. Clark, and L. A. Knoblich for conducting slow-strain-rate-tensile tests; R. V. Strain, D. L. Perry, A. G. Hins, and J. Y. Park for fractographic examination by scanning electron microscopy; and J. E. Sanecki for grain-boundary analysis by Auger electron spectroscopy. The authors are grateful to C. Moyer and A. Lee for many helpful discussions. The authors would like to express special thanks to W. H. Cullen, Jr. of U.S. Nuclear Regulatory Commission who managed this project.

### 1 Introduction

In recent years, failures of reactor core internal components have increased after accumulating a fluence  $>0.5 \times 10^{21} \text{ n} \cdot \text{cm}^{-2}$  (E >1 MeV), or  $\approx 0.7$  dpa, in boiling water reactors (BWRs),<sup>1</sup> and at approximately one order of magnitude higher fluences in some pressurized water reactors (PWRs).<sup>2</sup> The general pattern of the observed failures indicates that, as nuclear plants age and neutron fluence increases, various nonsensitized austenitic stainless steels (SSs) become susceptible to intergranular (IG) failure. Some components are known to have cracked under minimal applied stress. Although most failed components can be replaced, some safety–significant structural components (e.g., the BWR top guide, shroud, and core plate) would be very costly or impractical to replace. Therefore, the structural integrity of these components at high fluence or near the end of life (EOL) has been a subject of concern, and extensive research has been conducted to provide an understanding of this type of degradation, commonly known as irradiation-assisted stress corrosion cracking (IASCC).

Irradiation affects profoundly both local coolant-water chemistry and component microstructure. The primary effects of irradiation on materials include alteration of local microchemistry, microstructure, and mechanical properties of the core internal components, which are usually fabricated from American Iron and Steel Institute (AISI) Types 304, 316, or 348 SS. Irradiation produces defects and defect clusters in grain matrices, alters the dislocation and dislocation loop structures, and produces defect-impurity and defect-cluster-impurity complexes, leading to radiation-induced hardening and, in many cases, flow localization via dislocation channeling. Irradiation also leads to changes in the stability of second–phase precipitates and the local alloy chemistry near grain boundaries, precipitates, and defect clusters. Grain–boundary microchemistry that significantly differs from bulk composition can be produced in association with not only radiation–induced segregation but also thermally driven segregation of alloying and impurity elements.<sup>1-3</sup>

For many years, irradiation-induced grain-boundary depletion of Cr has been considered the primary metallurgical process that causes IASCC. One of the most important factors that has been considered by many investigators to be a support for the Cr-depletion mechanism is the similarity in dependence on water chemistry (i.e., oxidizing potential) of both intergranular stress corrosion cracking (IGSCC) of unirradiated, thermally sensitized material and IASCC of solution-annealed and quenched, BWR-irradiated material.<sup>4,5,6</sup> However, contrary to expectations based on the Cr-depletion mechanism, cracking incidents of control rod cladding and baffle bolts have been reported in numerous PWRs (i.e., under nonoxidizing potential). Also, the susceptibility of PWR-irradiated components to IASCC has been shown clearly by expanding-mandrel,<sup>2,7</sup> slow-strain-rate tensile (SSRT) tests,<sup>8,9,10</sup> and C-ring compression tests<sup>11</sup> in PWR water<sup>2,7</sup> or simulated PWR-like water,<sup>8-11</sup> although PWR water chemistry falls well within the range of protective electrochemical potential (ECP). Also, it has been reported that crack propagation by intergranular separation occurs in highly irradiated compact-tension specimens even in hydrogen-water chemistry (HWC) in a simulated BWR environment.<sup>12</sup> These observations indicate that the phenomenon of IASCC cannot be explained well by the grain-boundary Cr depletion and slip dissolution mechanism.

Another important aspect of IASCC characteristics that does not support the grain-boundary Crdepletion mechanism is the effect of postirradiation annealing. Several investigators have reported that postirradiation annealing at 450-550°C for relatively short periods of time is sufficient to partially or fully mitigate the susceptibility to IASCC of austenitic SSs irradiated under BWR<sup>13-15</sup> and PWR<sup>10,16</sup> conditions. During short annealing at such low temperatures (e.g., 5-24 h at 475°C<sup>14</sup> and 1 h at 550°C<sup>16</sup>), it is not possible for significant back diffusion of metallic elements (Fe, Cr, and Ni) to occur from the grain boundary. It has been shown that, during postirradiation annealing, IASCC susceptibility becomes negligible long before grain-boundary Cr concentration begins to change from its level in as-irradiated state.<sup>16,17</sup> These observations do not support the premise that grain-boundary Cr depletion is the direct cause of IASCC. The observations, however, suggest a possibility that a rapidly diffusing impurity element, segregated on the grain boundary during irradiation, plays an important role in annealing-induced restoration of the resistance to IASCC.

In general, IASCC is characterized by strong heat-to-heat variation in susceptibility, in addition to strong dependence on irradiation condition, material type, and grade, even among materials of virtually identical chemical composition. This behavior also indicates that the traditional interpretation based on the role of grain-boundary Cr depletion cannot explain well the mechanism of IASCC, because, for a given level of neutron fluence, heat-to-heat variation of Cr depletion is insignificant among a given type of solution-annealed and quenched steel.

Many investigators have implicated radiation-induced segregation of Si or P as the primary process that causes IASCC.<sup>1-3,7</sup> The superior resistance of one heat of Type 348 SS that is especially low in C, Si, and P seemed to provide support for this process,<sup>2</sup> and the same rationale was extended to Type 304 SS. However, in direct contradiction, several investigators reported that IASCC resistance of high-purity (HP) heats (low in C, Si, and P) of Type 304 SS is no better than that of commercial-purity (CP) counterparts.<sup>7-9,12,16,23</sup> Therefore, the role of grain-boundary segregation of C, Si, and P appeared poorly established.

Although C significantly increases the yield strength of irradiated stainless steels, higher C content (>0.06 wt.%) seems to be either benign or even conducive to lower susceptibility of irradiated austenitic SSs to IASCC.<sup>23,24</sup>

Deleterious effects of O on IGSCC of proton-irradiated steels have been reported by Cookson et al., who showed that oxides near the surface facilitate crack nucleation.<sup>25</sup> High concentrations of O and internal oxide precipitates were also reported for grain boundaries in the heat-affected zones of unirradiated Types 304 and 304L SSs and field-cracked BWR core shroud welds that were fabricated by the shielded-metal-arc procedure.<sup>26,27</sup> Yet, the exact role and the source of O on grain boundaries in IASCC of a solution-annealed material or intergranular failure of a BWR core internal weld remain poorly understood.

Indications of the deleterious effect of grain-boundary segregation of N have been reported for BWR neutron absorber tubes fabricated from high- and commercial-purity heats of Type 304 SS.<sup>23</sup> Similar reports suggested that a high concentration of N (e.g., >0.06 wt.%) is deleterious, at least under BWR conditions.<sup>1,28-30</sup> Indications of the deleterious role of N have also been reported for Types 304L and 316L SS that contain <0.024 wt.% C and have been irradiated in BWRs or test reactors at 240–300°C.<sup>22</sup> Kasahara et al.<sup>28</sup> also reported that higher N in Type 316L increased the susceptibility to IASCC, indicating that Type 316LN is a susceptible material. This observation appeared to be consistent with the behavior of 316NG SSs reported by Jacobs et al.<sup>1</sup> and Jenssen and Ljungberg.<sup>30</sup> In contrast to these observations, 316NG SSs irradiated at ≈50°C have been reported to be resistant to intergranular failure at ≈288°C in high-purity water that contained ≈32 ppm dissolved oxygen (DO).<sup>22</sup> Therefore, the role of N appears to be neither clear nor convincing.

The strong heat-to-heat variation in IASCC susceptibility has been customarily correlated with variations in irradiation-induced evolution of the microstructure and microchemistry of ordinary grain boundaries, i.e., grain boundaries not directly involved in crack-tip formation or crack propagation. This approach implicitly assumes that the crack tip, stress intensity, or the event of crack propagation itself does

not influence the microstructure and microchemistry of the grain boundary in front of the advancing crack tip. However, contrary to this assumption, results from several crack-tip microstructural studies indicate that the microstructure and microchemistry of a crack-tip grain boundary differ significantly from those of an ordinary grain boundary.<sup>31-34</sup>

Although intergranular failure in irradiated fuel cladding fabricated from austenitic SSs has been observed since the 1960s, extensive research and investigations on core internals were not conducted until the late 1980s. In spite of the extensive investigations, IASCC is still only poorly understood, and its mechanism remains elusive, and in some respects, more perplexing. In the context of this somewhat unusual background, we have initiated a comprehensive program to systematically investigate the effects of alloying and impurity elements (Cr, Ni, Si, P, S, Mn, C, O, and N) on the susceptibility of austenitic SSs to IASCC at several fluence levels. Construction of the test matrix was based on the method of Taguchi,<sup>35</sup> and susceptibility to IASCC was March 17, 2005 determined by well-controlled SSRT testing of the irradiated specimens in BWR-like water. Posttest fractographic examination was performed by scanning electron microscopy (SEM) and Auger electron spectroscopy (AES).

This report is the final topical report of the program; it summarizes the results obtained from 27 austenitic SS alloys that were irradiated in the Halden reactor at 288°C in helium to three fluence levels, i.e.,  $\approx 0.3 \times 10^{21} \text{ n} \cdot \text{cm}^{-2}$  ( $\approx 1.3 \text{ dpa}$ ), and  $\approx 2.0 \times 10^{21} \text{ n} \cdot \text{cm}^{-2}$  ( $\approx 3.0 \text{ dpa}$ ). Several reports have been issued previously to document the results obtained from specimens irradiated to the low,<sup>36,37</sup> medium,<sup>38,39</sup> and high<sup>40</sup> fluence levels.

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#### 2.1 Construction of Test Matrix

The test matrix for irradiation and stress corrosion testing for this project was constructed according to the method of Taguchi, which is described in detail in Ref. 39. The base matrix followed Taguchi's standard orthogonal array  $L_{18}$  (2<sup>1</sup> x 3<sup>7</sup>), which is an optimized matrix designed to systematically determine the effects of seven variables (i.e., bulk material concentrations of Cr, Si, P, S, Mn, C, and N) at three concentration levels, and one variable (bulk Ni concentration) at two levels. A possible synergistic interaction was assumed only between Ni and Si. Compositions assigned to the three levels of the seven alloying and impurity elements and two levels of Ni are given in Table 1. The basic test matrix, optimized on the basis of the orthogonal array  $L_{18}$  (2<sup>1</sup> x 3<sup>7</sup>), is given in Table 2.

		COMPOSITION (IN WT.%)					
Factor	Element	Ass	ASSIGNED TO LEVEL				
Ň0.	ASSIGNED	· 1	1 2				
1	NI	8.000	10.000	-			
2	Si	0.700	1.400	0.150			
<b>3</b> ·	Р	0.040	0.090	0.020			
4	S	0.006	0.020	0.002			
5	Mn	1.200	2.050	0.500			
6	С	0.040	0.090	0.010			
7	Ν	0.040	0.090	0.010			
8	Cr	17.500	18.500	16.000			

Table 1. Compositions assigned to medium, high, and low levels of alloying and impurity elements used to construct Taguchi's  $L_{18}$  (2<sup>1</sup> x 3<sup>7</sup>) array.

In addition to the 18 statistically optimized heats given in Table 2, six supplementary heats of commercial- and high-purity grade Types 304, 316, and 348 SS were included in the test matrix. Compositions of the six supplementary heats are given in Table 3. Of the 24 heats given in Tables 2 and 3, eight (i.e., Heats 1, 3, 9, 10, 12, 16, 19, and 21) were replaced by commercially purchased heats. Compositions of major impurities (i.e., Si, P, C, and N) of each of the eight commercial heats matched closely those of each corresponding heat given in Tables 2 and 3. The prefix "C" was added to the identification number of these eight commercial heats (see Table 4).

The remaining 16 heats in Tables 2 and 3 were fabricated in the laboratory (USDOE Materials Preparation Center, Ames Laboratory, Ames, Iowa). All of the laboratory-fabricated heats were designated with identification numbers that began with "L". To this matrix of 24 heats, three laboratory heats were added to test the effects of the fabrication procedure. Compositions of these three laboratory heats (i.e., Heats L25, L26, and L27 in Table 4) closely match those of the counterpart commercial heats (i.e., Heats C3, C19,

and C21), respectively. Elemental composition of the complete test matrix, comprising 27 austenitic SSs, is given in Table 4.

Oxygen content, which is not specified in AISI Specifications for Types 304, 316, and 348 SS and their low-C counterparts, was not systematically controlled. However, oxygen content in the 27 heats ranged from 0.0037 wt.% (L4) to 0.0274 wt.% (L7) except for one heat: Heat L20 contained an unusually high concentration of O (0.0940 wt.%) and exhibited an anomalous dendritic structure that is not characteristic of an austenitic SS.

HEAT	COMPOSITION (IN WT.%)							
No.	NI	Si	Р	S	Mn	С	N	CR
1	10.0	0.70	0.04	0.006	1.20	0.04	0.04	18.5
2	10.0	0.70	0.09	0.020	2.05	0.09	0.09	17.5
3	10.0	0.70	0.02	0.002	0.50	0.01	0.01	16.0
4	10.0	1.40	0.04	0.006	2.05	0.09	0.01	16.0
5	10.0	1.40	0.09	0.020	0.50	0.01	0.04	18.5
6	10.0	1.40	0.02	0.002	1.20	0.04	0.09	17.5
7	10.0	0.15	0.04	0.020	1.20	0.01	0.09	16.0
8	10.0	0.15	0.09	0.006	2.05	0.04	0.01	18.5
9	10.0	0.15	0.02	0.002	0.50	0.09	0.04	17.5
10	.8.0	0.70	0.04	0.002	0.50	0.09	0.09	18.5
11	8.0	0.70	0.09	0.006	1.20	0.01	0.01	17.5
12	8.0	0.70	0.02	0.020	2.05	0.04	0.04	16.0
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13	8.0	1.40	0.04	0.020	0.50	0.04	0.01	17.5
14	8.0	1.40	0.09	0.002	2.05	0.09	0.04	16.0
15	8.0	1.40	0.02	0.006	1.20	0.01	0.09	18.5
16	8.0	0.15	0.04	0.002	2.05	0.01	0.04	17.5
17	8.0	0.15	0.09	0.006	0.50	0.04	0.09	16.0
18	8.0	0.15	0.02	0.020	1.20	0.09	0.01	18.5

Table 2. Basic test matrix optimized according to Taguchi's orthogonal array  $L_{18}$  (2<sup>1 x</sup> 3<sup>7</sup>).

HEAT	SS TYPE	C	N	SI	Р	S	Mn	CR	NI	NB	Мо
19	CP 304	0.04	0.04	0.70	0.04	0.06	1.20	18.5	10.0	-	-
20	HP 304	0.01	0.01	0.15	0.02	0.02	0.50	18.5	10.0	-	-
21	CP 316	0.04	0.04	0.70	0.04	0.06	1.20	16.5	13.5	-	2.20
22	HP 316	0.01	0.01	0.15	0.02	0.02	0.50	16.5	13.5	-	2.20
23	CP 348	0.04	0.04	0.70	0.04	0.06	1.20	17.5	11.5	2.20	-
24	HP 348	0.01	0.01	0.15	0.02	0.02	0.50	17.5	11.5	2.20	-

Table 3. Composition (wt.%) of six supplementary commercial- and high-purity heats of Types 304, 316, and 348 stainless steel.

Table 4. Elemental composition of 27 commercial and laboratory model austenitic stainless steel alloys irradiated in Halden Reactor.

ANL	Source		Elemental Composition (wt.%)									
ID <sup>a</sup>	Heat ID	Ni	Si	Р	S	Mn	С	N	Cr	0	В	Mo, Nb
Ci	ASW-70378	8.12	0.50	0.038	0.0020	1.00	0.060	0.060	18.11	0.0102	<0.001	-
L2	BPC-4111	10.50	0.82	0.080	0.0345	1.58	0.074	0.102	17.02	0.0065	<0.001	-
C3	PNL-C1	8.91	0.46	0.019	0.0040	1.81	0.016	0.083	18.55	-	<0.001	-
N	BPC-488	10.20	0.94	0.031	0.0100	1.75	0.110	0.002	15.80	0.0037	<0.001	-
L5	BPC-4104	9.66	0.90	0.113	0.0283	0.47	0.006	0.033	21.00	0.0068	<0.001	-
Lo	BPC-4127	10.00	1.90	0.020	0.0051	1.13	0.096	0.087	17.10	0.0058	<0.001	-
L.,	BPC-4112	10.60	0.18	0.040	0.0384	1.02	0.007	0.111	15.40	0.0274	<0.001	-
LS	BPC-491	10.20	· 0.15	0.093	0.0100	1.85	0.041	0.001	18.30	0.0059	<0.001	-
C9	PNL-C6	8.75	0.39	0.013	0.0130	1.72	0.062	0.065	18.48	0.0102	<0.001	-
C10	ASW-23381	8.13	0,55	0.033	0.0020	1.00	0.060	0.086	18.19	0.0074	<0.001	-
L11	BPC-493	8.15	0.47	0.097	0.0094	1.02	0.014	0.004	17.40	0.0071	<0.001	-
C:2	ASW-23805	8.23	0.47	0.018	0.0020	1.00	0.060	0.070	18.43	-	<0.001	-
LIS	BPC-496	8.18	1.18	0.027	0.0222	0.36	0.026	0.001	17.40	0.0042	<0.001	-
L14	BPC-4129	7.93	1.49	0.080	0.0023	1.76	0.107	0.028	15.00	0.0045	< 0.001	-
L15	BPC-4126	8.00	1.82	0.010	0.0129	1.07	0.020	0.085	17.80	0.0110	< 0.001	-
C16	PNL-SS14	12.90	0.38	0.014	0.0020	1.66	0.020	0.011	16.92	0.0157	<0.001	Mo 2.30
L17	BPC-4128	8.00	0.66	0.090	0.0089	0.48	0.061	0.078	15.30	0.0090	<0.001	-
L18	BPC-498	8.13	0.14	0.016	0.0326	1.13	0.080	0.001	18.00	0.0055	<0.001	-
C19	ASW-74827	8.08	0.45	0.031	0.0030	0.99	0.060	0.070	18.21	0.0200	<0.001	-
L20	BPC-4101	8.91	0.017	0.010	0.0039	0.41	0.002	0.002	18.10	0.0940	<0.001	-
C21	4 SW-17455	10.24	0.51	0.034	0.0010	011	0.060	0.020	16.28	0.0112	<0.001	Mo 2.08
1 22	RPC-4100	13 30	0.024	0.034	0.0010	0.40	0.003	0.001	16 10	0.0103	<0.001	Mo 2.00
1 22	DIC-4100	12.04	0.024	0.015	0.0050	0.40	0.003	0.001	17 30	0.0103	~0.001	NB 1.06
1.2.5	DDC 4105	12.04	0.00	0.050	0.0475	0.30	0.071	0.072	16.00	0.0075	<0.001	ND 1.00
1.24	BPC-4103	12.30	0.03	0.007	0.0033	1.40	0.031	0.002	17.20	0.0129	<b>\0.001</b>	NO 1.72
L23	BrC-4133	6.93	0.92	0.020	0.0082	1.94	0.019	0.095	17.20	0.0085	0.010	-
L26	BPC-4131	8.09	0.79	0.004	0.0022	0.91	0.070	0.089	17.20	0.0080	<0.001	-
L27	BPC-4132	10.30	0.96	0.040	0.0018	0.97	0.057	0.019	15.30	0.0058	0.030	Mo 2.01

<sup>a</sup>The first letter "C" or "L" denotes, respectively, a commercial or a laboratory-fabricated heat.

### 2.2 Fabrication of Specimen

Slow-strain-rate tensile (SSRT) specimens were machined from solution-annealed and water-quenched plates, pipes, or sheets that were fabricated from the 27 model austenitic SS alloys. Thick plates or pipes of various thickness and diameter were obtained only for commercial heats, and the laboratory heats were in sheet form. The geometry of the SSRT specimens, 0.76 mm thick, 57.2 mm long, and 12.7 mm wide (gauge section 19.1 mm long and 3.1 mm wide), is shown in Fig. 1. Gauge length and plane of the specimen were parallel to the rolling direction and the plane of the sheet, respectively. Subsize compact-tension (1/4TCT) specimens were also fabricated and irradiated in tandem in the same capsules with the SSRT specimens to determine J-R fracture toughness properties and crack growth rate (CGR). The geometry of the CT specimen is shown in Fig. 2. Together, 96 SSRT and 24 CT specimens were prepared and irradiated in this study.

Before machining the specimens, the surface oxide layer was removed to minimize specimen-tospecimen variation in surface roughness and uncertainty in surface chemical composition. After the specimens were mechanically machined and polished, no additional heat treatment was applied to them. Electric discharge machining was not employed because of concern about potential contamination of the surface by Cu impurity. The machined specimens were degreased in acetone and cleaned ultrasonically in alcohol before they were encapsulated in clean Type 304 SS capsules. The capsules were filled with research-grade He for irradiation in the Halden HBWR located in Halden, Norway.



ALL DIMENSIONS IN MM

Figure 1. Geometry of slow-strain-rate tensile test specimen; all dimensions in mm.

The width of the SSRT specimens (12.7 mm, Fig. 1) was chosen similar to the width of the compacttension specimens (14.0 mm, Fig. 2) so that an optimum combination of the two types of specimens could be stacked in the irradiation capsule of the Halden reactor. In each capsule, four CT specimens were positioned at the bottom, and 16 SSRT specimens could be stacked vertically on top of the CT specimens. In this way, it was possible to make the best use of the irradiation space. For reference, the dimension of the compact tension specimens is given in Fig. 2, although the specimens were tested in a separate investigation.



Figure 2. Geometry of compact-tension specimen.

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## **3** Specimen Irradiation

Ninety-six SSRT and 24 CT specimens were encapsulated in six capsules, each capsule containing 16 SSRT and 4 CT specimens. A fixed 0.5-mm gap, filled with He, was allowed between the inner wall of the Type 304 SS capsule and the specimen surface. The size of the gap was selected to maintain specimen temperature at 289°C during irradiation in He. To prevent capsule wall creepdown and possible changes in gap size, spacers in the form of Type 304 SS wires (0.5-mm diameter) were placed between the specimens and the capsule inner wall. Type 304 SS filler bodies were inserted on both sides of the SSRT specimen stack to avoid overheating the thin gauge section. Table 5 shows the specimens categorized according to alloy, fluence level, and postirradiation test type. The six capsules were irradiated in the Halden HBWR starting on April 8, 1992. Fast neutron (E > 1 MeV) flux during the various irradiation cycles ranged from 1.80 x 10<sup>13</sup> n·cm<sup>-2</sup> s<sup>-1</sup> to 3.31 x 10<sup>13</sup> n·cm<sup>-2</sup> s<sup>-1</sup>. The irradiation history of the six capsules is summarized in Table 6. Specimen temperature during the irradiation, determined on the basis of the melting behavior of five to six types of low-melting alloys, was 288±2°C.

Alloy ID	SSRT Test			Cor	stant Load Te	st	J-R, Crack Growth Rate Test		
-	high <sup>a</sup>	medium <sup>a</sup>	low <sup>a</sup>	high	medium	low	high	medium	low
C:1	<u> </u>	1	1						
L2	1	1					1	1	
C'3	1	1	1				1	1	1
L4	1	1	1						
L5	1	1	1				1		
L6	1	1							
L7	1	1							
L8	1	1	1						
C9	1	1	1						
C10	1	· 1	1						
L11	1	1	1						
C12	1	1	1						
L13	1	1	1						
L14	1	1					1		
L15	1	1							
C16	1	1	1				1	1	
L17	1	1							
L18	1	1	1				1	1	
C19	5	1	1	4	4		1	1	1
L20	5	1	1	4	4		1	1	1
C21	1	1	1				1	1	1
L22	1	1	1				1	1	
L23	1	1					I		
L24	1	1					1		
L25	3								
L26	3								
L27	2								

Table 5.	Specimen	number per	<sup>,</sup> allov.	irradiation	fluence level	. and	postirradiation 1	lest type.
						,		

<sup>a</sup>Target fluence level in units of 10<sup>21</sup> n·cm<sup>-2</sup>, 0.3 (low), 0.9 (medium), and 2.5 (high).

Capsule ID	Fluence Level	Target Fluence <sup>a</sup>	Irradiation Cycles	Discharge Date	Discharge Fluence <sup>a</sup>
1	Medium	1.0	IFA 530-3 to -6, D07-004-2	Nov. 1996	0.9
4	Low	0.4	IFA 530-3	Oct. 1992	0.3
5	High	2.5	IFA 530-4 to -6, D07-004-1 to -3	Nov. 1999	2.0
6	High	2.5	IFA 530-4 to -6, D07-004-1 to -3	Nov. 1999	2.0
7	Medium	1.0	IFA 530-4 and -6, D07-004-1	May 1996	0.9
8	high	2.5	IFA 530-5 to -6, D07-004-1 to -3	Nov. 1999	2.0

Table 6.Summary of discharge date and fluence of model austenitic stainless steel alloys<br/>irradiated in Halden Heavy Boiling Water Reactor.

<sup>a</sup>In units of  $10^{21}$  n·cm<sup>-2</sup> (E >1 MeV).
# **4** Postirradiation Testing and Posttest Characterization

After retrieval of the irradiated specimens from the capsules, slow-strain-rate tensile tests and fractographic characterization by scanning electron microscopy were performed for each specimen listed in Table 5. In addition to the irradiated specimens, unirradiated control specimens were also tested under the same conditions.

All SSRT tests were conducted at 289°C in simulated BWR-like water that contained  $\approx 8$  ppm dissolved oxygen (DO). Conductivity and pH of the water at 23°C were kept at  $\approx 0.07-0.10 \ \mu S \ cm^{-1}$  and  $\approx 6.5$ , respectively. Strain rate was held constant at 1.65 x 10<sup>-7</sup> s<sup>-1</sup>. Electrochemical potential (ECP) was measured at the effluent side at regular intervals.

A limited number of specimens were selected and tested in air either at 23°C or 289-325°C in asirradiated state by the SSRT technique at a strain rate of  $1.65 \times 10^{-7} \text{ s}^{-1}$ . The objective of such testing was to determine the susceptibility of the selected heats to irradiation-assisted intergranular cracking in an inert environment at high fluence, and to compare the result with the susceptibility of the same heats to IASCC in 289°C water.

Some high-fluence specimens, irradiated to  $2.0 \times 10^{21}$  n cm<sup>-2</sup> ( $\approx 3$  dpa) and SSRT-tested in BWR-like oxidizing water at 289°C, were selected for rapid bending tests in 23°C air. Fracture in such a test was produced by simple bending of the gauge section near the specimen fracture tip that was produced during SSRT testing in 289°C water. Surprisingly, many high-fluence specimens were susceptible to intergranular cracking in 23°C air after 1-3 times of repeated bending to  $\approx 90^{\circ}$ .

After SSRT testing in water or in air, fracture surface morphology was imaged in a shield scanning electron microscope (SEM). From the medium-magnification (100-150X) images, a composite fracture surface was constructed for each tested specimen. The type of fracture surface morphology (i.e., ductile, intergranular, and transgranular) was identified from the high-magnification fractographs, which varied from region to region on the fracture surface. From the composite of the whole fracture surface, fractions of intergranular (IG) and transgranular (TG) fracture morphology were measured.

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# 5 SSRT SCC Test and Fractography on Unirradiated Specimens

Yield strength and total elongation, measured on the non-irradiated control specimens in the 8-ppm-DO water at 288°C, are plotted in Figs. 3 and 4, respectively. Yield strengths of the non-irradiated solutionannealed Type 304 SS heats were 170-240 MPa, whereas yield strengths of Type 316 and 348 SS heats were significantly higher, i.e., 280-300 MPa and 330-350 MPa, respectively. Total elongation measured in water on most non-irradiated specimens was 17-59% (see Fig. 4). However, heat-to-heat variation in ductility in BWR-like water was significant, even among the same type of steel. Of the 17 heats tested in 289°C water, five heats exhibited relatively low ductility. Compositional characteristics of these five heats are denoted in Fig. 4. Two Type 304 SS heats that contained unusually high concentrations of S exhibited unusually low ductility (5-8%), i.e., Heat L7 (S 0.038 wt.%) and Heat L2 (S 0.034 wt.%). Tables 7 and 8 summarize the results obtained from non-irradiated specimens.



### Figure 3.

Yield strength of non-irradiated control specimens of model stainless steel alloys tested at 288°C in BWR-simulated water containing ≈8 ppm DO.

## Figure 4.

Total elongation of nonirradiated control specimens of model stainless steel alloys tested at 288°C in BWRsimulated water containing ≈8 ppm DO.

ALLOY AND	)		FEEDWATE		SSRT PA	RAMETERS	FRACTURE BEHAVIOR					
Specimen		OXYGEN	AVERAGE	COND.		YIELD	Max.	UNIFORM	TOTAL			TGSCC+
IDENT.	SSRT	CONC.	ECP	ат 25°С	рН	STRESS	STRESS	ELONG.	ELONG.	TGSCC	IGSCC	IGSCC
No.	No.	(PPM)	(MV SHE)	(μS·CM <sup>1</sup> )	AT 25°C	(MPA)	(MPA)	(%)	(%)	(%)	(%)	(%)
L23-4	CHR-1	8.6	+228	0.07	6.65	332	480	15.6	17.0	15	0	15
L7-4	CHR-2	8.0	+217	0.07	7.37	195	370	2.5	5.2	20	0	20
L7-B1	CHR-7	TESTED IN AIR				282	676	42.3	43.9	0	0	0
L14-4	CHR-3	8.6	+208	0.07	7.37	240	474	41.8	44.2	0	0	0
L17-4	CHR-4	7.5	+262	0.06	7.09	189	412	11.6	13.3	60	0	60
L17-B1	CHR-19	7.8	+166	0.08	6.71	184	447	30.1	31.2	8	0	8
L6-4	CHR-5	7.9	+256	0.08	6.85	227	545	43.0	44.5	0	0	0
L27-4	CHR-6	9.3	+247	0.08	6.96	298	483	20.6	22.9	0	0	0
L26-4	CHR-8	9.4	+223	0.07	6.65	184	596	38.2	40.2	0	0	0
L2-4	CHR-9	8.6	+292	0.06	6.55	193	348	6.6	7.8	57	0	57
L25-4	CHR-10	8.2	+239	0.06	6.42	184	458	25.5	27.0	0	0	0
L15-4	CHR-11	8.2	+195	0.06	6.32	218	512	36.7	37.9	0	0	0
L24-4	CHR-12	8.4	+200	0.07	6.20	352	461	10.4	12.3	10	0	10
C1-15	CHR-13	8.1	+187	0.07	6.33	179	498	49.4	51.7	0	0	0
C19-B1	CHR-14	8.8	+179	0.08	6.29	178	501	47.4	49.2	0	0	0
C9-B1	CHR-15	8.5	+166	0.07	6.83	178	408	17.4	19.4	32	0	32
C12-B1	CHR-16	8.5	+124	0.07	6.18	182	511	46.0	47.6	0	0	0
C10-B1	CHR-17	9.2	+145	0.07	6.26	174	478	30.6	35.1	0	0	0
C21-9	CHR-18	9.2	+187	0.07	6.41	277	455	48.9	59.5	0	0	0

 Table 7.
 Results of SSRT test and SEM fractography for non-irradiated control specimens of model austenitic stainless steel alloys.

 Table 8.
 Composition (wt.%) of non-irradiated control specimens of model austenitic stainless steels correlated with results of SSRT test and SEM fractography.

Alloy ID	Ni	Si	P	s	Mn	с	N	Cr	Mo or Nb	O wppm	YS (MPa)	UTS (MPa)	UE (%)	TE (%)	TG (%)	IG (%)
L23	12.04	0.68	0.030	0.047	0.96	0.043	0.092	17.30	Nb 1.06	93	332	480	15.6	17.0	15	0
L7	10.60	0.18	0.040	0.038	1.02	0.007	0.111	15.40	-	274	195	370	2.5	5.2	20	0
L14	7.93	1.49	0.080	0.002	1.76	0.107	0.028	15.00	-	45	240	474	41.8	44.2	0	0
L17	8.00	0.66	0.090	0.009	0.48	0.061	0.078	15.30	-	90	189	412	11.6	13.3	60	0
L6	10.00	1.90	0.020	0.005	1.13	0.096	0.087	17.10	-	58	227	515	43.0	44.5	0	0
L27	10.30	0.96	0.040	0.002	0.97	0.057	0.019	15.30	Mo 2.01	58	298	483	20.6	22.9	0	0
L26	8.09	0.79	0.004	0.002	0.91	0.070	0.089	17.20	-	80	184	506	38.2	40.2	0	0
L2	10.50	0.82	0.080	0.034	1.58	0.074	0.102	17.02	-	66	193	348	6.6	7.8	57	0
L25	8.93	0.92	0.020	0.008	1.54	0.019	0.095	17.20	-	85	184	458	25.5	27.0	0	0
L15	8.00	1.82	0.010	0.013	1.07	0.020	0.085	17.80	-	110	218	512	36.7	37.9	10	0
L24	12.30	0.03	0.007	0.005	0.48	0.031	0.002	16.90	Nb 1.72	129	352	461	10.4	12.3	0	0
Cl	8.12	0.50	0.038	0.002	1.00	0.060	0.060	18.11	-	102	179	498	49.4	51.7	0	0
C19	8.08	0.45	0.031	0.003	0.99	0.060	0.070	18.21	-	200	178	501	47.4	49.2	0	0
C9	8.75	0.39	0.013	0.013	1.72	0.062	0.065	18.48	-	102	178	408	17.4	19.4	32	0
C12	8.23	0.47	0.018	0.002	1.00	0.060	0.070	18.43	-	-	182	511	46.0	47.6	0	0
C10	8.13	0.55	0.033	0.002	1.00	0.060	0.086	18.19	-	74	174	478	30.6	35.1	0	0
C21	10.24	0.51	0.034	0.001	1.19	0.060	0.020	16.28	Mo 2.08	112	277	455	48.9	59.5	0	0_

Load-elongation behavior of Type 304 SS Heat L7 exhibited significantly deleterious effects of water even in the nonirradiated state, as shown in Fig. 5. Approximately 20% of the fracture surface of the specimen was characterized by transgranular fracture morphology that was mostly covered with unidentified corrosion debris (Fig. 6). Although it was suspected that such corrosion debris contained a high concentration of S, positive identification of the debris could not be determined.



The five heats that exhibited relatively low ductility were characterized by relatively high susceptibility to TGSCC. Other heats were very ductile and exhibited predominantly ductile dimple fracture surface morphology. However, no evidence of susceptibility to IGSCC was observed in any of the 17 heats that were tested in the BWR-like water at 289°C in the nonirradiated state.

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# 6 SSRT SCC Test and Fractography on Low-Fluence Specimens (≈0.4 dpa)

## 6.1 Tabulation of Test Results

Results of SSRT tests on specimens irradiated to  $\approx 0.3 \times 10^{21} \text{ n} \cdot \text{cm}^{-2}$  (E > 1 MeV) ( $\approx 0.4$  dpa) are summarized in Tables 9 and 10. Feedwater chemistry (i.e., DO, ECP, conductivity, and pH) and SSRT behavior in water (i.e., 0.2%-offset yield strength, maximum strength, uniform strain, and total strain) are given in Table 9. Also shown in these tables are results of SEM fractographic analysis of the failure mode (i.e., ductile, intergranular, and transgranular fracture surface morphology) of the specimens. In Table 10, the results of SSRT and SEM fractographic analysis (percent TGSCC and percent IGSCC) are correlated with the compositions of the low-fluence specimens.

Table 9. Results of SSRT tests in 289°C water (DO  $\approx$ 8 ppm) and SEM fractography of steels irradiated to  $\approx$ 0.3 x 10<sup>21</sup> n·cm<sup>-2</sup> (E > 1 MeV).

Specimen	Specimen Oxygen		Average	Cond.	pН	Yield	Max.	Uniform	Total	Fracture	
Ident.	SSRT	Conc.	ECP	at 25°C	at	Stress	Stress	Elong.	Elong.	TGSCC	IGSCC
No.	No.	(ppm)	(mV SHE)	(μS cm <sup>-1</sup> )	25°C	(MPa)	(MPa)	(%)	(%)	(%)	(%)
C1-1	HR-1	8.3	+184	0.07	7.03	490	680	13.4	16.6	4	0
L5-1	HR-2	9.7	+208	0.07	6.89	413	539	29.5	32.7	2	2
L22-1	HR-3	8.0	+236	0.07	6.80	360	596	6.6	9.4	50	15
C3-1	HR-4	8.7	+161	0.07	6.68	338	491	27.7	31.6	5	0
C16-1	HR-5	8.3	+204	0.08	6.74	370	527	17.6	20.6	2	0
L4-1	HR-6	<b>9.0</b> .	+202	0.08	6.70	367	542	19.7	22.3	46	0
L18-1	HR-7	9.0	+203	0.08	6.33	503	572	6.3	8.8	54	0
C10-1	HR-8	8.2	, +174	0.07	6.35	523	640	17.4	18.9	6	0
C21-1	HR-9	8.1	+149	0.08	6.49	480	620	15.9	19.4	4	0
L11-1	HR-10	9.0	+157	0.08	6.17	487	599	2.3	3.8	62	0
L13-1	HR-11	8.7	+164	0.08	6.17	248	461	22.1	24.8	18	0
L20-1	HR-12	8.4	+174	0.07	6.20	454	552	2.9	5.1	denc	irite
C19-1	HR-13	9.5	+132	0.12	6.36	554	682	10.5	14.7	7	0
C9-1	HR-14	8.0	+192	0.11	6.30	522	607	13.4	14.6	24	0
C12-1	HR-15	9.0	+195	0.08	6.40	404	589	20.4	24.2	5	0
L8-1	HR-16	9.0	+215	0.08	6.60	411	571	15.6	17.9	64	0

#### 6.2 Susceptibility to IGSCC

Even at the low damage level of  $\approx 0.4$  dpa, heat-to-heat variations in susceptibility to IGSCC and TGSCC were significant among the steels tested and listed in Table 10. A high-purity Heat L22 of Type 316L SS, which contains very low concentrations of Si (0.024 wt.%) and C (0.003 wt.%), exhibited relatively low ductility and the highest susceptibility to IASCC (i.e., the highest percent IGSCC) among the 16 tested heats. The high-purity Heat L20 of Type 304L SS, which contains very low concentrations of Si (0.017 wt.%) and C (0.002 wt.%), also exhibited an indication of slight susceptibility to IGSCC. Susceptibility of all other heats to IGSCC was insignificant (see Fig. 7).

At this low fluence, the limited observation in Fig. 7 seems to indicate that very low concentrations of Si and C are deleterious from the standpoint of IASCC. The susceptible behavior of the two high-purity heats in the figure appears to be consistent with similar behavior of BWR neutron absorber tubes fabricated from

high-purity 304L SSs which exhibited relatively higher susceptibility during SSRT tests performed after irradiation to  $0.2 \times 10^{21} \text{ n} \cdot \text{cm}^{-2}$  to  $1.4 \times 10^{21} \text{ n} \cdot \text{cm}^{-2}$  (E > 1 MeV).<sup>19</sup> However, field failure by IASCC in a BWR usually occurs at higher fluence levels, e.g., >1.5 x  $10^{21} \text{ n} \cdot \text{cm}^{-2}$  (E >1 MeV). Therefore, a comprehensive database at higher fluence levels is necessary to be able to evaluate the roles of Si, C, and other impurity elements.

Figure 8 shows that limited intergranular fracture surface morphology is surrounded by transgranular morphology in Heat L22, indicating that transgranular fracture made a transition to intergranular fracture. Heat L22 also exhibited relatively high susceptibility to IGSCC after irradiation to higher fluence (Section 7).

Spec.										YS	UTS	UE	TE	TG	IG
ID	Ni	Si	P	<u>S</u>	Mn	С	N	Cr	Others	(MPa)	(MPa)	(%)	(%)	(%)	(%)
C1-1	8.12	0.50	0.038	0.002	1.00	0.060	0.060	18.11	O 0.0102	490	680	13.4	16.6	4	0
L5-1 <sup>a</sup>	9.66	0.90	0.113	0.028	0.47	0.006	0.033	21.00	O 0.0068	413	539	29.5	32.7	2	2
									≈3% δ <sup>a</sup>						
L22-1	13.30	0.024	0.015	0.004	0.40	0.003	0.001	16.10	O 0.0103	360	596	6.6	9.4	50	15
									Mo 2.04						
C3-1	8.91	0.46	0.019	0.004	1.81	0.016	0.083	18.55	-	338	491	27.7	31.6	5	0
C16-1	12.90	0.38	0.014	0.002	1.66	0.020	0.011	16.92	O 0.0157	370	527	17.6	20.6	2	0
									Mo 2.30						
L4-1	10.20	0.94	0.031	0.010	1.75	0.110	0.002	15.80	O 0.0037	367	542	19.7	22.3	46	0
L18-1	8.13	0.14	0.016	0.033	1.13	0.080	0.001	18.00	O 0.0055	503	572	6.3	8.8	54	0
C10-1	8.13	0.55	0.033	0.002	1.00	0.060	0.086	18.19	O 0.0074	523	640	17.4	18.9	6	0
C21-1	10.24	0.51	0.034	0.001	1.19	0.060	0.020	16.28	O 0.0112	480	620	15.9	19.4	4	0
									Mo 2.08						
L11-1	8.15	0.47	0.097	0.009	1.02	0.014	0.004	17.40	O 0.0071	487	599	2.3	3.8	62	0
L13-1	8.18	1.18	0.027	0.022	0.36	0.026	0.001	17.40	O 0.0042	248	461	22.1	24.8	18	0
L20-1	8.91	0.017	0.010	0.004	0.41	0.002	0.002	18.10	O 0.0940	454	552	2.9	5.1	den	drite
C19-1	8.08	0.45	0.031	0.003	0.99	0.060	0.070	18.21	O 0.0200	554	682	10.5	14.7	7	0
C9-1	8.75	0.39	0.013	0.013	1.72	0.062	0.065	18.48	O 0.0102	522	607	13.4	14.6	24	0
C12-1	8.23	0.47	0.018	0.002	1.00	0.060	0.070	18.43	-	404	589	20.4	24.2	5	0
L8-1	10.20	0.15	0.093	0.010	1.85	0.041	0.001	18.30	O 0.0059	411	571	15.6	17.8	64	0

Table 10. Composition (wt.%) of steels irradiated to  $\approx 0.3 \times 10^{21} \text{ n-cm}^{-2}$  (E > 1 MeV) correlated with results of SSRT tests and SEM fractography.

<sup>a</sup>Heat L5 contains ≈3 vol.% globular delta ferrite.



Figure 7. Percent IGSCC of model SS alloys irradiated in He in the Halden reactor to fluence of  $\approx 0.3 \times 10^{21}$ n·cm<sup>-2</sup> (E > 1 MeV) and tested at 289°C in BWR-like water.

Figure 8. Limited intergranular fracture surface surrounded by transgranular surface in 316L-like SS Heat L22 irradiated to  $\approx 0.3 \times 10^{21} \text{ n} \cdot \text{cm}^{-2}$  (E > 1 MeV,  $\approx 0.4 \text{ dpa}$ ) and tested at 289°C in BWR-like water.

## 6.3 Effect of Si on Strength

The relatively higher susceptibility of the high-purity heat of Type 316L SS (i.e., Heat L22) when compared with the commercial counterpart (Heat C21) is of particular interest. This behavior is similar to that observed for BWR neutron absorber tubes fabricated from high- and commercial-purity heats of Type 304 SS.<sup>23</sup> In another investigation similar to that of the present study, Jenssen and Ljunberg irradiated U-notched rod specimens that had been fabricated from two heats of Type 316 SS, then conducted postirradiation SSRT tests in a BWR loop under conditions of normal oxidizing water chemistry.<sup>30</sup> In their study, a commercial-purity heat of Type 316 SS (Heat K) was resistant to IASCC, whereas a high-purity heat of Type 316L SS (Heat F) was susceptible. The two relatively more susceptible heats (i.e., Heats L22 and F) are characterized by an unusually low Si concentration of <0.26 wt.%, whereas the two relatively more resistant heats (i.e., Heats C21 and K) contain a higher Si concentration of >0.5 wt.%.

An interesting observation from Table 10 is the effect of Si content on the alloy strength that was measured in water before and after irradiation. Yield strength of unirradiated commercial heats of Types 304 and 304L SS, measured in water, was similar to that measured in air,<sup>1</sup> i.e., 174-182 MPa vs. 170-175 MPa, indicating that the effect of water is insignificant. The yield and maximum strengths (in water) are plotted in Fig. 9 as function of Si content for the unirradiated and low-fluence specimens. Before irradiation, 304- and 304L-type steels did not exhibit a systematic effect of Si content. However, after irradiation to  $\approx 0.3 \times 10^{-21}$ n·cm<sup>-2</sup> (E > 1 MeV), yield and maximum strengths decreased as Si content increased to 0.8-1.2 wt.%.

The influence of Si content on the material strength indicates that, at high concentrations, Si atoms suppress formation of some types of irradiation-induced hardening centers and contribute to greater ductility, at least at low fluence levels. Tsukada et al.<sup>29</sup> irradiated two SSRT specimens of high-purity Type 304L SS (C 0.003, Si 0.01, Mn 1.36, P 0.001, S 0.0014, and N 0.0014 wt.%) in He at  $\approx 240^{\circ}$ C in the JRR-3 reactor to a fluence of  $\approx 0.67 \times 10^{21}$ n cm<sup>-2</sup> (E > 1 MeV). One specimen had been doped with  $\approx 0.69$  wt.% Si and the other had not been doped. After SSRT testing of the irradiated specimens at  $\approx 300^{\circ}$ C in high-purity water (DO  $\approx 32$  ppm), they observed that the Si-doped specimen exhibited significantly greater ductility than the undoped specimen. Silicone effect at 1.3 dpa is discussed in Section 7.4.

#### 6.4 Correlation of S Content and Susceptibility to TGSCC of 304 and 304L SSs

For the fluence level of  $\approx 0.3 \times 10^{21} \text{ n} \cdot \text{cm}^{-2}$  (E > 1 MeV), SSRT tests in 289°C water and fractographic analysis were performed for 11 heats of Type 304 or 304L SS, three heats of Type 316 or 316L SS, and one heat of austenitic-ferritic duplex SS. Susceptibilities to TGSCC of the 11 heats of Type 304 or 304L SS varied strongly from heat to heat. However, it was found that bulk S content correlates reasonably well with percent TGSCC, as is shown in Fig. 10. Heats that contain  $\geq 0.009$  wt.% S exhibited high susceptibility to TGSCC, whereas heats that contain  $\leq 0.004$  wt.% S exhibited low susceptibility to TGSCC.

For a given heat of all types of austenitic SSs at low fluence, significant susceptibility to TGSCC is often observed after an SSRT test, even though susceptibility to IGSCC is insignificant. As fluence level increases, the susceptibility to TGSCC peaks at a certain fluence specific to the heat, and at the same time, the susceptibility to IGSCC increases at the expense of TGSCC. Considering this trend, the good correlation observed between S content and percent TGSCC in Fig. 10 is considered an important indication that may provide useful insight into the susceptibility to IGSCC at higher fluence levels.



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# 7. SSRT Test and SEM Fractography on Medium-Fluence Specimens (≈1.3 dpa)

### 7.1 Tabulation of Test Results

Slow-strain-rate tensile tests in 289°C BWR-like water and SEM fractographic analysis were performed on 26 medium-fluence specimens irradiated to  $\approx 0.9 \times 10^{21} \text{ n} \cdot \text{cm}^{-2}$  (E > 1 MeV) ( $\approx 1.3 \text{ dpa}$ ). Feedwater chemistry (i.e., DO, ECP, conductivity, and pH) and results from SSRT testing (i.e., 0.2%-offset yield strength, maximum strength, uniform plastic strain, and total plastic strain) are summarized in Table 11. Also shown in the table are results of SEM fractographic analysis of the failure mode (i.e., intergranular and transgranular fracture surface morphology) of the specimens. In Table 12, the results of SSRT and SEM fractographic analysis are correlated with compositional characteristics of the specimens.

For Heat L20, which contains an unusually high concentration of O (0.094 wt.) and a dendritic grain structure, fractographic analysis was not performed, although SSRT testing was conducted. This heat, which is not a prototypic austenitic SS, exhibited poor ductility when compared with other steels, and its behavior was excluded in the analyses of SCC test results.

### 7.2 Heat-to-Heat Variations in Susceptibility to TGSCC and IGSCC

As an example, effects of fluence and test environment on load-elongation behavior are shown for Type 304 SS commercial Heat C19 in Fig. 11. As fluence increased from  $\approx 0.3 \times 10^{21} \text{ n} \cdot \text{cm}^{-2}$  to  $\approx 0.9 \times 10^{21} \text{ n} \cdot \text{cm}^{-2}$ , maximum strength increased slightly (from 690 MPa to 800 MPa). In contrast, total elongation decreased significantly (from 1.45% to 0.7%).

Figures 12 and 13 show, respectively, heat-to-heat variations in percent TGSCC and IGSCC for the two tested fluence levels, i.e.,  $\approx 0.3 \times 10^{21} \text{ n} \cdot \text{cm}^{-2}$  and  $\approx 0.9 \times 10^{21} \text{ n} \cdot \text{cm}^{-2}$ . As fluence increased, percent TGSCC decreased significantly, and, at the same time, the percent IGSCC increased significantly for many heats, e.g., L22, L18, L11, and L8. The fact that the IGSCC fracture surfaces in these alloys were surrounded by TGSCC fracture surfaces indicates that an IGSCC fracture surface emerges from a portion of a TGSCC fracture surface. In addition, for many alloys, high susceptibility to TGSCC is a precursor to susceptibility to IGSCC at a higher fluence. However, the threshold fluence for the transition from TGSCC to IGSCC differed significantly from alloy.

		_	ECP	Cond.	pН	Yield	Max.	Uniform	Total			TG+IG
Speci.	SSRT	DO	(mV	at 25°C	at	Stress	Stress	Elong.	Elong.	TGSCC	IGSCC	SCC
ID	No.	(ppm)	SHE)	(µS cm <sup>-1</sup> )	25°C	(MPa)	(MPa)	(%)	(%)	(%)	(%)	(%)
L22-02	HR-17	8.0	+181	0.08	6.77	475	549	4.20	5.82	30	35	65
L11-02	HR-18	8.0	+191	0.08	6.55	820	856	0.43	1.65	50	14	64
L18-02	HR-19	8.0	+193	0.10	6.07	710	755	3.98	5.05	38	14	52
L20-02	HR-28		test in	289°C air		826	845	0.31	2.09	dendritic	struc	ture
L20-05	HR-26	9.0	+182	0.09	6.32	670	743	0.37	1.03	dendritic	struc	ture
L20-06	HR-27	8.0	+274	0.07	6.05	632	697	0.85	2.72	0	0	0
C9-02	HR-21	8.0	+240	0.07	6.47	651	679	1.42	2.50	62	22	84
L17-02	HR-22	8.0	+198	0.07	6.42	574	654	2.02	3.08	44	41	85
L7-02	HR-23	8.0	+215	0.07	6.03	553	561	0.24	2.44	38	54	92
C10-02	HR-24	7.0	+221	0.07	5.26	651	706	6.35	9.25	14	0	14
C3-02	HR-25	8.0	+240	0.07	6.34	632	668	16.72	19.74	9	4	13
C19-05	HR-58		test in	23°C air		1130	1260	15.50	15.70	-	-	-
C19-02	HR-30		test in	289°C air		888	894	6.41	10.21	1	0	1
C19-04	HR-31	8.0	+252	0.07	6.18	750	769	6.06	8.79	1	0	1
L6-02	HR-32	8.0	+250	0.07	6.40	493	546	2.45	3.77	8	27	35
L14-02	HR-33	8.0	+246	0.08	6.07	649	684	1.90	4.67	84	2	86
L13-02	HR-34	7.0	+222	0.09	6.85	602	624	1.67	4.95	55	12	67
L4-02	HR-35	7.0	+259	0.08	6.54	634	680	1.07	2.02	58	12	70
L5-02	HR-36	7.0	+243	0.07	6.85	665	725	3.07	4.57	3	5	8
C16-02	HR-37	7.0	+230	0.07	6.62	562	618	11.99	15.80	7	1	8
L8-02	HR-38	8.0	+242	0.07	6.57	838	838	0.12	3.12	15	22	37
C21-02	HR-39	8.0	+231	0.08	6.21	643	716	15.38	18.30	1	2	3
L2-02	HR-40	7.0	+239	0.07	7.11	839	849	0.88	1.56	31	11	42
L24-02	HR-41	8.0	+239	0.06	6.40	725	725	0.15	2.45	2	1	3
L23-02	HR-42	7.0	+237	0.08	6.60	787	818	0.38	1.24	3	24	27
C12-02	HR-43	7.0	+227	0.07	6.19	747	756	14.96	18.57	4	0	4
C1-02	HR-44	8.0	+229	0.07	6.30	707	763	13.36	17.04	2	0	2

Table 11. Results of SSRT test and fractography for model austenitic stainless steels irradiated to  $\approx 0.9 \times 10^{21} \text{ n cm}^{-2}$  (E > 1 MeV) ( $\approx 1.3 \text{ dpa}$ ).

#### 7.3 Effect of Sulfur

The information given in Tables 11 and 12 led us to carefully examine the relationship between fracture properties and compositional characteristics of the alloys in an effort to identify the alloying and impurity elements that strongly influence stress corrosion cracking behavior. Of the steel heats tested and listed in Table 12, 17 heats are Type 304 or 304L, three heats are Type 316 or 316L, two heats are Type 348 or 348L, and one heat is duplex steel that contains  $\approx 3\%$  delta ferrite.

Of the 17 heats of Types 304 or 316 SS, steels that contain low concentrations of S (<0.004 wt.% S) exhibited high resistance to IGSCC (see Fig. 14) and high uniform and total elongation (see Fig. 15), whereas steels that contain relatively higher concentrations of S (>0.005 wt.% S) exhibited poor resistance to IGSCC and low uniform and total elongation.

The observations summarized in Figs. 14 and 15 indicate strongly that, for Types 304 and 304L SS, a high concentration of S (>0.005 wt.%) is significantly detrimental, and that a sufficiently low concentration of S (e.g., <0.004 wt.%) is beneficial to render good resistance to IASCC at  $\approx$ 1.3 dpa.

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Spec.						_		~			YS	UTS	UE	TE	TG	IG
	Ni	Sı	4	<u> </u>	Mn	<u>C</u>	<u>N</u>	Cr	0	Other	(MPa)	(MPa)	<u>(%)</u>	(%)	<u>(%)</u>	(%)
L22-0?	13.30	0.024	0.015	0.004	0.40	0.003	0.001	16.10	0.0102	Mo 2.04	475	549	4.20	5.82	30	35
L11-02	8.15	0.47	0.097	0.009	1.02	0.014	0.004	17.40	0.0071	-	820	856	0.43	1.65	50	14
L18-02	8.13	0.14	0.016	0.033	1.13	0.080	0.001	18.00	0.0055	-	710	755	3.98	5.05	38	14
L20-05	8.91	0.017	0.010	0.004	0.41	0.002	0.002	18.10	0.0940	-	670	743	0.37	1.03	dendr	ite
L20-06	8.91	0.017	0.010	0.004	0.41	0.002	0.002	18.10	0.0940	-	632	697	0.85	2.72	dendr	ite
C9-02	8.75	0.39	0.013	0.013	1.72	0.062	0.065	18.48	0.0102	-	651	679	1.42	2.50	62	22
L17-02	8.00	0.66	0.090	0.009	0.48	0.061	0.078	15.30	0.0090	-	574	654	2.02	3.08	44	41
L7-02	10.60	0.18	0.040	0.038	1.02	0.007	0.111	15.40	0.0274	-	553	561	0.24	2.44	38	54
C10-02	8.13	0.55	0.033	0.002	1.00	0.060	0.086	18.19	0.0074	-	651	706	6.35	9.25	14	0
C3-02	8.91	0.46	0.019	0.004	1.81	0.016	0.083	18.55		-	632	668	16.72	19.74	9	4
C19-04	8.08	0.45	0.031	0.003	0.99	0.060	0.070	18.21	0.0200	-	750	769	6.06	8.79	1	0
L6-02	10.00	1.90	0.020	0.005	1.13	0.096	0.087	17.10	0.0058	-	493	546	2.45	3.77	8	27
L14-02	7.93	1.49	0.080	0.002	1.76	0.107	0.028	15.00	0.0045	-	649	684	1.90	4.67	84	2
L13-02	8.18	1.18	0.027	0.022	0.36	0.026	0.001	17.40	0.0042		602	624	1.67	4.95	55	12
L4-02	10.20	0.94	0.031	0.010	1.75	0.110	0.002	15.80	0.0037	-	634	680	1.07	2.02	58	12
L5-02 <sup>a</sup>	9.66	0.90	0.113	0.028	0.47	0.006	0.033	21.00	0.0068	<sup>a</sup> 3% δ	665	725	3.07	4.57	3	5
C16-02	12.90	0.38	0.014	0.002	1.66	0.020	0.011	16.92	0.0157	Mo 2.30	562	618	11.99	15.80	7	1
L8-02	10.20	0.15	0.093	0.010	1.85	0.041	0.001	18.30	0.0059	-	838	838	0.12	3.12	15	22
C21-02	10.24	0.51	0.034	0.001	1.19	0.060	0.020	16.28	0.0112	Mo 2.08	643	716	15.38	18.30	1	2
L2-02	10.50	0.82	0.080	0.034	1.58	0.074	0.102	17.02	0.0065	-	839	849	0.88	1.56	31	11
L24-02	12.30	0.03	0.007	0.005	0.48	0.031	0.002	16.90	0.0129	Nb 1.72	725	725	0.15	2.45	2	1
L23-02	12.04	0.68	0.030	0.047	0.96	0.043	0.092	17.30	0.0093	Nb 1.06	787	818	0.38	1.24	3	24
C12-02	8.23	0.47	0.018	0.002	1.00	0.060	0.070	18.43	-	-	747	756	14.96	18.57	4	0
<u>C1-02</u>	8.12	0.50	0.038	0.002	1.00	0.060	0.060	18.11	0.0102		707	763	13.36	17.04	2	0

Table 12. Composition (wt.%) of stainless steels irradiated to  $\approx 0.9 \times 10^{21} \text{ n cm}^{-2}$  (E > 1 Me\') (1.3 dpa) correlated with results of SSRT test and fractography.

<sup>a</sup>L5-02 contains ≈3 vol.% globular-shaped delta ferrite.

### 7.4 Effect of Silicon

The yield strength of the model steel alloys, measured in BWR-like water at 289°C (DO  $\approx$ 8 ppm), v/as nearly constant at  $\approx$ 200 MPa in the unirradiated state and was more or less independent of Si concentration (see Fig. 16). However, as fluence was increased to  $\approx$ 0.3 x 10<sup>21</sup> n·cm<sup>-2</sup> and  $\approx$ 0.9 x 10<sup>21</sup> n·cm<sup>-2</sup>, the degree of increase in yield strength was significantly lower for alloys that contain >0.9 wt.% Si than for alloys that contain <0.8 wt.% Si. This behavior indicates that the nature of irradiation-induced hardening centers and the degree of irradiation hardening are influenced by Si. Silicon atoms in austenitic stainless steels occupy substitutional sites; therefore, they are likely to interact preferentially with irradiation-induced vacancy sizes. This effect is likely to inhibit the formation of vacancy clusters or vacancy-impurity complexes, and is, therefore, conducive to a less significant degree of irradiation-induced hardening. An effect similar to that of Si was, however, not observed for C and N.



Figure 11. Effects of fluence and test environment on load elongation behavior of Type 304 SS commercial heat C19.

Figure 12. Effects of fluence on percent TGSCC measured in 289°C water containing ≈8 ppm DO, 0.4 to 1.3 dpa.

C01

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C07\_



Figure 15. Effect of S on uniform and total elongation of Type 304 and 304L steels irradiated to ≈0.9 x 10<sup>21</sup> n·cm<sup>-2</sup> (E > 1 MeV); one group of steels contains <0.004 wt.% S and the other group contains >0.005 wt.% S.

Among the laboratory-fabricated heats of Types 304 and 304L SS that contain relatively high concentrations of S (0.009-0.038 wt.%), alloys that contain  $\leq 0.66$  wt.% Si exhibited significant susceptibility to IGSCC, whereas alloys with  $\geq 0.82$  wt.% Si exhibited relatively lower susceptibility to IGSCC (see Fig. 17). However, Heat L6, which contains  $\approx 1.9$  wt.% Si, exhibited susceptibility to IGSCC (percent IGSCC  $\approx 27$ ). These observations indicate that Si concentration in the range of  $\approx 0.8$  to  $\approx 1.5$  wt.% is beneficial in delaying the onset of or suppressing the susceptibility to IASCC, at least up to 1.3 dpa.

However, it is not clear if such a beneficial effect of Si (in the range of  $\approx 0.8$  to  $\approx 1.5$  wt.%) is also significant at higher fluence levels. Considering the strongly deleterious effect of S shown in Figs. 14 and 15, it is possible that, as fluence increases to higher levels, the deleterious effect of S more than offsets the beneficial effect of Si. As such, the effect of Si could be secondary, that is, manifested only at low fluence levels.

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# 8. SSRT SCC Test and Fractography on High-Fluence Specimens (≈3.0 dpa)

## 8.1 Tabulation of Test Results

Twenty-four steel alloys were tested by the SSRT technique after irradiation to  $\approx 2.0 \times 10^{21}$  n cm<sup>-2</sup> (E > 1 MeV) ( $\approx 3.0$  dpa) in BWR-like oxidizing water at 289°C (DO  $\approx 8$  ppm). Of the 24 alloys, 18 were 304- or 304L-type steels, 3 were 316- or 316L-type steels, 2 were 348- or 348L-type steels, and one alloy was austenitic-ferritic duplex steel (delta ferrite  $\approx 3$  vol.%). At this fluence, some of the specimens (Heats L13 and L8) were so brittle that they failed in 23°C air during assembly of the test train. Table 13 summarizes the test conditions and results of the SSRT testing and fractography. Table 14 summarizes correlation of the fracture characteristics with chemical composition of the steels.

Table 13. Results of SSRT tests and SEM fractography for model austenitic stainless steels irradiated to  $\approx 2.0 \times 10^{21} \text{ n} \cdot \text{cm}^{-2}$  (E > 1 MeV) ( $\approx 3.0 \text{ dpa}$ )

Test		Oxygen				Yield	Max.	Uniform	Total		
Speci.	SSRT	Conc.	ECP	Cond.	pH at	Stress	Stress	Elong.	Elong.	TG	IG
ID	No.	(ppm)	(mV SHE)	(µScm <sup>-1</sup> )	25°C	(MPa)	(MPa)	(%)	(%)	(%)	(%)
L4-03	HR-45	9.0	+198	0.09	6.5	876	1068	2.49	3.27	5	95
C12-03	HR-46	8.0	+208	0.09	6.5	922	<b>9</b> 96	1.28	2.73	8	2
C9-03	HR-47	8.0	+212	0.11	6.5	earl	ly failure in	289°C wate	r	6	94
L5-03	HR-48	8.0	+204	0.10	6.5	953	985	0.59	2.97	2	4
C19-08	Hr-59		test in	289°C air		1001	1001	0.27	1.95	-	-
C19-09	HR-60		• test in	325°C air		<del>9</del> 98	998	0.27	2.02	-	-
C19-03	HR-49	8.0	+171	0.11	6.5	787	801	0.89	3.32	2	62
C16-03	HR-50	8.0	. +202	0.10	6.5	766	803	0.83	1.84	2	29
C10-03	HR-51	8.0	+167	0.11	6.5	1062	1065	3.15	4.51	3	0
L18-03	HR-52	8.0	+169	0.11	6.5	795	779	0.35	1.75	5	86
L13-03	HR-53	-	_	brittle fractur	e during	test preparat	tion in air		_	-	_
C1-03	HR-54	8.0	+161	0.10	6.5	802	833	3.38	5.27	2	0
C3-03	HR-55	8.0	+160	0.10	6.5	796	826	5.05	7.31	0	26
C21-03	HR-56	8.0	+156	0.10	6.5	893	924	1.89	4.85	2	1
L8-03	HR-57	-		brittle fractur	e during t	lest preparat	ion in air		-		-
L26-01	HR-61	8.0	+221	0.11	6.5	795	821	0.58	1.83	32	2
L25-01	HR-62	8.0	+220	0.09	6.5	867	867	0.05	0.48	8	43
L2-03	HR-63	8.0	+204	0.10	6.5	916	916	0.26	0.50	58	32
L7-03	HR-64	8.0	+196	0.11	6.5	889	889	0.02	0.05	10	52
L23-03	HR-65	8.0	+189	0.11	6.5	887	887	0.06	0.41	16	22
L17-03	HR-66	8.0	+188	0.11	6.5	1105	1116	0.23	0.28	56	20
L14-03	HR-67	8.0	+174	0.11	6.5	917	951	1.64	2.01	16	0
L27-01	HR-68	8.0	+175	0.11	6.5	776	808	0.73	1.02	12	2
L15-03	HR-69	8.0	+180	0.11	6.5	946	946	0.78	1.17	8	32
L6-03	HR-70	8.0	+171	0.11	6.5	794	805	2.02	2.44	12	36
L24-03	HR-71	8.0	+173	0.11	6.5	862	862	0.11	0.86	3	0

Test				·							Unif	Total	
Speci											Elong.	Elong.	IG
ID	Ni	Si	Р	S	Mn	С	N	Cr	0	Other	(%)	(%)	(%)
L4-03	10.20	0.94	0.031	0.0100	1.75	0.110	0.002	15.80	0.0037	_	2.49	3.27	95
C12-03	8.23	0.47	0.018	0.0020	1.00	0.060	0.070	18.43	-	-	1.28	1.73	2
C9-03	8.75	0.39	0.013	0.0130	1.72	0.062	0.065	18.48	0.0102	-	-	-	94
L5-03 <sup>a</sup>	9.66	0.90	0.113	0.0283	0.47	0.006	0.033	21.00	0.0068	≈3 % δ <sup>a</sup>	0.59	2.97	4
C19-03	8.08	0.45	0.031	0.0030	0.99	0.060	0.070	18.21	0.0200	-	0.89	3.32	62
C16-03	12.90	0.38	0.014	0.0020	1.66	0.020	0.011	16.92	0.0150	Mo 2.30	0.83	1.84	29
C10-03	8.13	0.55	0.033	0.0020	1.00	0.060	0.086	18.19	0.0074	-	3.15	4.51	0
L18-03	8.13	0.14	0.016	0.0326	1.13	0.080	0.001	18.00	0.0055	-	0.35	1.75	86
L13-03	8.18	1.18	0.027	0.0222	0.36	0.026	0.001	17.40	0.0042	-	-		-
C1-03	8.12	0.50	0.038	0.0020	1.00	0.060	0.060	18.11	0.0102	-	3.38	5.27	0
C3-03	8.91	0.46	0.019	0.0040	1.81	0.016	0.083	18.55	-	-	5.05	7.31	26
C21-03	10.24	0.51	0.034	0.0010	1.19	0.060	0.020	16.28	0.0112	Mo 2.08	1.89	4.85	1
L8-03	10.20	0.15	0.093	0.0100	1.85	0.041	0.001	18.30	0.0059	-	-	~	-
L26-01	8.09	0.79	0.004	0.0022	0.91	0.070	0.089	17.20	0.0080	-	0.58	1.83	2
L25-01	8.93	0.92	0.020	0.0082	1.54	0.019	0.095	17.20	0.0085		0.05	0.48	43
L2-03	10.50	0.82	0.080	0.0345	1.58	0.074	0.102	17.02	0.0065	-	0.26	0.50	32
L7-03	10.60	0.18	0.040	0.0384	1.02	0.007	0.111	15.40	0.0274	-	0.02	0.05	52
L23-03	12.04	0.68	0.030	0.0475	0.96	0.043	0.092	17.30	0.0093	Nb 1.06	0.06	0.41	22
L17-03	8.00	0.66	0.090	0.0089	0.48	0.061	0.078	15.30	0.0090	-	0.23	0.28	20
L14-03	7.93	1.49	0.080	0.0023	1.76	0.107	0.028	15.00	0.0045	-	1.64	2.01	0
L27-01	10.30	0.96	0.040	0.0018	0.97	0.057	0.019	15.30	0.0058	Mo 2.01	0.73	1.02	2
L15-03	8.00	1.82	0.010	0.0129	1.07	0.020	0.085	17.80	0.0110	-	0.78	1.17	32
L6-03	10.00	1.90	0.020	0.0051	1.13	0.096	0.087	17.10	0.0058	-	2.02	2.44	36
L24-03	12.30	0.03	0.007	0.0055	0.48	0.031	0.002	16.90	0.0129	Nb 1.72	0.11	0.86	0

Table 14. Composition (wt.%) of stainless steels irradiated to  $\approx 2.0 \times 10^{21} \text{ n cm}^{-2}$  (E > 1 MeV) ( $\approx 3.0 \text{ dpa}$ ) correlated with results of SSRT test and SEM fractography.

<sup>a</sup>L5-03 contains globular-shaped  $\approx$ 3 vol.% delta ferrite.

Examples of load-elongation behavior are shown in Fig. 18; they were obtained on SSRT specimens fabricated from a commercial heat of Type 304 SS (Heat C19) and tested in water at 289°C and in air at 23, 289, and 325°C. Effects of neutron damage levels of 0, 0.4, 1.3, and 3 dpa are also shown in the figure. Several observations can be inferred from the results: (a) the increase in yield strength from 1.3 dpa to 3.0 dpa is small, indicating that irradiation hardening reaches near saturation at  $\approx 2$  dpa; (b) by the time the damage level reaches  $\approx 3$  dpa, work hardening capability of the steel is insignificant in water, and strain-softening occurs to some extent in air at 289-325°C; (c) at  $\approx 3$  dpa, tensile behavior in 289°C and 325°C air is similar; (d) ductility, i.e., uniform and total elongations in 1.3-dpa specimens tested in air, is lower at 289°C than at 23°C. The mechanism that leads to the last behavior is not known. However, the degree of strain aging or localized deformation via dislocation channeling or twinning could be more pronounced at the higher temperatures than at 23°C.

#### 8.2 Correlation of Percent IGSCC with Elemental Composition

From the results summarized in Table 14, percent IGSCC obtained on 304- and 316-type steels was plotted as a function of bulk concentration of Cr, Ni, Mn, Si, P, N, O, C, and S to see if a good correlation can be obtained with any of the elements. Such plots are shown in Figs. 19-27, respectively. In these plots, the results obtained on the 348-type steels (Alloys L23 and L24) and the austenitic-ferritic steel (i.e., Alloy L5 that contains  $\approx$ 3 vol.% delta ferrite) were not included.





A careful examination of the data in Fig. 19 shows that relative IASCC susceptibility (i.e., percent IGSCC from the SSRT tests in this study) cannot be correlated well with bulk Cr concentration. That is, for a given Cr content between 15 and 19 wt.%, percent IGSCC is independent of Cr content, although by the time the damage level reaches 3 dpa, grain-boundary Cr depletion should be significant in all the tested steels and should be near the saturated levels. Therefore, the results inFig. 19 do not support the premise that Cr atoms in bulk or on grain boundaries play a direct (primary) role in IASCC. In relation with this observation, it is interesting to note that the grain-boundary Cr-depletion mechanism is not supported by the results of postirradiation-annealing studies reported for BWR<sup>14-15</sup> and PWR<sup>10,16</sup> conditions. It has been shown in these studies that, during post-irradiation annealing at 400-550°C, IASCC susceptibility becomes negligible long before grain-boundary Cr concentration begins to change from its level in the as-irradiated state.

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Similar examination of Figs. 20-26 strongly indicates that Ni, Mn, Si, P, N, O, or C do not play a primary role.

Of the 19 alloys analyzed in Figs. 19-27, of particular interest is the behavior of seven 304- or 316-type alloys that were found to be resistant to IASCC, i.e., Alloys C1, C10, C12, C21, L14, L26, and L27. Understanding why these heats are resistant is considered to hold an important clue to understanding the mechanism of IASCC. From Figs. 19-26, it is evident that IASCC susceptibilities (i.e., percent IGSCC) of the seven alloys are zero or insignificant, regardless of the concentrations of Cr, Ni, Mn, Si, P, N, O, or C. The only exception is S (see Fig. 27).

The S and C content of the seven resistant heats is given in Figs. 19-26. The S and C content of C16, which is susceptible to IASCC, is also given. Sulfur content of all other heats not denoted in the figures is 0.003 wt.% or higher. It can be deduced from the figures that, whereas the IASCC-resistant behavior of the seven heats is independent of the concentration of the element of each figure, they possess a unique and common character. That is, S concentration is very low (<0.002 wt.%) and, at the same time, C concentration is relatively high (>0.05 wt.%).

Furthermore, Fig. 27 shows that all 304- or 316-type steels containing >0.002 wt.% S and tested in this investigation were susceptible, whereas steels containing  $\leq 0.002$  wt.% S were either susceptible or resistant. Therefore, the results in Figs. 19-27 strongly indicate that S concentration of  $\leq 0.002$  wt.% is a necessary but not a sufficient condition for a 304- or 316-type steel to be resistant to IASCC under BWR-like conditions.



Figure 19. Percent IGSCC at ≈3 dpa correlated with bulk Cr concentration, 304- and 316-type steels.



Figure 20. Percent IGSCC at ≈3 dpa correlated with bulk Ni concentration, 304- and 316-type steels.

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Figure 21. Percent IGSCC at ≈3 dpa correlated with bulk Mn concentration, 304- and 316-type steels.



Figure 22. Percent IGSCC at ≈3 dpa correlated with bulk Si concentration, 304- and 316-type steels.



Figure 23. Percent IGSCC at ≈3 dpa correlated with bulk P concentration, 304- and 316-type steels.



Figure 24. Percent IGSCC at ≈3 dpa correlated with bulk N concentration, 304- and 316-type steels.

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Figure 25. Percent IGSCC at ≈3 dpa correlated with bulk O concentration, 304- and 316-type steels.



Figure 26. Percent IGSCC at ≈3 dpa correlated with bulk C concentration, 304- and 316-type steels.

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In Fig. 26, the behavior of Heat C16, which contains a very low concentration of S (0.002 wt.%) and a relatively low concentration of C (0.02 wt.%), is compared with the behavior of other similar heats that contain  $\leq 0.002$  wt.% S but higher concentrations of C. In spite of the very low concentration of S, the heat was susceptible. This finding indicates that C concentration higher than a threshold value (somewhere between 0.02 and 0.057 wt.%) is also a necessary (but not a sufficient) condition for a 304- or 316-type steel to be resistant to IASCC.

However, a combined condition that S concentration be less than  $\approx 0.002$  wt.% and, at the same time, that C concentration be higher than a threshold value (e.g., >0.03 wt.%) appears to be a sufficient condition for a 304- or 316-type steel to be resistant to IASCC. This observation strongly indicates that S is the primary contributor to IASCC susceptibility; therefore, the effect of S and relevant data were analyzed more systematically.

#### 8.3 Effect of Fluence

Figure 28 shows the effect of fluence on the IASCC-resistant behavior of the seven heats of 304- or 316-type SSs that contain  $\leq 0.002 \text{ wt.}\%$  S and > 0.05 wt.% C. For damage levels of 0.4, 1.3, and 3.0 dpa, no appreciable susceptibility to IASCC was observed in these heats in 289°C BWR-like high-purity water that contains  $\approx 8 \text{ ppm DO}$  (ECP  $\approx 160\text{-}220 \text{ mV}$  SHE). The IASCC-resistant behavior of these heats is expected to be valid also in similar water that contains lower levels of DO and ECP.

In contrast, a strong effect of fluence was observed for two groups of 304- or 316-type steels that are susceptible to IASCC, i.e., heats that contain high S ( $\geq 0.003 \text{ wt.\%}$ ) and heats that contain low S (<0.002 wt.%) and low C (<0.03 wt.%). The effect of fluence for the 16 susceptible heats of 304- or 316-type SSs tested in this study is shown in Fig. 29. The effect of fluence of the susceptible heats varied significantly from heat to heat. However, the root cause of the strong variation in fluence effect is not understood. Strong variation is evident for both laboratory and commercial heats.

#### 8.4 Role of S in IASCC at High Fluence

In contrast to the laboratory-fabricated model SSs that have compositions outside of the AISI specifications, for all eight commercial heats of Type 304, 304L, 316, and 316L austenitic SSs investigated in this study, bulk S content provided a good monotonic correlation with percent IGSCC (see Fig. 30). Four commercial heats of Type 304 and 316 SS that contain very low concentrations of S ( $\leq 0.002$  wt.%) exhibited only negligible susceptibility to IGSCC.



Figure 28. Effect of neutron damage on IASCC susceptibility (percent IGSCC) of Type 304 or 316 steels that contain low concentrations of S and high concentrations of C (i.e., <0.0023 wt.% S and >0.05 wt.% C).



Figure 29. Effect of neutron damage on IASCC susceptibility (percent IGSCC) of Type 304 or 316 steels that contain low S and low C concentrations (i.e., ≤0.002 wt.% S and <0.03 wt.% C), high S and high C concentrations (i.e., >0.003 wt.% S and >0.05 wt.% C), or high S and low C concentrations (i.e., >0.003 wt.% S and <0.03 wt.% C).</p>

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However, as S content increased to  $\geq 0.003$  wt.%, susceptibility increased drastically. At an S content of >0.005 wt.%, the specimens from the commercial heats failed in virtually complete IG mode, just as in IASCC of field components. The steep increase of susceptibility at S levels >0.003 wt.% indicates that an S-related critical phenomenon plays an important role in IASCC.



Figure 30. Effect of bulk S content on IASCC susceptibility (percent IGSCC) of commercial heats of Types 304, 304L, 316, and 316L steels.

The deleterious effect of S can be discerned more directly by comparing the IASCC susceptibilities of a pair of high- and low-S heats of the same grade that contain otherwise similar elemental compositions, e.g., Heat C12 (S 0.002 wt.%) and Heat C9 (S 0.013 wt.%). As fluence increased, the susceptibility of Heat C12 remained negligible, whereas the susceptibility of Heat C9 increased drastically (see Fig. 31).

The relative behavior of Heats C9 and C12 is consistent with the observation reported by Tsukada et al. on their HP and HP+S heats.<sup>24,29</sup> They reported that an HP heat that contained 0.0318 wt.% S and 0.002 wt.% C (Heat HP+S, see Fig. 32) was significantly more susceptible than an HP heat that contained 0.0014 wt.% S and 0.003 wt.% C (Heat HP). They also reported that the latter heat was more susceptible than a C-doped HP heat that contains 0.0020 wt.% S and 0.098 wt.% C (Heat HP+C). A deleterious effect of S is also indicated from the expanding-tube-test data reported by Kasahara et al.,<sup>28</sup> Jacobs et al.,<sup>21</sup> and Garzarolli et al.<sup>2</sup> (see Figs. 33-35, respectively).

The tests used to obtain the data in Figs. 31 and 33-35 are based on the "binary-like" approach. In this approach, S content was varied high or low while the composition of other elements was kept similar to that of commercial steels used in reactor components. In contrast, the result in Fig. 32 is based on the "single-element" approach in which high-purity heats are used.



Figure 31. IASCC susceptibility cf low-S (Heat C12) and high-S (Heat C19) heats of comparable chemical composition Type 304 SS.

Figure 32. IASCC susceptibility of high-purity heats of Type 304L SS doped with C or S, from Tsukada et al., Ref. 29.



Although the latter approach is advantageous when used to isolate more clearly the effect of a single element, it is difficult to determine which element plays a dominant or secondary role and to determine if the

effect of that particular element is influenced by other elements present in a steel similar to those used in a reactor. Therefore, a finding from such an approach must be confirmed for the type of steels used to fabricate actual field components. Tests on a sufficiently large number of commercial heats, such as those in Fig. 30, serve this purpose. Also, any potential effect of uncommon parameters, such as the presence of ferrite or dendrites, should be screened carefully. The deleterious effect of S indicated from the three approaches is consistent, i.e., the binary-like single-element approach with commercial steels (Figs. 31 and 33-35), the single-element approach with HP steels (Fig. 32), and the multiple-elements approach with commercial steels (Fig. 30).





#### 8.5 Effect of Carbon on Deleterious Role of Sulfur

For high-carbon heats of Types 304 and 316 SS, it was shown in Fig. 28 that S content of  $\leq 0.002$  wt.% provides good resistance to IASCC. However, whether the same is true or not with low-carbon heats of Types 304L, 316L, or high-purity SSs needs careful examination.

In this investigation, Type 316 SS Heat C21 that contains 0.001 wt.% S and 0.060 wt.% C was resistant to IASCC. In contrast, a similar heat of Type 316L SS (Heat C16) that contains 0.002 wt.% S and 0.020 wt.% C was susceptible (Fig. 30). This behavior appears to be consistent with the data of Tsukada et al.,<sup>29</sup> i.e., the behavior of Heats HP+C vs. HP shown in Fig. 32. Tanaka and coworkers performed SSRT tests on 304L- and 316L-type SS heats that contained S in nominal concentrations between 0.000 and 0.003 wt.% after irradiation in a BWR to  $\approx 1.9$  to  $2.1 \times 10^{21} \text{ n} \cdot \text{cm}^{-2}$  (E > 1 MeV).<sup>41</sup> A 304-type heat that contained 0.002 wt.% S and 0.060 wt.% C was resistant to IASCC at  $1.3 \times 10^{21} \text{ n} \cdot \text{cm}^{-2}$  (E > 1 MeV), whereas a 304Ltype heat that contained 0.002 wt.% S and 0.013 wt.% C was susceptible to IASCC at  $2.1 \times 10^{21} \text{ n} \cdot \text{cm}^{-2}$ . Of the nine 316L-type heats investigated by Tanaka et al., three exhibited negligible susceptibility to IASCC, whereas the other six exhibited susceptibility. Fukuya et al. reported similar results.<sup>20</sup> Summarizing the results from the four investigations, Figs 36 and 37 show, respectively, relative IASCC susceptibilities (i.e., percent IGSCC from SSRT test) of low-S heats (S  $\leq 0.002$  wt.%) of high-C steels (i.e., Types 304 and 316 SS) and those of their low-C counterparts. The trend in the figures indicates that low-S heats (S  $\leq 0.002$  wt.%) of 304- and 316-type SS are resistant to IASCC. However, all low-S heats of 304L- and 316L-type SS that contain  $\leq 0.002$  wt.% S are not necessarily resistant.

A similar trend is also evident for 348- and 348L-type SSs investigated by Garzarolli and coworkers by swelling-tube tests<sup>2,7,42</sup> in a PWR (see Figs. 38 and 39). In their experiments, average diametral strain was measured for intact and cracked tubes from the same heat after irradiation to one to three fuel cycles. Sulfur concentration in the tubes ranged from 0.002 to 0.007 wt.%. Their one-cycle data in Fig. 38 show that, under the same irradiation conditions, high-C heats (C 0.027-0.043 wt.%) could tolerate more diametral expansion without incurring a failure than low-C heats (C 0.003-0.015 wt.%), in spite of the fact that the high-C heats contained somewhat higher levels of S than the latter (i.e., 0.005-0.007 wt.% vs. 0.003-0.004 wt.%). A similar trend is observed from the one-to-three cycle data shown in Fig. 39. That is, high-C heats (C 0.027-0.048 wt.%) could withstand more diametral expansion without a failure than low-C heats (C 0.003-0.015 wt.%), in spite of S than the latter and that fluence was higher in the former than in the latter.

The consistent behavior from the six investigations summarized in Figs. 36-39 shows that high C content suppresses the deleterious effect of S, if the S concentration is very low. In 304- or 316-type SSs, the beneficial effect of C (in concentrations  $\geq 0.03$  wt.%) is manifested significantly if S concentration is less than  $\approx 0.004$  wt.%. In the absence of such a beneficial effect, 304L- or 316L-type steels seem to be inherently less resistant than their high-C counterparts; this seems to lead to significant susceptibility even in some low-S heats, e.g., those shown in Figs. 36 and 37.




Figure 37. Percent IGSCC from SSRT test of low-S high-C and low-S low-C heats of Type 316 or 316L SSs.







Figure 39. Diametral strain at failure for low-S high-C and low-S low-C heats of Type 348 SSs. Data from Garzarolli et al. Refs. 2, 7, and 42, one-to-three-cycle irradiation.

When S concentration is high, e.g.,  $\geq 0.004$  wt.% in 304- or 316-type SSs and  $\geq 0.01$  wt.% in 348-type SSs, it appears that the deleterious effect of S is so dominant that the effect of C is negligible or insignificant, even at high concentrations of C.

#### 8.6 Effect of Si at High Fluence

As pointed out in Section 7, for a medium level of damage ( $\approx 1.3$  dpa), the data in Fig. 17 indicate that Si in high concentrations either delays the onset of IGSCC or suppresses the susceptibility to IGSCC. However, by the time the damage level reaches  $\approx 3$  dpa, it appears that the deleterious effect of S is so dominant that the effect of Si is either insignificant or secondary at best. This conclusion can be deduced from Fig. 22, in which percent IGSCC is plotted as a function of Si content.

Three 304-type laboratory heats that contained unusually high concentrations of Si were tested in 289°C water in this investigation after irradiation to  $\approx$ 3 dpa, i.e., Heats L14 (Si 1.49 wt.%), L15 (Si 1.82), and L6 (Si 1.90). These Si concentrations well exceed the AISI-specified limit of 1.0 wt.%. As shown in Table 14, the three heats exhibited relatively higher level of uniform elongation compared to other laboratory heats of 304- or 316-type SSs. However, in spite of this, the susceptibility of the three heats appeared to be determined largely by the concentrations of S and C. This is shown in Fig. 40. At  $\approx$ 3 dpa, there was no clear evidence that high Si content suppresses the susceptibility to IASCC of the steels investigated in this study.





Some of the low-C, low-S heats of 316L-type SSs (damage level  $\approx$ 3 dpa), shown in Fig. 37, contain an unusually low Si concentration of  $\leq 0.05$  wt.%.<sup>19,41</sup> When compared with the other heats in the figure that contain higher levels of Si (0.38-0.59 wt.%), the IASCC behavior of the high- and low-Si heats was similar; this is shown in Fig. 41.

As in the case of 304- and 316-type SSs, IASCC susceptibility of 348-type SS at high fluence ( $\approx$ 3 dpa) seems to be determined largely by the concentrations of S and C, and the effect of Si appears secondary.





## 9 Representation of IASCC-Resistant or –Susceptible Behavior of 304and 316-Type Steels in Sulfur-Carbon Map

Typical in-service cracking of BWR core internal components occurs at >2 dpa and is characterized by a virtually full intergranular fracture surface, sometimes accompanied by a small amount of transgranular fracture surface. In this respect, the IASCC behavior of SSs at  $\approx 3$  dpa (Section 8) is more relevant than that at  $\approx 1.3$  dpa (Section 7). As described in Section 8, IASCC-resistant or -susceptible behavior at  $\approx 3$  dpa is largely determined by the bulk concentrations of S and C. Similar analysis was performed for data reported in the literature. Virtually all literature data have been reported after 1987. Figures 42 and 43 show a twodimensional map of bulk S and C contents to show the range in which 304- or 316-type steels are either resistant or susceptible to IASCC. The figure also includes similar data reported in the literature since 1987. The only data missing from the figure are those obtained for BWR internals for which S and C contents are not available, e.g., an IASCC-resistant 304 SS control blade sheath reported in Ref. 23 and an IASCCsusceptible 304L SS dry tube reported by Shen and Chang.<sup>43</sup>



Bulk Sulfur Content (wt.%)

Figure 42. Range of bulk S and C contents in which Type 304 or 316 steels are resistant or susceptible to IASCC in BWR-like oxidizing water.

In constructing the maps in Figs. 42 or 43, it was assumed that a heat susceptible at <2.5 dpa is equally or more susceptible at  $\geq 3$  dpa under otherwise similar conditions. It was also assumed that a heat susceptible at a DO concentration of <0.4 ppm is more susceptible at a DO concentration of  $\geq 8$  ppm under otherwise identical conditions. Note that, in the figure, the plotted ranges of S and C contents extend beyond the AISI limits specified for Types 304 and 316 SS, i.e., maximum S, 0.030 wt.% and maximum C, 0.08 wt.%.



Figure 43. Expanded view of low-S portion of Fig. 42 showing data in detail.

In Figs. 42 and 43, data obtained on neutron-, ion-, or proton-irradiated steels from SSRT, crackgrowth-rate (CGR), or tube-expansion tests are shown for comparison with those obtained by SSRT in this investigation. As shown in Fig. 43, the results obtained on He-ion-irradiated steels (Fukuya et al.<sup>3</sup>) and proton-irradiated steels (Cookson et al.<sup>44</sup> and Busby et al.<sup>45</sup>) are also consistent with the IASCC-susceptible range shown in the S-C map. The earlier efforts by the latter group of investigators focused on "ultra-highpurity" (UHP) heats. However, their "UHP" heats contained either insufficiently low concentrations of S, insufficiently high concentrations of C, or both, to render the steel resistant to IASCC. Figure 43 also includes the results of CGR tests reported by Karlsen and Hauso<sup>46</sup> and Jenssen et al.,<sup>12</sup> in which intergranular crack propagation was confirmed by SEM fractography.

In Fig. 44, which is similar to Fig. 42, data points for four susceptible heats tested after irradiation to 1.3 dpa (i.e., Heats L22, L11, L13, and L8; see Fig. 29) are also included. It was assumed that the four heats are also susceptible to IASCC at 3 dpa, although they were not tested at this damage level.



Figure 44. Range of bulk S and C contents in which Type 304 or 316 steels are resistant or susceptible to IASCC, including four susceptible heats tested at 1.3 dpa in this study.

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## 10 IASCC Behavior of Type 348 Steels

Besides the 304- and 316-type steels, two heats of 348-type SS were tested in 289°C water in this investigation after irradiation to  $\approx$ 1.3 and  $\approx$ 3 dpa. One heat (L23) contains a very high concentration of S (0.0475 wt.%) and the other (L24), a low concentration of S (0.0055 wt.%). Percent IGSCC of the two heats is compared in Fig. 45. The high-S heat was susceptible to IASCC, but the low-S heat was resistant at both levels of neutron damage. This behavior is consistent with the behavior of 304- or 316-type SSs described in Section 8.

Similar to the correlation established for 304- and 316-type SSs in Fig. 42, the IASCC-resistant range of 348-type and an Nb-containing SS was plotted in a two-dimensional S-C map in Fig. 46. This map includes all such data in the literature that were obtained under BWR- and PWR-like conditions. As indicated in the figure, it appears that the IASCC-resistant range of 348-type and Nb-doped SSs in PWR- or BWR-like water can be described well in an S-C map that differs somewhat from S-C map of 304- and 316-type SSs. Behavior of the Nb-doped 316-type steel was reported by Jacobs et al. for BWR conditions.<sup>21</sup> Although the correlation in Fig. 46 should be considered only preliminary, it is qualitatively consistent with the correlation constructed for 304- and 316-type SS for BWR-like conditions. It appears that 348-type SSs exhibit higher tolerance to S than 304- or 316-type steels.



Figure 45. Percent IGSCC of high- and low-sulfur laboratory heats of Type 348 SS after irradiation to 1.3 and 3 dpa, SSRT test in BWR-like water.

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Figure 46. Range of bulk S and C contents in which Type 348 stainless steels are resistant or susceptible to IASCC.

#### 11 IASCC Behavior of Austenitic-Ferritic Steel

In this investigation, one austenitic-ferritic duplex SS alloy was tested by SSRT in 289°C water after irradiation to 0.4, 1.3, and 3.0 dpa. The alloy (L5), which contains 21 wt.% Cr and  $\approx$ 3 vol.% delta ferrite, was resistant to IASCC, even though its S content was very high (i.e., 0.028 wt.%). Figure 47 shows an insignificant level of percent IGSCC measured on the alloy at all the three damage levels.



The morphology of the delta ferrite in the alloy was globular, as shown in Fig. 48. Annealing twins are also visible in the optical metallograph. The fracture surface of the alloy at 3 dpa showed small broken islands of delta ferrite surrounded by dominantly ductile regions (see Fig. 49).

The excellent resistance of this alloy to IASCC appears to be consistent with the behavior of cast austenitic SSs of Types CF-3 and CF-8, which, in the unirradiated state, are well known for their resistance to IGSCC. Most CF-3 and CF-8 duplex SSs contain a delta ferrite content of >10 vol.%, usually in the form of acicular or elongated grains. If the volume fraction of delta ferrite is too high, or if the distribution of ferrite is continuous, significant embrittlement due to ferrite cleavage may lead to unacceptable degradation of fracture toughness after irradiation to high fluence. However, with a ferrite content of only  $\approx3\%$ , as in Alloy L5, small globular ferrite islands remain isolated, and overall material embrittlement due to ferrite cleavage appears insignificant. Supporting this premise, yield strength and total elongation of the alloy in 289°C water were significant even after irradiation.

It appears that the mechanism through which a small amount of delta ferrite may suppress the susceptibility to IASCC in Alloy L5 can be explained well by the effect of delta ferrite on S distribution in the steel. As shown in the equilibrium Fe-S diagram in Fig. 50, the solubility limit of S is several times higher in the  $\delta$  ferrite phase than in the austenitic ( $\gamma$ ) phase. At  $\approx$ 1365°C, the solubility limits of S in delta ferrite and austenite are 0.18 and 0.05 wt.%, respectively. Therefore, during ingot melting, solidification, and cooling of

a material such as L5, S atoms are expected to migrate toward and be incorporated into the delta ferrite globules, which act as efficient trapping sites for S atoms. As a result, segregation of S atoms to austenite grain boundaries during fabrication or irradiation will be suppressed because of the presence of ferrite globules and the large austenite-ferrite phase boundaries.

Also, continuous austenite grain boundaries are rare in this type of steel, because ferrite grains intersect austenite grain boundaries. As a consequence, continuous crack propagation along austenite grain boundaries seems to be more difficult, and, as a consequence, IASCC susceptibility seems to be suppressed significantly.



Figure 48. Optical metallograph of IASCC-resistant Alloy L5 that contains ≈3 vol.% delta ferrite.



Figure 49. Predominantly ductile fracture surface morphology of IASCCresistant Alloy L5 that contains ≈3 vol.% delta ferrite, 3 dpa.

Figure 50. Fe-rich side of Fe-S phase diagram; from Hansen, Ref. 47.

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### 12 Fractographic Analysis

Fractographs produced in 289°C water were obtained in a shielded SEM for all 3-dpa specimens listed in Table 14. Fractographs were also obtained for specimens that were irradiated to 0.4 and 1.3 dpa; some of these specimens exhibited significant susceptibility to IASCC. Typically, 10-15 fractographs were obtained for each specimen at various magnifications ranging from 20X to 800X. For each specimen, a composite fractograph was then constructed for the whole fracture surface at a magnification ranging from 100X to 150X. This section summarizes results of the fractographic analysis for 3-dpa specimens.

# 12.1 Fracture Surface Morphology of IASCC-Resistant Heats of 304, 316, and 348 Type Steels at 3 dpa

Typical fracture surface morphologies of the IASCC-resistant seven heats of 304- or 316-type SSs (see Fig. 28) and one heat of 348 SS (see Fig. 45) are shown in Figs. 51-58. All of the eight heats, which were irradiated to 3 dpa, exhibited predominantly ductile fracture surface morphology, sometimes mixed with a limited amount of transgranular fracture. Percent intergranular fracture was negligible in the eight heats (i.e.,  $\leq 2\%$ ).





Transgranular fracture morphology was often observed near the specimen free surfaces, which were covered with thin oxide layers. In the IASCC-resistant heats, a transition from transgranular morphology to ductile morphology was often observed not far from the specimen free surface. Such examples are visible in Figs. 51, 56, and 57. The fracture surface morphology of Laboratory Heat L26 (Fig. 57) shows

predominantly ductile morphology and a limited amount of transgranular morphology that surrounds an insignificant amount of intergranular morphology of  $\approx 2\%$ .



Figure 52. Dominantly ductile fracture surface morphology of IASCCresistant Commercial Heat C10 of Type 304 SS; 3 dpa.

Figure 53. Dominantly ductile fracture surface morphology of IASCC-resistant Commercial Heat C12 of Type 304 SS; 3 dpa.



(a) (b) Figure 54. Fracture surface morphology of IASCC-resistant Laboratory Heat L14 of Type 304 SS (3 dpa), showing predominant ductile morphology (a) and limited transgranular morphology (b); intergranular morphology is negligible.



Figure 55. Dominantly ductile fracture surface morphology of IASCC-resistant Commercial Heat C21 of Type 316 SS; 3 dpa.



Figure 56. Fracture surface morphology of IASCC-resistant Laboratory Heat L24 of Type 348 SS (3 dpa); both low- (a) and high-magnification (b) images show that ductile morphology is dominant, and intergranular morphology is negligible.



Figure 57. Fracture surface morphology of IASCC-resistant Laboratory Heat L26 of Type 304 SS (3 dpa), showing (a) limited amount of transgranular morphology that surrounds insignificant amount of intergranular morphology (≈2% IGSCC) and (b) dominantly ductile morphology.



Figure 58. Ductile fracture morphology of IASCC-resistant Laboratory Heat L27 (Type 316 SS, 3 dpa); both low- (a) and high-magnification (b) images show that intergranular morphology is negligible.

# 12.2 Fracture Surface Morphology of IASCC-Susceptible Heats of 304, 316, and 348 Type Steels at 3 dpa

Typical fracture surface morphologies of the 12 IASCC-susceptible heats of 304- or 316-type SSs (3 dpa) are shown in Figs. 59-70. Similar morphology for one susceptible 348 SS heat is shown in Fig. 71. All of the 13 heats exhibited a significant amount of intergranular fracture surface morphology, sometimes close to 100%. From the results of fractographic analysis, several characteristics were observed:

- (1) Most of the intergranular fracture surfaces show numerous secondary cracks along the grain boundaries. These cracks branch at sharp angles from the plane of the main crack propagation.
- (2) In some cases, virtually whole grains appear to be encircled by intergranular separation, e.g., Figs. 59, 60, 63, and 64. This phenomenon has been commonly referred to as "grain encirclement" by some investigators. Sometimes, such a grain "drops out," like a grain of sand, during metallographic polishing.
- Intergranular fracture surfaces are often covered with an unknown type of corrosion debris, e.g., Figs. 59, 60, 62, 63, 66, 67, 68, 70, and 71. The shape of the debris is mostly spherical, but in some case, it is tetrahedral, e.g., Fig. 62.
- (4) Little corrosion debris is observed on ductile or transgranular fracture surfaces.



## Figure 59. Intergranular fracture surface morphology of IASCC-susceptible Heat C16 (316L SS, 3 dpa). Note secondary cracks along grain boundaries at high angles to plane of main crack propagation.

Figure 60. Intergranular fracture surface morphology of IASCC-susceptible Heat C19 (304 SS, 3 dpa). Note corrosion debris and secondary cracks along grain boundaries at high angles to plane of main crack propagation.



Figure 61. Intergranular fracture surface morphology of IASCC-susceptible Heat L4 (304 SS, 3 dpa). Note secondary cracks along grain boundaries at high angles to plane of main crack propagation.



Figure 62. Intergranular fracture surface morphology of IASCC-susceptible Heat C9 (304 SS, 3 dpa). Note corrosion debris and secondary cracks along grain boundaries at high angles to plane of main crack propagation.



Figure 63. Intergranular fracture surface morphology of IASCC-susceptible Heat L18 (304 SS, 3 dpa). Note corrosion debris and secondary cracks along grain boundaries at high angles to plane of main crack propagation.



Figure 64. Intergranular fracture surface morphology of IASCCsusceptible Heat C3 (304L SS, 3 dpa). Note secondary cracks along grain boundaries.



Figure 65. Intergranular and ductile fracture surface morphologies of IASCC-susceptible Heat L25 (304L SS, 3 dpa). Note secondary cracks along grain boundaries.

Figure 66. Intergranular fracture surface morphology of IASCC-susceptible Heat L2 (304 SS, 3 dpa). Note corrosion debris and deformation steps on grain boundaries.



Figure 67. Intergranular fracture surface morphology of IASCC-susceptible Heat L7 (304L SS, 3 dpa). Note corrosion debris and high-angle secondary cracks.

Figure 68. Intergranular fracture surface morphology of IASCC-susceptible Heat L17 (304 SS, 3 dpa). Note corrosion debris and high-angle secondary cracks.



Figure 69. Intergranular and ductile fracture surface morphologies of IASCC-susceptible Heat L15 (304L SS, 3 dpa).

Figure 70. Intergranular fracture surface morphology of IASCC-susceptible Heat L6 (304 SS, 3 dpa). Note corrosion debris and high-angle secondary cracks.



Figure 71. Intergranular fracture surface morphology of IASCC-susceptible Heat L23 (348 SS, 3 dpa). Note corrosion debris and high-angle secondary cracks.

## 13. Analysis of Grain-Boundary Microchemistry by Auger Electron Spectroscopy

In the consideration of the importance of the roles of S and C in IASCC described in Sections 7-9, the behavior of grain-boundary segregation of the two elements was characterized by Auger electron spectroscopy (AES). Similar analysis by advanced analytical electron microscopy (AAEM) is difficul: or impractical for an ordinary grain boundary in an irradiated steel, although S-rich thin films on a crack-tip grain boundary can be detected.<sup>33,34</sup> Auger electron spectroscopy is probably the only practical technique that can characterize well the profile and degree of grain-boundary segregation of S and C in an irradiated steel.

The materials selected for the AES analysis were taken from BWR neutron absorber tubes. The absorber tubes were fabricated from two commercial heats of Type 304 SS (Heats A and B) and were irradiated to  $2.0 \times 10^{21}$  n cm<sup>-2</sup> (E > 1 MeV) ( $\approx$ 3 dpa) during service in a BWR. The behavior of grainboundary segregation in such material was considered most relevant to BWR IASCC. The neutron absorber tubes: were susceptible to field cracking. Unfortunately, neither documented chemical composition of the absorber tubes nor the archive, unirradiated ingots could be found.

Needle-like specimens were prepared from the selected BWR neutron absorber tubes. After cathodically charging with hydrogen at  $\approx 50^{\circ}$ C in a solution that contained 100-mg/L NaAsO<sub>2</sub> dissolved in 0.1-N H<sub>2</sub>SO<sub>4</sub>, the needle-like specimen was fractured in a 23°C vacuum in a scanning Auger electron microscope (SAM). In-situ analysis was performed in a JEOL JAMP-10 SAM equipped with automated Arsputtering and depth-profiling devices. Four to six ductile fracture surfaces and 12-15 intergranular (IG) fracture surface locations were selected for spot analysis in each specimen. Sulfur and C signals from the selected surfaces were then recorded and compared.

After each spot analysis, a depth-profile analysis was performed on a selected IG fracture surface. The procedure for automated depth-profiling analysis has been described elsewhere.<sup>18,19</sup> Progressive sputtering of the intergranular surfaces with Ar ions and acquisition of the Auger electron signals in the ultra-high vacuum environment of the SAM were controlled by computer.

#### 13.1 Grain-Boundary Segregation of Sulfur and Carbon

An example of an IG fracture surface, produced at 23°C in the H-charged specimens from an absorber tube (304 SS Heat B), is shown in Fig. 72. The IG fracture surfaces reveal numerous deformation steps and a precipitate (diameter  $\approx 15 \,\mu$ m) denoted with number "6". Spots denoted with numbers "5" and "20" were the spots selected for acquisition of Auger electron signals. The results of spot analyses obtained for Tubes A-1 and E are shown in Figs. 73 and 74, respectively. In Fig. 73(a), the peak heights of Auger electrons of S obtained from four ductile fracture surfaces are presented with their counterparts from 12 IG fracture surfaces in Tube A-1. The S intensities from the IG fracture surfaces are significantly higher than their counterparts from ductile-fracture surfaces (i.e., spots not on a grain boundary). This behavior shows that S segregated significantly to grain boundaries (i.e., IG fracture surface) in Tube A-1. Similar results for C are also plotted in Fig. 73(b), which shows that C segregated significantly to grain boundaries in the same tube. Essentially the same behavior was observed for S and C segregation in Tube B; see Fig. 74.

A careful examination of Figs. 73 and 74 reveals that some spots that contain high concentrations of S also contain high concentration of C, e.g., intergranular Spot 16, Tube A-1 and intergranular Spot 12, Tube B. However, whether C segregates preferentially to a grain-boundary region that contains a high concentration of S, or vice versa is not conclusive. This behavior could not be conclusively determined by AES in this investigation; it needs further study by a more direct technique such as field-emission atom-probe technique.



Figure 72. High-magnification fractograph of Hcharged BWR neutron absorber tube, 304 SS, Heat B, ≈3 dpa; fracture was produced at 23°C in ultra-high vacuum environment in a scanning Auger electron microscope.

Figure 75 shows the results of depth-profile analysis for the two tubes fabricated from 304 SS Heats A-1 and B. The figure confirms significant grain-boundary segregation of S in Tubes A-1 and B and C segregation in Tube B.



e 73. Summary of AES analysis of grain-boundary segregation of S (a) and C (b) in BWR neutron absorber tube fabricated from Type 304 SS Heat A and irradiated to 3 dpa: S and C peak heights from ductile (denoted "D") and IG (denoted "I") fracture surfaces are compared. The horizontal line denotes the average signal strength on a ductile fracture surface.



Figure 74. Summary of AES analysis of grain-boundary segregation of S (a) and C (b) in BWR neutron absorber tube fabricated from Type 304 SS Heat B and irradiated to 3 dpa: S and C peak heights from ductile (denoted "D") and IG (denoted "I") fracture surfaces are compared. The horizontal line denotes the average signal strength on a ductile fracture surface.



Figure 75. Result of AES depth-profiling analysis of grain-boundary segregation of S and C in BWR neutron absorber tubes fabricated from Type 304 SS Heats A and B and irradiated to ≈3 dpa.

The results in Figs. 73-75, however, do not clarify whether the grain-boundary segregation of S or C occurred via a thermal process (during fabrication), an irradiation-induced process, or both. The former type of segregation is expected to be significantly influenced by several steps during fabrication, e.g., ingot cooling, inter-rolling annealing and cooling, final annealing, and quenching. Thermal segregation is also influenced by plate thickness.

Results of AES analysis of grain-boundary segregation of S have been reported by Jacobs et al.<sup>48</sup> for fuel-cladding-like tubes that were irradiated in a BWR. A tube fabricated from a commercial heat of Type 348 SS was irradiated to  $1.5 \times 1021 \text{ n} \cdot \text{cm}^{-2}$  (E > 1 MeV), and a tube fabricated from a high-purity heat of Type 348 SS was irradiated to  $3.4 \times 10^{21} \text{ n} \cdot \text{cm}^{-2}$  (E > 1 MeV). Results of spot analysis similar to that of Figs. 73 and 74 showed significant grain-boundary segregation of S in both tubes. Depth-profiling analysis similar to that of Fig. 75 was, however, not reported from their investigation.

Thermal segregation of S to grain boundaries for unirradiated ultra-high-purity (UHP) austenitic SS doped with S has been reported by Andresen and Briant.<sup>49</sup> They concluded that S plays an important role in producing the IG crack path in thermally sensitized non-irradiated steels. Because S atoms are thermally segregated on grain boundaries, more S ions are released into water from the grain boundary than from the grain matrix, and it was thought that an IG crack path is promoted in steels that contain high concentrations of S. Thus, the authors essentially viewed the role of S as accelerating corrosive attack (i.e., dissolution of grain

boundary metal) of a Cr-depleted grain boundary, exacerbated by S ions released into the crack tip water. According to this model, as long as grain boundaries are significantly depleted of Cr, the production of an IGSCC path is predicted, even in a steel free of S.

However, a similar model of the role of S does not appear to explain our results on IASCC susceptibility of irradiated steels. One difficulty is how to explain the observation that IASCC susceptibility becomes negligible when S concentration is very low even though Cr depletion is significant at  $\approx 3$  dpa (see Figs. 28 and 43). Another difficulty is how to explain the trend that the S effect is strongly influenced by fluence at >0.003 wt.% S, e.g., in 304- and 316-type steels (see Fig. 29).

#### 13.2 Analysis of Solubility of Carbon in Sulfides

The results in Figs. 42-44 indicate that, when S concentration in a 304- or 316-type steel is  $\leq 0.002$  wt.%, a high concentration of C (>0.03 wt.%) promotes a decrease of the susceptibility to IASCC. When S concentration exceeds  $\approx 0.003$  wt.%, such a "beneficial" effect of high C concentration appears to be insignificant or absent. This observation suggests that some kind of synergism occurs between S and C and that the effect of such synergism is significant only at sufficiently low concentrations of S (<0.002 wt.%).

With respect to the above observation, our attention was focused on the composition and properties of Ni- and S-rich thin films that were observed on grain boundaries under three situations: i.e., at crack-tip grain boundaries in an unirradiated thermally sensitized crack-growth specimen tested in sulfated BWR-like water (304 SS), reported by Dumbill;<sup>31</sup> at crack-tip grain boundaries in a field-failed BWR top guide and a PWR baffle bolt (304 SS), reported by Thomas and Bruemmer;<sup>33,34</sup> and at ordinary grain boundaries of S-doped Ni specimens that failed during SSRT tests in air at 23°C, reported by Heuer et al.<sup>50</sup> The latter investigators showed that S-doped Ni specimens lose their tensile strength when the grain-boundary concentration of S exceeds  $\approx 15$  at.%, which causes an IG failure. Then, it was thought in our study that, if C is dissolved in a Ni<sub>x</sub>S<sub>y</sub>-type film in a significant amount, the mechanical properties of such a film are likely to be altered significantly. Accordingly, available information and data were evaluated to determine if C is soluble in sulfides.

The AES signal from the precipitate denoted with number "6" in Fig. 72 is shown in Fig. 76. The AES signal indicates that the precipitate is a CuS-type precipitate that contains a significant amount of C. Similar data are shown in Figs. 77 and 78 for two more CuS-type precipitates that were present in the same specimen.



Figure 76. AES signal showing high concentrations of Cu, S, and C in CuStype precipitate denoted with number "6" in Fig. 72.

The only direct evidence of significant solubility of C in Ni sulfide could be observed from the data reported by Dumbill.<sup>31</sup> The data (Fig. 6c in his report), obtained by field-emission-gun advanced analytical electron microscopy (FEG-AAEM), show X-ray intensities of Ni, S, and C contained in a Ni- and S-rich thin film that was observed at an intergranular crack in a thermally sensitized, unirradiated Type 304 SS. The FEG-AAEM X-ray spectra strongly indicate that C can be dissolved in significant amounts in Ni- and S-rich thin films at  $\approx 290^{\circ}$ C. Unfortunately, similar data were not reported for the field-cracked (irradiated) steels examined by Thomas and Bruemmer.<sup>33,34</sup> However, irradiation is not expected to significantly decrease the solubility of C in the Ni- and S-rich thin films or islands that they observed.

Based on the information described above, it is predicted that C atoms compete with S atoms to become incorporated in the unit cell of NiS-type sulfides under BWR conditions. The degree of such C incorporation, which replaces part of the S atoms in the sulfide, is predicted to be more pronounced for a higher concentration of C at the grain boundary, i.e., for a high C content in the steel and for a higher degree of grain-boundary segregation of C.



Figure 77. AES signal showing a high concentration of C in CuS-type precipitate in BWR neutron absorber tube, 304 SS, Heat B,  $\approx$ 3 dpa.



### 14. Discussion and Model Development

#### **14.1 General Discussion**

The results of this investigation shows that, as neutron fluence increases to a high level (e.g.,  $2.0 \times 10^{21}$  n cm<sup>-2</sup> [E > 1 MeV], or 3 dpa), the deleterious effect of S on IASCC susceptibility is dominant over the effects of other elements. The IG fracture surface morphology observed in commercial heats in this study was similar to that of field-cracked BWR core internals. This study also shows that a sufficiently low concentration of S (<0.002 wt.%) is the primary material factor necessary to ensure good resistance to IASCC. Even at slightly higher S concentrations of >0.003 wt.%, Type 304, 304L, 316, and 316L steels were susceptible to IASCC at 3 dpa. This strong dependence on S concentration, shown in Fig. 30 as an example, suggests that some kind of threshold phenomenon related to S is involved in IASCC.

It is important to note that, for 304- and 316-type steels that contain a low concentration of C of <0.03 wt.% (e.g., Types 304L, 316L, and their high-purity counterparts), an S concentration of <0.002 wt.% is not a sufficient material condition to ensure good resistance to IASCC. This behavior is in distinct contrast to the behavior of 304- and 316-type steels that contain high C concentrations of >0.03 wt.% (e.g., Types 304 and 316-type steels that contain high C concentrations of >0.03 wt.% (e.g., Types 304 and 316 steels) and S concentration of <0.002 wt.%. The latter class of steels exhibited good resistance to IASCC. This contrasting behavior indicates that, for S concentration <0.002 wt.%, a high concentration of C plays a beneficial role. However, when S concentration is high, i.e., >0.002 wt.%, it seems that the deleterious effect of S is so pronounced that the effect of a high concentration of C is largely obscured.

At lower damage levels (i.e., <1 dpa), a beneficial effect of a high concentration of Si (i.e., 0.8-1.5 wt.%) was observed. At higher damage levels, however, such effect appears to be obscured by the deleterious effect of S. Therefore, from the standpoint of BWR IASCC, which usually occurs at damage levels >2.5 dpa, we believe that the typical variation in Si concentration in Types 304 and 316 SS (i.e., 0.4 to 1.0 wt.%) is not an important factor. It is not clear if, in addition to a sufficiently low concentration of S (i.e., <0.002 wt.%) and a sufficiently high concentration of C (i.e., >0.03 wt.%), an unusually high concentration of Si (e.g., >1.5 wt.%) provides an extra benefit when fluence is high (e.g., >2.5 dpa). The two heats that belong in this category, i.e., 304 SS Heat L14 (S  $\approx$ 0.002 wt.%, C 0.107 wt.%, and Si 0.96 wt.%) and 316 SS heat L27 (S  $\approx$ 0.002 wt.%, C 0.057 wt.%, and Si 1.82 wt.%) exhibited good resistance at 3 dpa. On the other hand, 304 SS Heat C19, which contains a slightly higher S concentration (i.e., 0.003 wt.%) and a relatively low concentration of Si (i.e., 0.45 wt.%), exhibited a significant susceptibility at 3 dpa despite a high concentration of C (i.e., 0.06 wt.%) (see Table 14). Also, 304 SS Heat L6, which contains 0.005 wt.% S and 0.096 wt.% C, exhibited a significant susceptibility at 3 dpa despite a high concentration of Si (i.e., 0.45 wt.%).

Most field-cracked LWR core internals and steels used in SSRT tests or to evaluate crack-growth behavior in the IASCC community contain S concentrations >0.002 wt.%. Therefore, it is not surprising to see that most of them were susceptible to IASCC in one way or another. For S concentrations >0.002 wt.%, the degree of IASCC susceptibility in this study varied significantly from heat to heat, and it was difficult to predict the degree quantitatively. This behavior, consistent with literature data, is not unexpected when one considers the fact that, for a given fluence level, several factors can influence the degree of S segregation to grain boundaries, e.g., annealing and cooling history during fabrication, Mn content, precipitation of sulfides such as MnS and CuS, and solubility of the sulfides in BWR-like water.

The observation that steels that contain a very low or negligible amount of S are resistant to IASCC provides an important clue to the mechanism of IASCC. To understand the mechanism, we believe it is important to consider the following:

- (1) The strong dependence of IASCC susceptibility on fluence
- (2) The strong dependence of IASCC susceptibility on S concentration and an indication that some kind of critical phenomenon involving S occurs
- (3) A high concentration of C (>0.03 wt.%) suppresses IASCC susceptibility at sufficiently low S concentrations (<0.002 wt.%) but not at S concentrations >0.003 wt.%.
- (4) Very low solubility of S in an austenitic SS at  $\approx 300^{\circ}$ C
- (5) Segregation of S to grain boundary
- (6) Parallel grain-boundary Cr depletion and Ni segregation
- (7) Strong influence of water chemistry on BWR IASCC
- (8) Strong effect of strain rate on percent IGSCC from SSRT tests

Considering the finding that the S effect is dominant at high fluence and the fact that grain-boundary Cr depletion is in some respect a mirror image of grain-boundary Ni segregation, we believe it is important to understand the peculiar properties of Ni-S thin film reported by Heuer et al., who investigated the effect of ion-implanted S on disorder-induced amorphization of Ni.<sup>50</sup> Those authors also investigated the effect of thermal segregation of S on grain boundaries on the mechanical properties of unirradiated binary Ni-S. In the latter investigation, they found that, as the grain boundary concentration of S exceeds  $\approx 10$  at.% ( $\approx 5.6$  wt.%), the mechanical properties of S-segregated Ni start to degrade drastically in 23°C air, i.e., there is a precipitous decrease in total elongation, tensile strength, modulus of toughness, reduction-in-area, and a sharp increase in percent intergranular cracking (IGC). This means that, at a sufficiently high level of S segregation, a grain boundary in the Ni-S system loses metallic strength and behaves more like a clay. Okamoto and his coworkers also showed that the volume fraction of amorphization of S-ion-implanted specimens starts to increase drastically when the bulk S concentration exceeds  $\approx 5.6$  wt.%.<sup>50,51</sup> Based on these observations, they concluded that S-induced intergranular fracture in binary Ni-S is explained well by disorder-induced melting of a Ni-S-rich thin film that formed on grain boundaries. They proposed that this process is strongly influenced by S and H concentrations, stress, temperature, and irradiation damage. We believe that the results of Okamoto and his coworkers constitute a very useful framework that can shed light on the mechanistic understanding of IASCC.

Similar to the observation of Heuer et al.,<sup>50</sup> predominantly IG fracture was observed in one of our highstrain-rate bending tests in 23°C air, specifically, a Type 304 SS Heat L13, which contained an unusually high concentration of S (0.022 wt.%) and an unusually low concentration of Mn (0.36 wt.%). The specimen was neither exposed to water nor charged with hydrogen before the bending test. Therefore, the intergranular fracture in the specimen cannot be explained on the basis of H-induced embrittlement of grain boundaries. The high-magnification intergranular fractograph of the specimen, shown in Fig. 79, is characterized by a
smooth grain boundary separation, a virtual absence of deformation steps, and an indication of significant grain encirclement. These features are similar to those of IASCC failures produced in 289°C water. Because the specimen was irradiated in high-purity helium, it is also difficult to explain the intergranular fracture by grain boundary oxidation or O segregation. The fracture behavior seems to differ from limited amount of near-surface intergranular separation reported by Onchi and his coworker<sup>52,53</sup> for thermally sensitized and irradiated 304 SSs that were tested in inert gas at  $\approx 290$ °C. Nevertheless, the experiment of Fig. 79 shows that purely mechanical intergranular failure similar to that reported in the unirradiated Ni-S system can occur in a highly irradiated austenitic SS that contains an unusually high concentration of S.

Considering the observations discussed above, we believe that it is important to better understand the structure and properties of Ni- and S-rich thin films and islands that have been reported by Thomas and Bruemmer<sup>32-34</sup> and Dumbill.<sup>31</sup> The former investigators conducted extensive studies on crack-tip microstructure and microchemistry of field-cracked BWR and PWR core internal components.<sup>33,34</sup> Similar studies were also performed by Dumbill<sup>31</sup> and Bruemmer and Thomas<sup>32</sup> on unirradiated, thermally sensitized 304<sup>31</sup> and 316<sup>32</sup> SSs that were tested in sulfated<sup>31</sup> or deoxygenated<sup>32</sup> BWR-like water at 289°C.



Figure 79. Fracture surface morphology of high-S, low-Mn Type 304 SS Specimen L13-03B (S, 0.022 wt.%; Mn, 0.36 wt.%; ≈3 dpa in helium at 289°C) produced during bending in 23°C air without previous exposure to water or H charging.

In Ref. 33, Thomas and Bruemmer report extensively on the crack-tip grain boundary microstructural characteristics of a PWR baffle bolt and a BWR top guide, both of which cracked during service. A layer of

Cr-Fe-type spinel oxide sandwiched between the two adjacent grain matrices was always observed ahead of an open crack tip. Such an oxide was fine-grained and contained tiny pores (diameter 2-10 nm). In the case of the BWR top guide, numerous irregularly shaped microcavities, as large as  $\approx 100$  nm in length and  $\approx 50$  nm in width, were observed, usually at the metal/oxide boundary and near the tip of the spinel oxide.

Direct evidence of  $Ni_xS_y$ -type thin films produced between such a spinel oxide and the metal matrix is observed in several photomicrographs, e.g., Figs. 3-10 and 3-13 in Ref. 33 for the PWR baffle bolt. Small islands of Ni or  $Ni_xS_y$ -type phase are also observed. The width of such  $Ni_xS_y$ -type thin film is extremely small, i.e., <7 nm, based on data from energy-dispersive spectroscopy (EDS) and <3 nm based on bright-field image. In the case of the BWR top guide, a similar type of thin film (thickness  $\approx$ 3 nm), which exhibits the same imaging contrast is observed at the boundary between the metal matrix and the spinel oxide, e.g., Figs. 4-11, 4-12, 4-14, 4-17, and 4-20 of Ref. 33.

Results of analysis by energy dispersive spectroscopy on the top guide indicate that such film contains a high concentration of Ni. However, data for S distribution near the film were not reported. The distance between EDS spots in the analysis was 5-25 nm, significantly larger than the width of such a thin film. Under such conditions, one can easily miss probing a spot correctly placed in the middle of a thin film that is only  $\leq$  nm wide. Therefore, it is not easy to measure the distribution of S near such a film. However, microstructural features such as spinel oxide, voids, cavities, the position and shape of an open crack, and Ni<sub>x</sub>S<sub>y</sub>-type thin film can be detected readily through bright-field imaging and tilting. A dark-field image of a  $\leq$ 3-nm-wide Ni<sub>x</sub>S<sub>y</sub>-type thin film, which is a more valuable piece of information, would be very difficult to obtain in an irradiated core internal. Dumbill reported results of X-ray mapping that show a matching distribution of Ni and S for a crack tip in his unirradiated sensitized 304 SS specimen.<sup>31</sup> The Ni- and S-rich thin film in his specimen was relatively thicker (i.e., width  $\approx$ 100 nm), and a continuous EDS line scan and matching elemental mapping that clearly showed the nature of such film could be obtained.

Thomas and Bruemmer provided a comprehensive interpretation of the crack-tip microstructures observed in their studies.<sup>33,34</sup> They suggested that Cr- and Fe-rich spinel oxide, observed ahead of an open crack tip, was produced as a corrosion byproduct that filled the crack opening after the crack had advanced along the grain boundary. Thus, the spinel oxide tip was interpreted to be the same as the crack tip. Formation of Ni-rich thin films and islands was attributed to a mechanism of selective dissolution in which Cr and Fe atoms are dissolved preferentially over Ni atoms in the crack-tip water.

Nevertheless, photomicrographs and EDS data given in the reports of Thomas and Bruemmer<sup>32-34</sup> or Dumbill<sup>31</sup> shed light on several new clues that are helpful to the understanding of the metallurgical processes that occur near a crack tip. Thus, we thought that a more detailed integrated evaluation is timely. Several important observations in the following discussion suggest a mechanism in which spinel oxide and Ni and Ni<sub>x</sub>S<sub>y</sub> thin films and islands visible near the crack tip were produced, not after the crack advanced to the position of the spinel oxide tip, but, instead, before the crack advanced in association with internal grainboundary oxidation which occurred ahead of the crack tip.

The first observation is that a crack tip is often visible adjacent to a grain boundary rather than in line with the grain boundary. The offset distance between the line of crack advance and the grain boundary (imaged edge on) was  $\approx 20$  nm in the field-cracked BWR top guide; see Figs. 4-19 and 4-21 in Ref. 33. In the case of unirradiated thermally sensitized 304 and 316 SSs tested in deoxygenated BWR-like water, the offset distance was  $\approx 90$  nm and  $\approx 15$  nm, respectively, (Ref. 32). This observation is difficult to explain on the basis

of preferential dissolution of the grain-boundary metal; it is equally difficult to explain on the basis of a preroise that the grain boundary itself shifts continuously ahead of an advancing crack tip. The offset distance is, in fact, the same as the distance between the grain boundary and the metal/spinel boundary, in which a Ni<sub>x</sub>S<sub>y</sub>-type thin film and islands are present, often accompanied by high-density voids and cavities. More voids and cavities tend to appear in regions where thin films or islands of Ni<sub>x</sub>S<sub>y</sub>-type or Ni-rich phases occur. This observation suggests that the voids and cavities were products of localized melting of Ni- and S-rich films or islands. This view is essentially consistent with the results of experiments and the theory advanced by Okamoto and his coworkers on the behavior of S-rich thin film in the Ni-S binary system.<sup>50,51</sup>

The second observation is dark-field morphology of spinel oxide present ahead of an open crack tip. Such a dark-field image was reported in Ref. 32 for a crack tip in an unirradiated, thermally sensitized 316 SS. In the image, extremely small particles of spinel oxide (diameter  $\approx 5$  nm) are observed decorating the grain boundary  $\approx 290$  to  $\approx 450$  nm ahead of the open crack tip, and twin boundaries  $\approx 350$  nm ahead of the crack tip. This observation provides direct evidence of internal oxidation on grain boundaries and twin boundaries in a BWR-like environment, and it is reasonable to expect a similar phenomenon in a BWR core internal. For ordinary grain boundaries (i.e., grain boundaries that are not involved in crack-tip advancement), evidence of grain-boundary internal oxidation has been reported only for the heat-affected zone (HAZ) in BWR core shroud welds that were prepared by shielded-metal-arc or submerged-arc procedure.<sup>26,54</sup> Oxygen contamination in such a HAZ is pronounced, and because O solubility in the grain matrices is extremely low near room temperature, O atoms segregate to grain boundaries during cooling of the HAZ. Internal oxide particles produced by this process have been characterized by Auger electron spectroscopy in Refs. 26 and 54 by transmission electron microscopy in Ref. 26.

The third observation is pores and voids that decorate previous twin boundaries located within a spinel oxide ahead of an open crack tip. Such a feature, visible in a top guide crack tip in Figs. 4-9 and 4-10 of Ref. 33, cannot be explained on the basis of a premise that the spinel oxide filled the crack opening up to the tip of the spinel oxide. Such a feature is, however, explained well based on a premise that a grain boundary region that contains a twin was oxidized internally before the crack advanced to that particular location.

The fourth observation notes the tendency that more cavities are observed near the tip of the spinel oxide layer than near the metal/spinel boundary, e.g., the top guide crack-tip structure shown in Figs. 4-20 to 4-25 of Ref. 33. Also, more Ni and Ni-sulfide islands tend to be present near the tip of a spinel oxide, e.g., Figs. 3-14 (baffle bolt) and 4-18 (top guide) in Ref. 33. This behavior is not explained well based on a premise that the spinel oxide filled the crack opening but is explained well based on a premise that the spinel oxide filled the crack opening but is explained well based on a premise that the spinel oxide internal oxidation of the grain-boundary. In the latter process, Cr and Fe atoms are expected to be selectively oxidized by the oxygen atoms that diffuse down the grain boundary, thereby gradually enriching the metal near the tip of the oxide with Ni. At the same time, S atoms are predicted to remain in the metal ahead of the spinel tip rather than being incorporated in the spinel oxide in which their solubility is lower (than in the metal).

The above observations lend strong support to a mechanism in which a spinel oxide layer and Ni- and S-rich thin films and islands are produced ahead of an advancing crack tip, and localized melting of the Niand S-rich films and islands plays a primary role in crack propagation. The result of such melting is the production of voids and cavities at the metal/spinel boundary and near the tip of the spinel layer.

#### 14.2 An IASCC Model

On the basis of several key observations in this study and elsewhere, and on the basis of the discussions above, we propose a new IASCC mechanism for irradiated austenitic SS. The model, initially reported in Ref. 55 and schematically illustrated in Fig. 80, is based on the following steps, some postulated and some supported by observations in this study or other investigations:

- (1) Many incipient cracks are nucleated in the brittle surface oxide layer that is in contact with water.
- (2) When one such incipient crack encounters a grain boundary, the metal in front of the crack tip is oxidized preferentially along the grain boundary, because stress and defects (crystallographic and irradiation-induced defects) are higher at the grain boundary. A grain boundary is a preferential path for faster diffusion of O from the water, especially in a highly irradiated steel.
- (3) The rate of O transport from water to grain-boundary metal is strongly influenced by the thickness and morphology of the Cr oxide layer that covers the crack-tip grain boundary. When grain boundary Cr depletion is significant, the Cr oxide layer is thinner and less protective, which allows a faster permeation and transport of O. If the Cr oxide layer that covers the crack-tip grain boundary is ruptured by twinning or dislocation channeling, O transport to the grain-boundary metal is facilitated.
- (4) The metal at the crack-tip grain boundary is gradually converted to Cr-Fe-type spinel oxide, because Cr and Fe atoms are preferentially (or selectively) oxidized over Ni atoms by the O atoms that diffused down the grain boundary. Nickel atoms are significantly more difficult to oxidize than Fe and Cr. The free energies of oxidation at ≈300°C of Ni, Fe, and Cr are, respectively, -92, -111, and -155 kcal/mole of O<sub>2</sub>. Thus, most Ni atoms remain unoxidized in the metal matrix. The result is that, Ni atoms, excluded from the growing Cr-Fe spinel oxide layer on the grain boundary, accumulate near the metal/spinel boundary and at the tip of the spinel oxide.
- (5) At the same time, S atoms segregated on the grain boundary are also pushed out of the spinel oxide, because the solubility of S is lower in the oxide than in the metal.
- (6) Eventually, Ni- and S-rich thin films or islands form at the metal/spinel boundary and at the tip of the growing spinel oxide layer. The Ni- and S-rich region can be in the shape of a continuous or discontinuous film or a small isolated island. Nickel and S atoms accumulate more at the tip of the long spinel oxide layer than at the metal/spinel boundary, because less oxidation occurs in the latter direction.
- (7) Some S ions accumulate in the crack tip water because of dissolution of soluble sulfides.<sup>56</sup> Some S atoms are released into the metal due to irradiation-induced instability of MnS<sup>57</sup> and subsequently diffuse to the thin region of metal that is in contact with the Ni- and S-rich film under the influence of high tensile stress.
- (8) When the concentration of S in the Ni- and S-rich thin film or island exceeds a threshold level, the film or island melts or is amorphized, thereby losing its metallic strength. When the thin film or island melts, voids and cavities are formed near the metal/spinel boundary and at the tip of the spinel layer.

- (9) The crack tip then advances along the weakened metal/oxide boundary, i.e., along the thin region in which the Ni- and S-rich thin film and islands melt or are amorphized, thereby producing voids and cavities.
- (10) Voids and cavities tend to be produced more ahead of the spinel tip than at the metal/spinel boundary because of the factor described in Step 6. When a large cavity(ies) form at the spinel tip, crack advance halts until a new spinel layer forms ahead of such a cavity(ies). At this time the line of crack advance can switch.
- (11) Once the Ni- and S-rich thin films or islands melt or are amorphized, they lose their crystalline structure. The polyhedral crystal structure of the phase, such as the cage structure described by Ashby and Spaegen,<sup>58</sup> is broken. Then, the S atoms incorporated in the polyhedral cages diffuse away into the metal matrix.
- (12) Depending on postcracking thermal history, the degree of such back-diffusion of S atoms varies. Then, the S-to-Ni ratio and the distribution of S and Ni near voids and cavities observed in a postcracking examination may vary significantly.

#### 14.3 Role of Carbon

The role of carbon that tends to suppress IASCC susceptibility at very low concentrations of S ( $\leq 0.002$  wt.%) is not well understood at this time. However, an attractive model that can explain the beneficial effect is based on a postulation that C atoms are dissolved in significant amounts in the Ni- and S-rich thin film. As shown by Heuer et al. a Ni- and S-rich thin film loses metallic strength when the S content in the film exceeds a threshold level.<sup>50</sup> Such a loss of metallic strength is attributed to severe localization of free electrons by S atoms incorporated in the Ni-S-type film, i.e., trapping of free electrons by S. When deprived of free electrons, the film no longer exhibits metallic behavior or metallic strength. A simple consideration suggests that C atoms can easily occupy the same octahedral sites in the unit cell of Ni sulfide that would be normally occupied by S atoms. Thus, when C concentration is high near the Ni- and S-rich thin film that is formed at the crack-tip grain boundary, S and C atoms compete to occupy the same octahedral sites. Under such a situation, C dissolved in the film is expected to promote a strong covalent bonding with Ni, in effect suppressing the effect of trapping free electrons by S. The limited information described in Section 13.2 indicates that the solubility of C in a CuS- and NiS-type thin film is significant, which is consistent with, but does not prove, the model outlined above. A further investigation is desirable to provide a better mechanistic understanding of the role of C atoms on BWR IASCC.



Figure 80. Schematic illustration of an IASCC model

CZ3

# **15 Conclusions**

Slow-strain-rate tensile tests in BWR-like water (dissolved oxygen  $\approx 8$  ppm) and fractographic analysis were conducted on commercial and laboratory-fabricated model heats of austenitic stainless steels irradiated at 289°C in helium to  $\approx 0.3 \times 10^{21}$  n cm<sup>-2</sup>,  $\approx 0.9 \times 10^{21}$  n cm<sup>-2</sup>, and  $\approx 2.0 \times 10^{21}$  n cm<sup>-2</sup> (E > 1 MeV) ( $\approx 0.4, 1.4,$  and  $\approx 3$  dpa). Susceptibility of the steels to irradiation-assisted stress corrosion cracking (IASCC) was determined on the basis of the degree of intergranular fracture surface morphology. The investigation has led to the following conclusions.

- (1) At ≈0.4 dpa, IASCC susceptibility was negligible for all tested heats, with the exception of a 316L-like high-purity alloy that contained 0.003 wt.% C and 0.004 wt.% S. At 0.4 and 1.4 dpa, significant transgranular fracture was mixed with intergranular fracture. At 3 dpa, the tendency to transgranular stress corrosion cracking was negligible, and fracture surface morphology was either dominantly intergranular, as in field-cracked core internal components, or dominantly ductile or mixed. This behavior indicates that percent IGSCC determined at ≈3 dpa is a good measure of IASCC susceptibility.
- (2) At 1.4 dpa, a beneficial effect of a high concentration of Si (0.8-1.5 wt.%) was observed. At 3 dpa, however, such effect was obscured by a strongly deleterious effect of S. From the standpoint of BWR IASCC, which usually occurs at >2.5 dpa, typical variation in Si concentration in the range of 0.4-1.1 wt.% in Types 304, 316, and 348 steel is, therefore, not an important factor.
- (3) Excellent resistance to IASCC was observed up to ≈3 dpa for seven heats of Types 304 and 316 steel that contain very low S concentrations of ≤0.002 wt.% and for one heat of Type 348 steel that contains 0.005 wt.% S. Susceptibility of Types 304 and 316 steel that contain >0.003 wt.% S increased crastically. These observations indicate that a sulfur-related critical phenomenon plays an important role in IASCC. A sufficiently low concentration of S (i.e., <0.002 wt.%) is the primary material factor necessary to ensure good resistance to IASCC.</p>
- (4) For Types 304L and 316L steel and their high-purity counterparts that contain <0.025 wt.% C, a sulfur concentration of <0.002 wt.% alone is not a sufficient material condition to ensure good resistance to IASCC. This is in distinct contrast to the behavior of the high-C counterparts (i.e., Types 304 and 316 steel that contain <0.002 wt.% S and >0.03 wt.% C), which exhibit good resistance to IASCC. The contrasting behavior indicates that, at sufficiently low concentrations of S, a high concentration of C is beneficial. However, at S concentrations >0.003 wt.%, the deleterious effect of S is so dominant that a high concentration of C is not an important factor.
- (5) A two-dimensional map was developed in which susceptibility or resistance to IASCC is shown as a function of bulk concentrations of S and C. Data reported in the literature are consistent with the map. The map is helpful to predict relative IASCC susceptibility of Types 304 and 316 steel. A similar map is helpful to predict IASCC behavior of Type 348 steel.
- (6) At S concentrations >0.003 wt.%, IASCC susceptibility is thought to be influenced significantly by various material-related factors, such as the degree of S segregation to grain boundaries, annealing and cooling during fabrication, component thickness, presence of delta ferrite, Mn content, precipitation of

sulfides, solubility of sulfides in the crack-tip water, and the thickness and morphology of the Cr oxide on the crack-tip grain boundary, which controls oxygen permeation and diffusion along the grain boundary.

- (7) A model alloy that contains ≈3 vol.% globular-shaped delta ferrite and ≈21 wt.% Cr exhibited excellent resistance to IASCC, despite the fact that it contained a very high S concentration of 0.028 wt.%. The excellent resistance is attributed to the effect of delta ferrite, which traps S atoms, and the effect of a high concentration of Cr, which forms Cr oxide and helps to block oxygen permeation and diffusion along the austenite grain boundary.
- (8) Grain-boundary segregation of S was observed for BWR neutron absorber tubes. It is not clear, however, whether the S segregation occurred via a thermal process, an irradiation-induced process, or both.
- (9) Based on the results of this study, an IASCC model has been developed. The model is based on several key steps known or postulated to occur at the crack-tip grain boundary, i.e., grain-boundary segregation of Ni and S, preferential oxidation of Cr and Fe atoms over Ni atoms by oxygen that diffuses along the grain boundary, formation of a Cr-Fe spinel oxide ahead of the crack tip, formation of Ni- and S-rich thin films and islands at the metal/spinel boundary, and melting or amorphization of the Ni-and S-rich thin films and islands. The model also predicts that the thickness and morphology of the Cr oxide that covers the crack-tip grain boundary strongly influence the rate of oxygen permeation and diffusion along the grain boundary.

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NRC FORM 335 (9-2004) NRCMD 3.7	U. S. NUCLEAR REGULATORY COMMISSION	1. REPORT NUMBER (Assigned by NRC. Add Vol., Supp., Rev., and Addendum Numbers, if any.)	
BIBLIOGRAPHIC DATA SHEET (See instructions on the reverse)		NUREG/CR-6892 ANL04/10	
		3 DATE REPORT PUBLISHED	
Irradiation-Assisted Stress Corrosion Cracking Behavior of Austenitic Stainless Steels Applicable to LWR Core Internals		MONTH	YEAR
		January	2006
		4. FIN OR GRANT NUMBER Y6388	
5. AUTHOR(S) H. M. Chung and W. J. Shack		6. TYPE OF REPORT Technical; Topical	
		7. PERIOD COVERED (Inclusive Dates)	
<ul> <li>8. PERFORMING O REANIZATION - NAME AND ADDRESS (If NRC, provide Division, Office or Region, U.S. Nuclear Regulatory Commission, and mailing address; if contractor, provide name and r ailing address.)</li> <li>Argonne National Laboratory</li> <li>9700 South Cass Avenue</li> <li>Argonne, IL 6()439</li> </ul>			
9. SPONSORING ORGANIZATION - NAME AND ADDRESS (If NRC, type "Same as above": if contractor, provide NRC Division, Office or Region, U.S. Nuclear Regulatory Commission, and mailing address.)			
Division of Engineering Technology Office of Nuclear Regulatory Research U.S. Nuclear Regulatory Commission Washington, DC 20555–0001			
10. SUPPLEMENTARY NOTES William H. Cullen, Jr., NRC Project Manager			
As LWRs age and neutron fluence increases, core internals become susceptible to irradiation-assisted stress corrosion cracking (IASCC). To better understand the mechanism of this type of degradation, austenitic stainless steels were irradiated in the Halden reactor under BWR conditions, and slow-strain-rate tensile tests were conducted in BWR-like water on 27 commercial and model laboratory heats. Fractographic and microchemical analyses were performed to characterize grain-boundary segregation of alloying and impurity elements. As neutron damage increases to $\approx 3$ dpa, sulfur atoms play a dominant role in causing IASCC. To provide good resistance to IASCC in Types 304 or 316 steel, it is necessary to ensure a sulfur content of <0.002 wt.% and a carbon content of >0.03 wt.%. A two-dimensional map was developed in which IASCC susceptibility or resistance of Types 304 or 316 is shown as a function of bulk concentrations of S and C. A similar map is applicable to Type 348 steel. Nine heats of Types 304, 316, and 348 steel were identified to be resistant to IASCC under BWR conditions. On the basis of the stress-corrosion-cracking tests and the microstructural characterization, a mechanistic IASCC model was developed.			
12. KEY WORDS/DES/CRIPTORS (List words or phrases that will assist researchers in locating this report.) Irradiation-induced degradation Stress corrosion cracking BWR core internals Grain boundary segregation		13. AVAILABI unlimited	LITY STATEMENT
		14. SECURITY CLASSIFICATION (This Page) unclassified (This Report)	
		unclassified 15. NUMBER OF PAGES	
		16.PRICE	
NRC FORM 335 (9-20)4)		PRINTED ON RECYCLED PAPER	

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### IRRADIATION-ASSISTED STRESS CORROSION CRACKING BEHAVIOR OF AUSTENITIC STAINLESS STEELS APPLICABLE TO LWR CORE INTERNALS

JANUARY 2006

UNITED STATES NUCLEAR REGULATORY COMMISSION WASHINGTON, DC 20555-0001

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