THE CONTROL OF SULFUR CONTENT IN NICKEL-BASE, SINGLE CRYSTAL SUPERALLOYS AND ITS EFFECTS ON CYCLIC OXIDATION RESISTANCE

Department of Materials Science & Engineering
University of Pittsburgh

J. L. Smialek
NASA Lewis Research Center

Abstract

State-of-the-art superalloys are useful for high temperature applications, in large part, because they form protective alumina surface films by the selective oxidation of aluminum from the alloy. The adherence of the alumina to the alloy is crucial to maintaining oxidation resistance, particularly under thermal cycling conditions. It is now well established that small additions of reactive elements, such as yttrium, hafnium, and cerium, substantially improve the adherence of alumina films to alloy substrates. While the effects produced by the reactive elements are widely known the mechanisms whereby they improve adherence are not completely understood. Over the last fifty years a number of mechanisms have been proposed. However, it has recently become clear that a major effect of the reactive elements is to tie up sulfur in the alloy and prevent it from segregating to the alloy/oxide interface and weakening an otherwise strong bond.

This paper describes the results of a study of the control of sulfur content in alumina-forming nickel-base superalloys and NiAl by three methods:
1. Addition of Reactive Elements (Y and Hf).
2. Desulfurization in the solid state.
3. Desulfurization in the liquid state.

Additionally, calculations have been performed to determine how much sulfur is available to segregate to the scale/alloy interface and how this quantity is influenced by the type and amount of reactive element in the alloy and the level to which the alloy is desulfurized. Finally, the results from experiments to desulfurize the alloys are described and cyclic oxidation measurements are used to evaluate the calculations.

Introduction

State-of-the-art superalloys and aluminide coatings are useful for high temperature applications, in large part, because they form protective alumina surface films by the selective oxidation of aluminum from the alloy. The adherence of the alumina to the alloy is crucial to maintaining oxidation resistance, particularly under thermal cycling conditions. It is now well established that small additions of reactive elements, such as yttrium, hafnium, and cerium, substantially improve the adherence of alumina films to alloy substrates [1]. While the effects produced by the reactive elements are widely known the mechanisms whereby they improve adherence are not completely understood. Over the last fifty years a number of mechanisms have been proposed. These include:

1. Reactive elements act as vacancy sinks to suppress void formation at the alloy/oxide interface [2,3].
2. Reactive elements form oxide pegs at the alloy/oxide interface [4].
3. Reactive elements alter the growth mechanism of the oxide resulting in reduced growth stresses [5].
4. Reactive elements segregate to the alloy/oxide interface and form a graded seal [6] or otherwise strengthen the alloy/oxide bond [7].
5. Incorporation of the reactive element into the oxide increases its plasticity [8].
6. Reactive elements tie up sulfur in the alloy and prevent it from segregating to the alloy/oxide interface and weakening an otherwise strong bond [9,10].

The importance of the latter mechanism has been illustrated in experiments where hydrogen annealing of nickel-base single crystals decreased the sulfur contents to very low levels and resulted in dramatic improvements in the adherence of alumina films to the alloys [11,12].

This paper presents the results of studies of the effects of reactive element additions to alumina-forming single crystal Ni-base superalloys and the effect of lowering the sulfur content of these alloys and NiAl on the oxide adherence.

Experimental

The alloys studied include three alumina-forming Ni-base single crystal superalloys (PWA 1480, PWA 1484, and PWA 1487) and equiatomic NiAl. The single crystal alloys had an initial sulfur content of 8-10 ppm and were undoped (1480), doped with 0.1Hf (1484), and doped with 0.1Hf+0.1Y (1487). Additional polycrystalline alloys with compositions similar to PWA 1484 but containing 1, 3, and 45 ppm sulfur, provided by Howmet Corporation, were also studied. The NiAl contained 20 ppm sulfur. Most specimens had approximate dimensions 1cm X 1cm X 1mm, however, additional specimens of selected alloys with thicknesses ranging from 0.3 to 3.0mm were annealed in Zr-gettered hydrogen for 100 hours at 1200°C to remove sulfur. All specimens were polished through 600 grit SiC paper, washed in soapy water, and rinsed in ethanol prior to oxidation or hydrogen annealing. Hydrogen annealed specimens were oxidized with no further surface treatment. Cyclic oxidation was performed in air at 1100°C with one cycle consisting of 45 minutes in the hot zone and 15 minutes cooling above the furnace. Selected alloys were also oxidized isothermally in air at 1100°C. The oxidized specimens were examined normal to the surface and in cross-section using optical and scanning electron metallography.
The residual stress in the oxide scale was measured at room temperature via the classical $\sin^2 \psi$ method [13] using a Philips X'pert diffractometer fitted with an open Eulerian cradle and parallel beam optics with a Ni-filter [14]. The x-ray source was a point source with a Cu target. The Tilting method combined with glancing incident angle ("Thin Film geometry") was employed to measure strain/stress in the $\alpha$-alumina layer. The geometrical set up of this technique is schematically shown in Figure 1. This technique was chosen because this setup provides greater intensity of the diffracted x-rays from the alumina layer as a result of increase in the interaction volume of the x-rays in the alumina layer. The specimen is tilted around an axis (parallel to the surface of the specimen and in the diffraction plane) which is the intersection of the diffraction plane with the surface of the specimen. The $(113)$ reflection of the $\alpha$-alumina was scanned with a fixed incidence of $\alpha = 1.75^\circ$ for the tilt angles $\psi = 0^\circ$, $28^\circ$ and $33^\circ$, $\psi = 0^\circ$ when the specimen normal is in the diffraction plane (Figure 1). The tilt angles were selected when $(113)$ diffraction lines could be measured. With this type of geometry, $\psi$ is not the angle of the tilt of the diffraction planes since these planes are inclined with respect to the surface of the specimen. The angle $\omega$ used in the $\sin^2 \psi$ plots is measured between the normal to $(113)$ planes and the normal to the specimen surface. It is given by equation (1) [15].

$$\cos \psi = \cos \omega \cos \psi \cos \alpha$$

(1)

in which $\alpha = \theta_b - \alpha$. $\theta_b$ is the bragg angle for the $(113)$ reflection. The absorption correction of the x-ray intensities is carried out by using the equation in reference 15. The line positions in the $2\theta$ scan were determined by fitting a parabola to the top of the peak [16].

Results And Discussion

Single Crystal Alloys

The results of cyclic oxidation experiments for the Ni-base single crystal alloys at 1100°C are presented in Figure 2. The oxidation resistance of the untreated alloys is in the order $1487 > 1484 > 1480$ which suggests that doping with 0.1Hf improves adherence but that combined doping with 0.1Hf + 0.1Y is more effective. Unfortunately an alloy doped only with Y was not available.

Sulfur removal improved the cyclic oxidation resistance of both 1480 and 1484. These results support the observations that sulfur removal improves the alumina adherence to alloys which do not contain reactive elements (1480). However, the improvement for 1484, which contains Hf, indicates that the sulfur removal somehow makes the Hf more effective. Whether this effect will occur for an alloy containing Y is not yet clear.

Figure 2 also indicates a dependence on specimen thickness for the desulfurized PWA 1480. The hydrogen anneal improved the adherence for the 3.0 mm thick specimen but not to the extent for the 0.3 mm thick specimen. This implies a limitation on the section thickness which can be desulfurized in the solid state. This problem has been addressed by Smialek and coworkers [17,18]. For thin specimens the average sulfur content, $c_s$, may be calculated as a function of time during hydrogen annealing from

$$c_s/c_{s,i} = (8/\pi^2)\exp(-\pi^2 D_s t/L^2)$$

(2)

where: $c_{s,i}$ is the initial sulfur content.
$D_s$ is the sulfur diffusivity in the alloy.
$t$ is the annealing time.
$L$ is the specimen thickness.

Equation (2) will be valid as long as the desulfurization process is diffusion controlled. Taking an initial sulfur content of 10 ppm and approximating $D_s$ as that in pure Ni [19] one calculates from Eqn. (2)
that the 100 hrs., 1200°C anneal should lower the sulfur content to 3 ppm for a 3.0 mm thick specimen and to 10 ppb for a 0.3 mm thick specimen. While the latter number is clearly unrealistic in that diffusion control will not persist to such low concentrations, it indicates that the thinner specimen would be desulfurized to a very low level. On the other hand, the thicker specimen contains enough residual sulfur to segregate and degrade adherence of the alumina scale. Smialek et al [18] have considered the adsorption behavior of S to a Ni surface as a function of $C_s$ and combined this with Eqn. (2) to express the relationship between initial sulfur content and the hydrogen annealing conditions required to lower the sulfur content to a level that will result in some number of monolayers of sulfur, $N_m$, during saturation of the surface:

$$C_{s,i} = (8.27 \times 10^{-2} \text{gm/cm}^2)^* (N_m \text{A/W}) * (2/\sigma) * \exp(x^2D_A/\sigma^2)$$

where: $C_{s,i}$ is expressed in ppmw.

$A$ is the specimen surface area in cm$^2$.

$W$ is the specimen mass in gm.

Taking one monolayer ($N_m = 1$) as the amount of segregation necessary to degrade adherence allows calculation of the annealing time required at 1200°C to adequately desulfurize the specimens. These times are 2.6 h. for the 0.3 mm thick specimen and 492 h. for the 3.0 mm thick specimen. These times indicate that substantially longer hydrogen annealing would be required for the thicker specimen while the 0.3 mm specimen would be adequately desulfurized after 100 h. However, it is interesting to note that the 0.3 mm thick specimen of 1480 still did not have the cyclic oxidation resistance exhibited by the 0.3 mm thick specimen of hydrogen-annealed 1484 which was doped with Hf. A similar result has also been reported for CMSX-4 which is also doped with Hf [20]. This suggests that the removal of sulfur makes the Hf more effective. The unannealed 1487 which contained Hf+Y also showed better cyclic oxidation resistance than the annealed 1480. This result suggests Y is more effective than Hf as a “sulfur getter” or may provide some benefit to alumina adherence beyond sulfur-gettering and that optimum performance may be achieved by combining desulfurization and reactive-element doping.

**Polycrystalline Alloys**

The starting microstructures of the alloys with the varying sulfur levels were similar. They exhibited a dendritic $\gamma+\gamma'$ microstructure with a grain size varying from 50 to 100μm. Small amounts of a Ta-rich phase, which was not present in single crystal 1484, were identified along the grain boundaries by EDS.

The isothermal oxidation rates of the single crystal and polycrystalline alloys were all similar at 1100°C yielding parabolic rate constants on the order of $5 \times 10^{-14} \text{g/m}^4 \text{s}$. However, on cooling after 168 hours of oxidation the scale spalled completely from the 45 ppm S alloy and spalled partially from the 8 ppm S single crystal. The scale remained completely adherent to the 1 and 3 ppm S alloys. The surface and cross-section of the 3 ppm S alloy are shown in Figure 3. Transient oxidation resulted in thicker oxide on certain grains and dendrites (e.g. dark grains in the SEM surface micrograph) as opposed to others (light grains). Bright spots in the SEM micrograph of the surface were Ta-rich oxides, as indicated by EDS. These apparently formed by the rapid oxidation of the Ta-containing phase on the grain boundaries. The cross-section indicates an adherent oxide in contact with an Al-depleted layer in the alloy along a wavy interface. The Al-depleted layer, which was observed for all the alloys, was indicated by XRD to contain only $\gamma$. X-ray diffraction and EDS indicated that the scale consisted of continuous $\alpha-\text{Al}_2\text{O}_3$ covered with a layer containing NiO, Cr$_2$O$_3$, Ni(Al,Cr)$_2$O$_4$, and NiTa$_2$O$_6$ for all the alloys. The surface and cross-section of the 45 ppm S alloy are presented in Figure 4. The oxide has spalled from a large fraction of the specimen area during cooling and spalled oxide fragments are seen lying on the surface. The cross-section shows that the scale is separated from the alloy even in areas where it has not completely spalled.
Figure 5 shows the results of cyclic oxidation experiments on 1484 alloys with various sulfur contents at 1100°C. These data show that, for this alloy, there is a critical sulfur content between 3 and 8 ppm below which adherent scales are formed. Comparison with the data for PWA 1480, in Figure 2, indicates that either the 0.1 wt% Hf in the polycrystalline specimens containing 1 and 3 ppm S lowers the content of sulfur which is free to diffuse to the alloy/oxide interface to below 0.1 ppm, or it provides a beneficial effect in addition to sulfur-gettering. Furthermore, the data for PWA 1487 (containing approximately 10 ppm S), in Figure 2, indicate that the combined presence of 0.1 wt% Hf and 0.1 wt% Y is able to mitigate the effects of higher sulfur contents. These results are significant in that they indicate that:

- the sulfur content must be reduced below 0.1 ppm if long-term scale adherence is to be achieved without the use of reactive elements and
- that, when reactive elements are used, the amount of sulfur which can be tolerated varies with the amount and type of reactive element used.

These two points will be discussed more quantitatively in the section on mechanisms.

Figure 5: Cyclic oxidation kinetics in air at 1100 °C (1 hour cycles) for polycrystalline alloys with different sulfur levels and desulfurized 1484 (< 0.1 ppm S).

Residual stress in the α-alumina layer underneath the transient oxide layer on a polycrystalline alloy containing 1 ppm sulfur after 24 hours oxidation at 1100°C in air was measured by XRD at room temperature. Figure 6 shows the oxide scale consisting of an α-alumina layer (2 μm thick) between a transient oxide layer and an Al-depleted substrate (г) layer. The oxide layer was adherent to the substrate after cooling to room temperature (there was no noticeable oxide spallation).

Figure 6: SEM (BEI) cross-section micrograph of a polycrystalline alloy (1 ppm sulfur) after isothermal oxidation at 1100 °C in air for 24 hours.

Assuming a biaxial (an isotropic) stress-state ($\sigma_{11} = \sigma, \sigma_{22} = \sigma, \sigma_{33} = 0, \sigma_{12} = \sigma_{23} = \sigma_{13} = 0$) and a homogeneous elastic medium, the strain in a
Residual stresses measured in the current study are almost equal to the calculated thermal stresses suggesting minor growth stresses or relaxation of the growth stresses. The alloy containing 1 ppm sulfur withstood high residual stresses during cooling indicating that the alumina/ alloy interface is inherently strong (adherent) while the alloy containing 8 ppm and 45 ppm sulfur showed extensive spallation during cooling indicating that sulfur weakened an otherwise strong interface. These results support the mechanism [9,10] by which the adherence of the alumina to the substrate is improved by addition of reactive elements such as Hf, and Y.

**Mass Change/Area (mg/cm²)**

<table>
<thead>
<tr>
<th>Mass Change/Area (mg/cm²)</th>
<th>As Received</th>
<th>H₂ Annealed</th>
</tr>
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<tbody>
<tr>
<td>0</td>
<td>-2</td>
<td>-2.5</td>
</tr>
<tr>
<td>100</td>
<td>-2</td>
<td>-3.4</td>
</tr>
<tr>
<td>500</td>
<td>-2</td>
<td>-4.8</td>
</tr>
<tr>
<td>1000</td>
<td>-2.5</td>
<td>-6.2</td>
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<tr>
<td>1500</td>
<td>-3</td>
<td>-7.6</td>
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<tr>
<td>2000</td>
<td>-3.5</td>
<td>-9.1</td>
</tr>
<tr>
<td>2500</td>
<td>-4</td>
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<tr>
<td>3000</td>
<td>-4.5</td>
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Figure 8: Cyclic oxidation kinetics in air at 1100 °C (1 hour cycles) for as-received and hydrogen annealed NiAl.

**NiAl**

Figure 8 shows the effect of desulfurization on the cyclic oxidation of NiAl. The as-received specimen contained 20 ppm S and exhibited extensive scale spallation. The sulfur content of the hydrogen-annealed specimen was estimated from Eqn. (2) to be on the order of 0.1 ppm and this alloy exhibited a substantial improvement in cyclic oxidation resistance.

**Figure 10:** SEM (SEI) surface micrograph of as-received NiAl isothermally oxidized at 1100 °C in air for one week illustrating void formation.
Figure 9: Polarized light optical surface micrographs of NiAl isothermally oxidized for 48 hours at 1100 °C. A. hydrogen annealed. B. as-received.

The hydrogen annealing also affected the morphology of the alloy/oxide interface. Figure 9 shows polarized light micrographs of the surfaces of as-received and hydrogen-annealed NiAl after isothermal oxidation at 1100°C for 48 hours. The bright areas are voids beneath the α-Al2O3 scale at the alloy/oxide interface. Voids have coalesced along grain boundaries in the NiAl in both specimens but there is profuse void formation over the grains in the as-received specimen whereas there are very few for the hydrogen annealed specimen. Figure 10 shows more detail of the voids in a spalled area of the as-received material after isothermal oxidation at 1100°C for one week. It is clear that most of the surface was separated from the scale by voids. Figure 11 shows one of the few spalled regions on the hydrogen-annealed specimen after the same oxidation treatment. The trench that formed on a grain boundary is evident, however, there are only a few voids within the adjacent grains. Most of the depressions on the surface of the grains are the imprints of oxide grains. The above results indicate that sulfur has detrimental effects on oxide adherence to NiAl, as it does to superalloys, and that control of sulfur in diffusion aluminide coatings can be expected to increase coating life.

Mechanisms

The above results clearly indicate that removing sulfur improves the adherence of the alumina scales to both Ni-base superalloys and NiAl. This also supports the proposals in previous studies [9-12,17,18,20] that the improvements in scale adherence produced by reactive element additions, e.g. Figs. 2 and 5, are, at the very least, partly the result of sulfur being combined with the reactive element so that it cannot diffuse to the alloy oxide interface or enter the scale. However, the question remains as to how the sulfur actually degrades oxide adherence. Hou and Sirringer [26] report Auger electron spectroscopy observations of sulfur segregation at intact FeCrAl/alumina interfaces and suggest that sulfur weakens the bond between the alloy and oxide. Grabke et al [27] argue that sulfur can only segregate to an alloy/gas interface and that, since sulfur lowers the surface free energy, it causes voids to grow along the alloy/oxide interface. No measurements of sulfur segregation were made in the present study but Figure 10 clearly shows that voids separated much of the scale from NiAl containing 20 ppm S whereas there are relatively few voids on desulfurized NiAl, Fig. 11, except at grain boundaries. These observations are more consistent with the void growth mechanism. However, the fact that desulfurization prevents voids, means that reactive elements acting as 'vacancy sinks' [2,3] are not necessarily required to suppress interface voids. As long as sulfur is not free (or available) to diffuse to the alloy/oxide interface other sites in the alloy, such as grain boundaries and dislocations, become more effective sinks than the alloy/oxide interface.

The work of Smialek et. al. [17, 18] has pointed to a critical sulfur content which must be achieved by desulfurization in order to achieve good adherence. If the major effect of reactive element additions is to tie up sulfur the same concept should apply to doped alloys. The effect of sulfide stability on surface adsorption of sulfur has been analyzed by Luthra and Briant [25] for binary M-S alloys and they indicated how their treatment could be extended to reactive element-doped alloys. However, the activity coefficients and surface free energies required for the calculations are generally not available. The analysis used here will extrapolate existing solubility product data to indicate how reactive element additions affect the amount of sulfur available for transport to the alloy/oxide interface.
The equilibrium between a reactive element sulfide (MS) and the elements in solution in the alloy is shown schematically in Figure 12 and may be represented by

$$MS(s) = M + S$$  \hspace{1cm} (5)

If the activity of the sulfide is taken as unity the activity coefficients of the elements in solution (taken here as independent of concentration) can be combined with the equilibrium constant for Eqn. 5 to yield a solubility product for MS.

$$K_{sp} = C_{M}^{\text{Residual}} C_{S}^{\text{Residual}}$$  \hspace{1cm} (6)

Here $C_{M}^{\text{Residual}}$ refers to the concentration of reactive element (in at% or wt.%) left in solution and will be given by

$$C_{M}^{\text{Residual}} = C_{M} - C_{M}^{\text{MS}}$$  \hspace{1cm} (7)

where $C_{M}$ is the total reactive element content of the alloy and $C_{M}^{\text{MS}}$ is the amount of reactive element tied up in the sulfide. Similarly,

$$C_{S}^{\text{Residual}} = C_{S}^{0} - C_{S}^{\text{MS}}$$  \hspace{1cm} (8)

where $C_{S}^{0}$ is the total sulfur content of the alloy and $C_{S}^{\text{MS}}$ is the amount of sulfur tied up in the sulfide. Note that $C_{M}^{\text{Residual}} = C_{M}^{\text{MS}}$.

Solubility products for sulfides in the alloys under study are not available. However, the use of representative values allows trends to be determined as indicated in Figure 13. (Note that sulfur concentrations have been converted to wt.%%.) A $K_{sp}$ value of 1.4x10^5(at%)^2 is reported for TiS at 1100°C in austenitic Fe [28] (which might approximate the solubility of TiS in a superalloy). The value of 10^10(at%)^2 might approximate the dissolution of YS which is a much more stable sulfide than TiS. The other values were chosen to show the effect of varying the solubility product. The trends indicated in Figure 13 and 14 are:

i. The residual sulfur content decreases with the amount of reactive element added.

ii. The residual sulfur content decreases with increased stability of MS i.e. a decrease in $K_{sp}$.

iii. The residual sulfur content is essentially independent of the initial sulfur content between two critical values of $C_{S}^{0}$.

The two critical values of $C_{S}^{0}$ mentioned in item (iii) are a low value, below which the sulfur activity is too low to form a sulfide, and a high...
value, above which there is more than the stoichiometric amount of sulfur to react with all the reactive element present.

\[ K_{SP} = 10^{-6} \text{ (wt%)}^2 \]

\[ C_s^0 \text{ residual (wt.%)} \]

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\[ C_s^0 \text{ residual} \]
The improvement in the cyclic oxidation resistance of PWA 1484 by lowering the initial sulfur content is qualitatively consistent with the above calculations. Similarly, the improvement in Hf-doped superalloys by hydrogen-annealing, PWA 1484 in Fig. 2 and CMSX-4 in Ref. 20, may be expected from an effective lowering of \( C_s^0 \) by the anneal. However, Figure 14 indicates that the residual sulfur content, which would stand in equilibrium with “HfS”, would not be lowered much by reducing \( C_s^0 \). The quantity which is lowered is \( C_s^\text{res} \), i.e. the amount of sulfide in the alloy. Therefore, the improvements in PWA 1484 and CMSX-4 may result from decreasing the amount of sulfur which can be released from sulfide as the alloy-oxide interface encroaches and the Hf is oxidized. Also, as illustrated in Figure 15, the value of \( C_s^\text{res} \) increases continuously as \( C_s^0 \) decreases. Therefore, lowering the initial sulfur content of PWA 1484 will leave more Hf in solution to tie up sulfur that is released during oxidation and to tie up other potentially detrimental impurities.

This approximate analysis is deficient in that accurate values for the solubility products are not known. For example, the results in Figure 5 would tend to indicate that the steep portion in the top diagram of Figure 13 and 14 should occur at less than \( C_s^0 = 8 \) ppm i.e. the solubility product of hafnium sulfide in PWA 1484 is actually more than an order of magnitude greater than the one used in Figure 14. However, it does indicate that evaluation of the sulfur-gettering capability of a given reactive element addition cannot be accomplished by a simple mass balance. The magnitude of \( K_{sp} \) and how the equilibrium may be affected by other reactive components, particularly oxygen and carbon, must be considered along with the amounts of sulfur present and reactive element added. This type of analysis has also been shown to be qualitatively explain the effects of reactive elements and sulfur on the cyclic oxidation behavior of alumina-forming FeCrAl alloys [31].

**Liquid-Phase Desulfurization**

The above results indicate the improvement in alumina adherence which can be achieved by removing sulfur and the need to reduce the sulfur content to the range of 1 ppm or lower to take full advantage of this approach. Although sulfur can be removed from solid alloys by annealing in purified hydrogen, this technique is both slow and inefficient. Recently an experimental program has been initiated to attempt to remove sulfur in nickel and nickel-based alloys to below 1 ppm in the liquid state. If successful, this will allow economical removal of sulfur from commercial sized heats.

The technique used is to melt 10 lb. of nickel in a vacuum induction furnace and allow the sulfur in the nickel to react with a reactive lining. The reactions will be of the type:

\[
\begin{align*}
MO + S &= MS + \text{Q} \\
MO + S + C &= MS + CO \\
M_2O_3 + S &= M_2S + CO \\
M_2O_3 + S + C &= M_2O_2S + CO
\end{align*}
\]

Under vacuum, the presence of carbon proceeding to form carbon monoxide will tend to push the reactions to the right. The presence of aluminum in the alloy would provide a similar impetus but would result in an oxide inclusion being formed. Thermodynamic calculations considering various active linings, such as CaO and Y_2O_3, indicate that it should be possible to remove the sulfur to well below the 1 ppm level.

Experiments carried out so far indicate that sulfur is maintained at 1 ppm and that the mechanism appears to proceed to produce calcium metal which bubbles through the liquid alloy, as indicated by the following reaction.

\[
\text{CaO} + \text{C} = \text{Ca(g)} + \text{CO}
\]

The reaction with sulfur in this case will involve

\[
\text{Ca(g)} + S = \text{CaS}
\]

Experiments have not been carried out using Y_2O_3 linings and currently the vacuum furnace is being modified to allow metal samples for analysis to be taken during the run. This will allow mechanisms, kinetics, and limiting sulfur levels to be determined.

**Summary And Conclusions**

The results of this study have shown that desulfurization of alumina forming Ni-base superalloys and NiAl by hydrogen annealing can result in improvements in cyclic oxidation comparable to that achieved by doping with reactive elements.

It is possible to estimate the amount of sulfur available to segregate to the alloy-oxide interface and how this is influenced by reactive element additions or hydrogen annealing. If these calculations can be made more quantitative it should be possible to engineer alumina-forming alloys for optimum resistance to cyclic oxidation. Preliminary experiments to desulfurize Ni-base alloys in the liquid phase have been discussed.
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References

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