Hot Corrosion Behavior of Superalloy in Different Corrosive Environments

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Abstract

Hot corrosion is a serious problem in aircraft, marine, industrial and land-base gas turbines. To obviate these problems, superalloy superfer 800H (Midihi grade) is one of the prominent materials for the high temperature applications. The present work investigates hot corrosion behaviour of superfer 800H exposed to two different molten salt environments Na₂SO₄ + 60% V₂O₅ and Na₂SO₄ + 5% V₂O₅ + 5% NaCl respectively, at 900°C under cyclic conditions. The weight change measurements made during the experiments are used to determine the kinetics of hot corrosion. It is observed that the formation of scale rich in Cr₂O₃, NiO and spinel NiCr₂O₄ has contributed for the better hot corrosion resistance of superfer 800H. Inspection for 10, 20, 35 and 50 cycles reveals that after 10 cycles Crack has been initiated in the scale and it is clearly visible in the presence of NaCl salt. On subsequent cycles, crack propagates perpendicular to surface and as it encounters inclusions.

Keywords

Hot Corrosion, Superalloy, Superfer 800H, Cyclic Conditions, Spinel NiCr₂O₄

1. Introduction

Degradation by high-temperature oxidation, hot corrosion and erosion are the main failure modes of components in the hot sections of gas turbines, boilers, industrial waste incinerators, etc. Superfer 800H have been developed
for high temperature applications, superfer finds application in the gas turbine industry, constituting over 50% of the gas turbine weight due to their good mechanical properties at elevated temperatures [1]-[4]. Superfer exhibits superior mechanical strength, surface stability, creep and fatigue resistance at high temperature [5]. The turbine engines are exposed to extremely high temperature and harsh environment and therefore tend to suffer from significant material degradation during service [6] [7]. Low-grade fuel oils and fossil fuels used in energy generation systems contains complex mixtures of molten sodium sulfate (Na₂SO₄) and vanadium pentoxide (V₂O₅) [8] [9]. The Na₂SO₄ can be ingested in the turbine intake air or it can be produced by a reaction between sodium chloride (NaCl) ingested with the intake air and sulfur impurities in the fuel [10]. Vanadium is present in the fuel in the form of vanadium porphyrin, which transforms during combustion into V₂O₃. V₂O₃ and Na₂SO₄ form low melting point inorganic compounds, which undergoes eutectic reaction below 600°C. NaCl is present in marine environment, and together with Na₂SO₄ it lowers the eutectic melting point to 620°C. When the temperature exceed melting point of the deposits [11], these compounds starts slowly depositing on the turbine blades, consequently corrosion rate rapidly increases due to faster transport phenomena in liquid phase which cause catastrophic corrosion phenomena [12]. Hence, for effective and efficient functioning of gas turbine components, it is highly essential to develop superalloy with high-temperature strength as well as good hot corrosion resistance [13]. However, there is still a lack of basic understanding of how these superalloys behave in severe (Na₂SO₄ + 60% V₂O₅) operating environment. Therefore, the hot corrosion behaviour of superfer 800H has been investigated at 900°C in two most corrosive environments (Na₂SO₄ + 60% V₂O₅ and Na₂SO₄ + 5% V₂O₅ + 5% NaCl) in the present work. The kinetics of corrosion of superalloy substrates was determined from the weight change of the samples calculated during the experiments. Inspections for 10, 20, 35 and 50 reveal the mechanism of hot corrosion via crack propagation. It was found that after 10 cycles Crack has been initiated in the scale and it is clearly visible in the presence of NaCl salt. On subsequent cycles, crack propagates perpendicular to surface and as it encounters inclusions, intermetallics it starts propagating parallel to surface. Through cracks, molten salts attack the substrate and cause hot corrosion. But extent of corrosion is less as the thickness of scale formed after 50 cycles is not more than 50 µm even in most severe environment (Na₂SO₄ + 60% V₂O₅). XRD, SEM and FE-SEM/EDAX, and X-ray mapping were used to characterize the corrosion products in order to render an insight into the corrosion mechanisms.

2. Experimental Procedure

2.1. Substrate Material

The Fe-based substrate material selected for this study, namely superfer 800H, were provided by Mishra Dhatu Nigam Limited, Hyderabad (India) in the rolled sheet form. The nominal chemical composition of the substrate materials is reported in Table 1.

2.2. Substrate Preparation

The specimens, with dimensions of approximately 20 × 15 × 5 mm³, were cut from the alloy sheets of superfer 800H. The specimens were polished using emery papers of 220, 400, 600 grit sizes and subsequently on 1/0, 2/0, 3/0 and 4/0 grades. Final polishing was carried out on a cloth polishing wheel machine with 1 µm alumina powder suspension. Subsequently the specimens were properly cleaned with acetone and dried in hot air (250°C) for 3 - 4 hours.

2.3. Molten Salt Corrosion Test

Cyclic studies were performed for the substrate in two different molten salt environments (Na₂SO₄ + 60% V₂O₅ and Na₂SO₄ + 5% V₂O₅ + 5% NaCl) for 50 cycles. Each cycle consisted of 1 hr of heating at 900°C in a silicon

<table>
<thead>
<tr>
<th>Alloy Midhani Grade (Similar Grade)</th>
<th>Chemical Composition (wt%)</th>
</tr>
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<tbody>
<tr>
<td>Superfer 800H (Incoloy 800H)</td>
<td>Fe  43.8 Ni 32.0 Cr 21.0 Ti 0.3 Al 0.3 Mo 1.5 Mn 1.0 Si Cu Ta C</td>
</tr>
</tbody>
</table>

Table 1. Chemical composition of the substrate material superfer 800H.
carbide tube furnace followed by 20 minutes of cooling at room temperature (25°C). The purpose of imposing cyclic conditions was to create an accelerated environment as observed in real cases for hot corrosion testing. The specimens were mirror polished down to 1 µm alumina on a cloth-polishing wheel before the corrosion run. A coating of uniform thickness with 3 - 5 mg/cm² of salt was applied with a camel hair brush on the preheated sample (250°C). The weight change measurements were taken at the end of each cycle using an electronic balance machine (Model 06120, Contech) with a sensitivity of 1 mg. The spalled scale was also included at the time of measurement of the weight change to determine the total rate of corrosion. The kinetics of corrosion was analyzed from the results of weight change measurements. The samples after hot corrosion were analyzed by XRD for the phase identification and FE-SEM/EDAX for surface morphological and compositional analysis and X-ray mapping for cross sectional compositional analysis.

3. Results

3.1. Visual Observations

The superalloys during hot corrosion in the molten salt environment at 900°C have shown spalling/sputtering right from the initial few cycles which intensified with the progress of study with lot of corrosion products formed in the boat. The surface became rougher with progressive exposure of time with a uniform pitting observed throughout the surface of the samples. A greenish scale appeared on the surface of superfer 800H; during initial cycles the colour was dark grey which turned greenish with grey patches. The oxide scale has two regions, with dark and light grey scales, dark grey scale has brownish tinges distributed randomly.

3.2. Cyclic Hot Corrosion Test

Figure 1 and Figure 2 show the weight gain/unit area for molten salt (Na₂SO₄ + 60% V₂O₅ and Na₂SO₄ + 5% V₂O₅ + 5% NaCl) environment, respectively, for the superalloys subjected to 900°C for 50 cycles. The parabolic rate constants calculated for the superfer 800H shows that the corrosion rate is higher in Na₂SO₄ + 60% V₂O₅ environment. The weight gain square (mg²/cm⁴) versus time (number of cycles) plots are drawn to establish the rate law for the hot corrosion. It is observed from the graph that superfer 800H follows nearly parabolic rate law in both the environment. The parabolic rate constant \(K_p\) was calculated by a linear least-square algorithm to a function in the form of \((W/A)^2 = K_p t\), where \(W/A\) is the weight gain per unit surface area (mg/cm²) and “t” indicates the number of cycles represents the time of exposure. The parabolic rate constants for both the environments (Na₂SO₄ + 60% V₂O₅ and Na₂SO₄ + 5% V₂O₅ + 5% NaCl) were calculated for 10, 20, 35 and 50 cycles and are reported in Table 2.

3.3. X-Ray Diffraction Analysis (XRD) of Scale

The XRD patterns for the superfer 800H subjected to molten salt (Na₂SO₄ + 60% V₂O₅ and Na₂SO₄ + 5% V₂O₅ + 5% NaCl) environments at 900°C after 10, 20, 35 and 50 cycles are shown in Figure 3 and Figure 4. The ma-

![Figure 1. Weight gain/area vs. number of cycles plot for superfer 800H subjected to cyclic hot corrosion in Na₂SO₄ + 60% V₂O₅ environment at 900°C for 10, 20, 35 and 50 cycles.](image)
Figure 2. Weight gain/area vs. number of cycles plot for super-fer 800H subjected to cyclic hot corrosion in Na₂SO₄ + 5% V₂O₅ + 5% NaCl environment at 900°C for 10, 20, 35 and 50 cycles.

Figure 3. X-ray diffraction patterns for the super 800H subjected to cyclic hot corrosion in Na₂SO₄ + 60% V₂O₅ environment at 900°C for 10, 20, 35 and 50 cycles.

Figure 4. X-ray diffraction patterns for the super 800H subjected to cyclic hot corrosion in Na₂SO₄ + 5% V₂O₅ + 5% NaCl environment at 900°C for 10, 20, 35 and 50 cycles.
Table 2. Variation of parabolic rate constant, kp with number of cycles.

<table>
<thead>
<tr>
<th>Number of Cycles</th>
<th>$K_p \times 10^{-9} , g^2 \cdot cm^{-4} \cdot s^{-2}$ $(Na_2SO_4 + 60% V_2O_5)$</th>
<th>$K_p \times 10^{-10} , g^2 \cdot cm^{-4} \cdot s^{-2}$ $(Na_2SO_4 + 5% V_2O_5 + 5% NaCl)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>1.684</td>
<td>9.44</td>
</tr>
<tr>
<td>20</td>
<td>2.931</td>
<td>1.986</td>
</tr>
<tr>
<td>35</td>
<td>2.437</td>
<td>4.580</td>
</tr>
<tr>
<td>50</td>
<td>3.119</td>
<td>4.767</td>
</tr>
</tbody>
</table>

jor phases identified in Na$_2$SO$_4$ + 60% V$_2$O$_5$ environments are NiO, Cr$_2$O$_3$, Ni(VO)$_3$, NiCr$_2$O$_4$ and $\gamma$-Fe$_2$O$_3$. NiCl$_2$ is the addition phase identified in the presence of NaCl salt.

3.4. FE-SEM/EDAX Analysis of the Scale

3.4.1. Surface Morphology of the Scale

FE-SEM/EDAX micrographs with EDS spectrum of the superfer 800H specimens after cyclic hot corrosion in molten salt (Na$_2$SO$_4$ + 60% V$_2$O$_5$ and Na$_2$SO$_4$ + 5% V$_2$O$_5$ + 5% NaCl) environments for 10, 20, 35, 50 cycles at 900°C are shown in Figure 5 and Figure 6, respectively. In superfer 800H, the formation of hemispherical shaped protrusion was observed, which mainly consists of NiO, Cr$_2$O$_3$ and Fe$_2$O$_3$ as the predominant phases.

3.4.2. X-Ray Mapping

The hot corroded samples were cut across the cross-section and mounted in transoptic mounting resin, mirror-polished and gold coated to facilitate X-ray mapping by FE-SEM/EDAX of the different elements present across the scale. X-ray mapping analyses of the scale formed after hot corrosion in (Na$_2$SO$_4$ + 60% V$_2$O$_5$ and Na$_2$SO$_4$ + 5% V$_2$O$_5$ + 5% NaCl) environments at 900°C for 10, 20, 35 and 50 cycles (Figure 7, Figure 8) reveals that the surface scale mainly consist of Cr, Ni and Fe in the upper most part of the scale, presence of oxygen in the top scale shows the formation of Cr$_2$O$_3$, NiO and Fe$_2$O$_3$ layer. In the subscale region, there is a thick band of chromium (Cr) which indicates the formation of relatively dense scale of Cr$_2$O$_3$, meager amount of S is penetrated deep in to the substrate.

4. Discussion

The weight change data for superfer 800H subjected to hot corrosion test in molten salt environments are plotted in Figure 1 and Figure 2. It indicates that the superfer 800H followed a parabolic rate law up for both the environment. The higher weight gain of the specimens during the first few cycles might be due to the rapid formation of oxides (formation of Cr$_2$O$_3$) at the boundaries and within the open pores due to the penetration of the oxidizing species. Further subsequent increase in weight is gradual. Values of parabolic rate constant $K_p \times 10^{-10} \, g^2 \cdot cm^{-4} \cdot s^{-2}$ were obtained from slope of the linear regression fitted line and are tabulated in Table 2. XRD-analysis of hot corroded samples indicate the formation of Fe$_2$O$_3$, NiO, Cr$_2$O$_3$ and spinel (NiCr$_2$O$_4$) as major phases. Surface morphology and composition analysis of corroded sample are shown in (Figure 5 and Figure 6). The EDAX analysis of the surface scale developed on superfer 800H in Na$_2$SO$_4$ + 60% V$_2$O$_5$ environment (Figures 5(a)-(d)) is crack free with dense clusters mainly consisting of Cr, Fe, Ni and O rich elements, thereby suggesting the formation of Cr$_2$O$_3$, Fe$_2$O$_3$, NiO and their spinel (NiCr$_2$O$_4$). Whereas scale developed on superfer 800H in Na$_2$SO$_4$ + 5% V$_2$O$_5$ + 5% NaCl environment (Figures 6(a)-(d)) mainly consisting of Cr, Fe, and O rich elements which suggesting the formation of Cr$_2$O$_3$ and Fe$_2$O$_3$. The formation of spinel of NiCr$_2$O$_4$ via solid phase reaction between NiO and Cr$_2$O$_3$ in the oxide scales helps to develop oxidation resistance as the spinel phase that usually has lower diffusion coefficients of the cations and anions than those in their parent oxides. The superfer 800H (Figure 5(d) and Figure 6(d)) show the formation of uneven protrusion in entire oxidised surface along the scale grain boundaries, similar protrusion behaviour was reported in the earlier literature [14] [15] during isothermal oxidation studies on plasma-sprayed NiCrAlY bond coat and oxidation and hot corrosion behaviours of superalloys at 900°C.

X-ray mapping (Figure 7) suggests that in molten salt environments, hot corrosion proceed through initiation and propagation of cracks, diffusion of molten salts through cracks and dissolution and re-precipitation of Cr$_2$O$_3$.
layer along the path of crack. In Na$_2$SO$_4$ + 60% V$_2$O$_5$ salt after 20 cycles, cracks are clearly revealed (Figure 7(b)). It is also observed that crack initiated at the surface (scale) first propagates perpendicular to the surface and as it encounters any obstacle (inclusions, inter-metallic) it starts propagating parallel to the surface. Figure 7(c) shows crack propagation and salt penetration through cracks into the substrate. After 50 cycles (Figure 7(d)), there is complete rupture of scale. In case of Na$_2$SO$_4$ + 5% V$_2$O$_5$ + 5% NaCl environment, presence of Cl$^-$ enhances the crack formation. Addition of NaCl accelerates the hot corrosion, as it reacts with the oxides thus releasing chlorine which in turn reacts with the oxides to form the volatile chlorides, as per the following reactions [16] [17].

$$8\text{NaCl(l)} + 2\text{Cr}_2\text{O}_3(s) + 5\text{SO}_2(g) = 4\text{Na}_2\text{CrO}_4(s) + 4\text{Cl}_2(g)$$
Figure 6. Oxide Scale morphology and variation of elemental composition across the cross-section of superfer 800H subjected to cyclic hot corrosion in Na$_2$SO$_4$ + 5% V$_2$O$_5$ + 5% NaCl at 900˚C after (a) 10 cycles, (b) 20 cycles (c) 35 cycles and (d) 50 cycles.

$\text{Cl}_2$ is able to penetrate through oxide scales quickly along cracks and react with substrate elements such as Cr. Then volatile chlorides are formed:

$$\text{Cr}(s) + \frac{3}{2}\text{Cl}_2(g) = \text{CrCl}_3(g)$$

The volatile chlorides thus formed tend to diffuse out through the grain boundaries to the surface, and in doing so numerous pits and voids are generated at the grain boundaries, which can be seen clearly in Figure 8. The chlorides may reoxidize on the surface of the scales:
Figure 7. FE-SEM/EDAX analysis along with EDS spectrum for the superfer 800H subjected to cyclic hot corrosion in Na₂SO₄ + 60% V₂O₅ at 900°C after (a) 10 cycles, (b) 20 cycles (c) 35 cycles and (d) 50 cycles.

Cl₂ is then regenerated and the corrosion repeated. Other alloy elements, such as Ni, Ti etc. also follow the above reactions. The pits and voids, produced due to the formation of volatile chlorides, provide paths for the further penetration of corrosive species resulting in internal oxidation and sulphidation of the substrates as being reported by Sidhu et al. [18].

It may be noticed that after 10 cycles in Na₂SO₄ + 5% V₂O₅ + 5% NaCl environment (Figure 8(a)), there is no any distinct scale substrate interface, but initiation of crack propagation has been started in the scale and crack propagation path is clearly visible. Through these cracks, molten salt diffuses and attacks the substrate.

\[
2\text{CrCl}_3(\text{g}) + \frac{3}{2}\text{O}_2(\text{g}) = \text{Cr}_2\text{O}_3(\text{s}) + 3\text{Cl}_2(\text{g})
\]
Figure 8. FE-SEM/EDAX analysis along with EDS spectrum for the superfer 800H subjected to cyclic hot corrosion in Na₂SO₄ + 5% V₂O₅ + 5% NaCl at 900°C after (a) 10 cycles, (b) 20 cycles (c) 35 cycles and (d) 50 cycles.

material by dissolving the Cr₂O₃ layer (Figure 8(c)). Propagation of crack into the material, diffusion of molten salt through the cracks and dissolution and reprecipitation of Cr₂O₃ along the crack path is clearly visible in Figure 8(c). Figure 8(d) further supports the hot corrosion mechanism via crack propagation. Intensive spalling/sputtering of the scale of the superfer 800H superalloy can be attributed to severe strain developed due to the precipitation of Fe₂O₃ from the liquid phase and interdiffusion of intermediate layers of iron oxide as has been reported by Sachs [19]. Further, the presence of different phases in a thin layer might impose severe strain on the
film, which may result in cracking and peeling of the scale. Thermal cycles might have also produced cracks in the scale due to different values of thermal expansion coefficients of the substrate and the oxides formed in the scale. These cracks may have allowed the aggressive molten liquid phase to reach the metal substrate and causes internal oxidation. X-ray mapping of corroded superfer 800H also supports internal oxidation via cracks as there is higher percentage of Cr along with Oxygen along the path of crack.

5. Conclusions

- The formation of protective oxide scales such as NiO, Cr₂O₃ and NiCr₂O₄ provide resistance to hot corrosion.
- Depletion of chromium occurs at scale metal interface, scale initially comprises of Cr₂O₃.
- During initial cycles Nickel concentration is high at metal interface region, as the number of cycles increase Nickel concentration increase in scale, scale has NiO and Cr₂O₃.
- Initial scale formed in Na₂SO₄ + 60% V₂O₅ environment is adherent to substrate, while in case of chloride environments (Na₂SO₄ + 10% NaCl, Na₂SO₄ + 5% NaCl + 5% V₂O₅) the initial scale formed is less adherent to substrate and crack propagation path is visible in initial cycles.
- Crack propagates via formation of Cr₂O₃. Region of crack propagation show Chromium and Oxygen in relatively large quantity.

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References

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