Long-Term Cyclic Oxidation Behavior of Uncoated and Coated Re108 and In939 at 980 and 870 °C

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LONG-TERM CYCLIC OXIDATION BEHAVIOR OF UNCOATED AND COATED Re108 AND In939 AT 980 AND 870 °C

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SUMMARY

Very long-term cyclic oxidation behavior of Re108 and In939 with and without a protective coating was evaluated at 980 and 870°C, respectively. Re108 and In939 without a protective coating began to show rapid weight loss at 3000 h due to scale spallation, indicating the need for an oxidation protective coating for longer than thousands of hours of oxidative life. NiAl-base coatings of a vapor phase aluminide (VPA), a pack aluminide (CODEP), and a slurry paint aluminide (SERMALOY J) were applied on Re108 and In939. VPA and CODEP on Re108 and all three coatings on In939 showed excellent cyclic oxidation resistance out to 10 000 hr. Coated alloys were annealed in an inert atmosphere to determine the loss of Al from the coating into the alloy substrate through diffusion. The Al loss from the coating through diffusion was twice as great as the Al loss through oxidation after 10000 h of cyclic exposure. Oxidation life of VPA-coated Re108 was estimated by calculating the amount of Al initially available for protective oxidation and the amount of Al lost through oxidation and diffusion.

INTRODUCTION

Oxidation protection of high temperature alloys relies on the formation of an external layer of stable, slow growing oxides, such as Cr₂O₃, Al₂O₃, or SiO₂, through the selective oxidation of Cr, Al, or Si, respectively (ref. 1). There is a critical value for each of these elements which is necessary to form a continuous external oxide layer of respective element in isothermal oxidation (ref. 1). Therefore, the protection conferred by Al₂O₃, for example, ends once the selective oxidation of Al reduces the Al content in the alloy below the critical value. Consequently, the oxidative lifetime of an alumina-forming alloy at a given temperature can be estimated by calculating the amount of Al initially available for protective oxidation and the loss of Al through oxidation, assuming rapid diffusional kinetics in the alloy (refs. 2 and 3).

Enhanced efficiency and performance of gas turbine engines requires hot section structural components with higher temperature capability and longer life. In a typical alumina-forming high temperature superalloy, there is only a limited amount of excess Al above the critical value because too much of Al degrades the mechanical strength of superalloys. Under thermal cycling, protective scales crack and spall due to the difference in thermal expansion between the alloy and the scale, accelerating the loss of Al compared with that in isothermal exposure. Thus, an external coating with a large reservoir of Al above the critical content is needed for a long cyclic oxidative life at elevated temperatures. In coated alloys, diffusion between the coating and the substrate is another source for the loss of the element responsible for the protective oxidation (ref. 4).

In this study, the long-term cyclic oxidation kinetics of alumina-forming Re108 and chromia-forming In939 with and without an external coating (NiAl-base coatings) were investigated. Coated alloys were also exposed in an inert atmosphere to investigate the chemical interdiffusion between the coating and the superalloy substrate. Oxidation life of coated Re108 was estimated on the basis of oxidation kinetics and diffusion data.
2. EXPERIMENTAL PROCEDURE

Oxidation coupons of Re108* and In939† (2.5 × 1.25 × 0.3 cm) were cut from a large plate of each respective alloy. The as-cast Re108 has a surface layer (~25 µm thick) depleted in Cr, W, and Co (ref. 5). The surface of a second set of coupons was ground off to remove the depleted surface layer. Some as-cast coupons were coated with a vapor phase aluminide (VPA),* a pack aluminide (CODEP)* and a slurry paint aluminide (SERMALOY J)*. VPA and CODEP were about 40 µm thick and SERMALOY J was about 60 µm thick. All three coatings are based on NiAl. Tables I and II, respectively, show the nominal composition of Re108 and In939 and the composition of the three coatings as determined in this study using electron microprobe analysis (EMPA). For the coating, the average composition of each element across the coating measured at 5 µm intervals is reported (table II). Uncoated coupons were polished to a 600-grit surface finish and all coupons were rinsed in acetone and methyl alcohol in an ultrasonic cleaner prior to the cyclic oxidation or chemical diffusion test.

Cyclic oxidation test was performed using an automated cyclic furnace (ref. 6). Each cycle consisted of 1 h at temperature and a minimum of 20 min at room temperature. Typically, samples reached the test temperature within 2 min and the ambient temperature within 5 min in each cycle. Oxidized coupons were weighed periodically using a microbalance (+0.02 mg) to determine the oxidation kinetics. Chemical diffusion was carried out in a horizontal furnace in a flowing Ar-5%H₂ to minimize the oxidation of coupons. Both cyclic oxidation and chemical diffusion tests were performed at 980 and 870 °C, respectively, for Re108 and In939.

X-ray diffraction (XRD) was used to determine the phases present in the oxide scale and the coating and scanning electron microscopy (SEM) equipped with energy dispersive spectroscopy (EDS) was used to examine the cross-sections of post-oxidation and post-diffusion coupons. EMPA was used to determine the compositional profile of various elements along the cross-section of post-diffusion coupons.

3. RESULTS AND DISCUSSION

3.1 Oxidation of Uncoated Re108 and In939

Figure 1 shows the oxidation kinetics of as-cast and ground Re108 after cyclic oxidation at 980 °C. Both coupons showed an initial parabolic oxidation followed by a constant weight period and eventually a weight loss after 3000 hr. Except for the slightly higher weight loss rate after 3000 hr in the ground coupon, the two coupons showed a similar oxidation behavior. Figure 2 shows the cross-section of Re108 after 4500 h. The oxide scale consists of a NiO top layer, an intermediate spinel layer, and an inner Al2O3 layer. There are also islands of HfO2 particles mostly within the spinel layer. X-ray diffraction on the Re108 coupon after 4500 h showed NiO, HfO2, spinel, Al2O3, and TiO2.

Figure 3 shows the oxidation kinetics of In939 after cyclic oxidation at 870 °C. The weight change behavior was very similar to that of Re108 at 980 °C, with the ground In939 showing a slightly higher weight loss rate. Figure 4 shows the cross-section of In939 after 4500 h. The oxide scale consists of a NiO top layer, an intermediate spinel layer, and an inner Cr2O3 layer, similar to Re108 except that the inner Al2O3 was replaced by Cr2O3. X-ray diffraction showed NiO, spinel, Cr2O3, and TiO2.

The kinetics data shown in figures 1 and 3 are typical of alumina- or chromia-forming superalloys under cyclic oxidation, i.e., initial weight gain followed by a weight loss due to scale spallation (paralinear oxidation). The rapid weight loss rate of Re108 and In939 indicates the need for protective coatings for an oxidation life beyond thousands of hours.

3.2. Oxidation of Coated Re108 and In939

NiAl-base coatings of VPA, CODEP and SERMALOY J were applied on Re108 and In939 for extended oxidation protection. Coating composition determined by EMPA in this study is shown in table II.

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*Supplied by GE Aircraft Engines, Cincinnati, Ohio.
†Supplied by Pratt & Whitney, West Palm Beach, Florida.
**Re108.**—Figure 5 shows the oxidation kinetics of coated Re108 after cyclic oxidation at 980 °C. The two sets of data for CODEP and VPA are replicates. CODEP and VPA showed a typical parilinear weight change of alumina-forming alloys under cyclic oxidation. No significant scale spallation was observed during the weight gain period, indicating that the weight change in this period was predominantly due to the formation of alumina scale. SERMALOY J gained substantially higher weight than VPA or CODEP when the test was terminated at 3500 h. A significant amount of spinel was observed in the scale of SERMALOY J coating, which presumably was responsible for the higher weight gain of this coating. The transition from weight gain to weight loss occurred at around 4000 hr for VPA and CODEP. The weight loss was fairly linear out to 10000 h (fig. 6).

Figure 7 shows the cross-section of VPA-coated Re108 after 1000 h. The scale was mostly Al2O3 with some isolated patches of HfO2. The brightest phase in the coating was rich in Hf and Ta, from which HfO2 formed. The bright phase in the coating was rich in W and Cr (fig. 7(b)). The diffusion zone contained significant amounts of Hf/Ta- and W/Cr-rich phases. Below the diffusion zone was a zone rich in aluminum, which contained a needle-shaped second phase rich in W and Cr (fig. 7(a)).

**In939.**—Figure 8 shows the oxidation kinetics of coated In939 after cyclic oxidation at 870 °C. All three coatings showed an initial rapid weight gain (transient oxidation) followed by a very slow, steady-state weight gain. SERMALOY J and VPA showed higher initial weight gains than CODEP; however, the steady state oxidation rate was similar in all three coatings. Note that the weight gain of SERMALOY J-coated In939 at 870 °C was significantly lower than that of SERMALOY J-coated Re108 at 980 °C (fig. 5). The oxide scale of SERMALOY J coating contained a smaller amount of spinel at the lower temperature.

### 3.3. Chemical Diffusion

**VPA-Coated Re108.**—Figures 9(a) and (b) show the concentration profiles of various elements along the coating/diffusion zone/substrate after 0 and 9000 h annealing, respectively, at 980 °C. Two points are worth mentioning. After the heat treatment, concentration profiles of all elements became fairly flat and the diffusion zone extended significantly. The flattening of elemental concentration profiles occurred within the first 1000 h annealing. Figure 10 shows the plot of Al concentration in the coating versus annealing time for VPA-coated Re108. Note the rapid drop of Al concentration within the first 1000 h after which it leveled off at 20–21 wt%, indicating the initial rapid diffusion of Al out of the coating. Figures 11 and 12 compare the cross-sections of as-processed and annealed (9000 h) VPA-coated Re108. As-processed VPA coating was β phase (fig. 11), while γ second phase appeared after annealing (fig. 12(a)). The diffusion zone contained two second phases, i.e., Hf/Ta-rich phase (brightest) and W/Cr-rich phase (bright). Note that the second phases significantly coarsened with annealing (fig. 12(b)). An Al-rich zone, containing a high concentration of a needle-shaped phase, developed below the diffusion zone (fig. 12(a)). This second phase was rich in W and Cr.

**SERMALOY J-Coated In939.**—Figures 13(a) and (b) show the concentration profiles of various elements along the coating/diffusion zone/substrate after 0 and 9000 h annealing, respectively, at 870 °C. Similar behavior to the VPA coating was observed, i.e., the concentration profiles of all elements became fairly flat after annealing and the diffusion zone thickness increased significantly. Figures 14 and 15 compare the cross-sections of as-processed and annealed (9000 h) SERMALOY J-coated In939. A high concentration of a second phase rich in Cr, Co and W precipitated in the coating after annealing (fig. 15). The diffusion zone, after annealing, consisted of at least three phases (fig. 15), the detailed analysis of which was not attempted in this study. Similar to VPA, the second phase below the diffusion zone was needle shaped (fig. 15). Figure 10 shows the plot of Al concentration in the coating versus annealing time for SERMALOY-coated In939. Similar to VPA, rapid diffusion of Al occurred within the first 1000 h, after which it leveled off at 18–19 wt%.

### 3.4. Oxidation Life Projection of VPA-Coated Re108

Oxidation life of an alumina-forming alloy ends when the Al concentration in the alloy drops below the critical Al concentration which is necessary for the formation of a continuous, protective Al2O3 scale (ref. 1). In other words, oxidation life ends when all the Al available (excess Al above the critical value) is lost. There are two sources for the loss of Al, i.e., oxidation to form Al2O3 scale and diffusion between the coating and the substrate. Since the scale formed on VPA was mostly Al2O3, the loss of Al through oxidation was determined from the oxidation weight change. The Al loss during the weight gain period was determined from the stoichiometry of the reaction of Al with O2 to form Al2O3 scale, assuming that scale spallation was negligible during this period. The Al loss rate during the weight loss period was taken from the average slope of the
weight loss curve, and was assumed to be applicable to the end of oxidation life. The projected oxidative lifetime of bulk NiAl using this assumption agreed fairly well with the lifetime determined by experiments (ref. 2). The change in Al concentration with annealing (fig. 10) was used to determine the loss of Al through diffusion.

Calculation of Al Available

Consider a unit volume (1 x 1 x h cm; h = coating thickness) of VPA coating. The fraction of Al initially available for protective oxidation is related to the initial fraction of Al and the critical Al concentration according to equation (1).

\[ f_{Al,\text{avail}} = f_{Al}^0 - f_{Al}^* \]  

\( f_{Al,\text{avail}} \) weight fraction of Al initially available for protective oxidation  
\( f_{Al}^0 \) initial weight fraction of Al  
\( f_{Al}^* \) weight fraction of Al at the time of oxidative failure

Thus, the weight of Al initially available for protective oxidation in the unit area of coating is given by

\[ W_{Al,\text{avail}} = \left( f_{Al}^0 - f_{Al}^* \right) \rho \cdot h \text{ mg/cm}^2 \]  

\( \rho \) coating density (mg/cm\(^2\))  
\( h \) coating thickness (cm)

Calculation of Al Lost

**Al Loss during the Weight Gain Period.**—The Al loss during the weight gain period is calculated from the parabolic oxidation equation, assuming that the scale spallation was negligible during this period. The parabolic rate constant was determined from the slope of (weight gain)\(^2\) versus time plot. The loss of Al through spallation needs to be taken into consideration for more rigorous calculation of Al loss during the weight gain period of a cyclic oxidation (ref. 7).

\[ \left( W_p \right)^2 = k_p \cdot t \]  

\( W_p \) weight gain by parabolic oxidation  
\( k_p \) parabolic rate constant  
\( t \) oxidation time

Rearranging equation (3) and taking the stoichiometry of alumina formation reaction into consideration, the loss of Al during the weight gain period is given by

\[ W_{Al,p} = 1.125 \left( k_p t \right)^{0.5} \text{ mg/cm}^2 \ (t < t^*) \]  

\( t^* \) transition time from weight gain to linear weight loss period.

**Al Loss during the linear weight loss period.**—During the weight loss period, the rate of Al loss is taken from the average slope of the weight loss curve, and is assumed to be applicable to the end of oxidation life. Thus, the loss of Al during this period is given by

\[ W_{Al,l} = k_l (t - t^*) \text{ mg/cm}^2 \ (t > t^*) \]  

\( k_l \) average weight loss rate (mg/cm\(^2\) h)
**Loss of Al through diffusion.**—Two key assumptions in this calculation are that the loss of Al through diffusion is negligible at \( t > 1000 \) h (based on the data in fig. 10) and the coating density is constant throughout the coating life. Although the coating density undoubtedly will change to some degree as the coating composition changes, this assumption may be justifiable considering the approximate nature of life projection in this study. Thus, the weight fraction of Al in the coating after diffusion anneal in an inert atmosphere is given by

\[
f_{\text{Al}}^{**} = f_{\text{Al}} - f_{\text{Al,d}}
\]  

- \( f_{\text{Al}}^{**} \): weight fraction of Al in the coating after diffusion anneal in inert atmosphere
- \( f_{\text{Al,d}} \): weight fraction of Al lost by diffusion

Rearranging equation (6), the weight fraction of Al lost by diffusion is given by

\[
f_{\text{Al,d}} = f_{\text{Al}}^{0} - f_{\text{Al}}^{**}
\]  

Thus, the weight of Al lost by diffusion from the unit area of coating is given by

\[
W_{\text{Al,d}} = \left( f_{\text{Al}}^{0} - f_{\text{Al}}^{**} \right) \rho \cdot h \text{ mg/cm}^2 (>1000 \text{ h})
\]  

**Oxidation Life Projection**

Oxidation life ends when the available Al, \( W_{\text{Al,avail}} \) (eq. (2)), equals the Al lost, i.e., \( W_{\text{Al,p}} \) (eq. (4)) + \( W_{\text{Al,i}} \) (eq. (5)) + \( W_{\text{Al,d}} \) (eq. (8)). In other words, the oxidation life ends when the following condition is met.

\[
(f_{\text{Al}}^{0} - f_{\text{Al}}^{**}) \rho \cdot h = 1.125(k_{p,t})^{0.5} + k_{1}(t - t^{*}) + (f_{\text{Al}}^{0} - f_{\text{Al}}^{**}) \rho \cdot h
\]  

All parameters in equation (9) are known except for \( f_{\text{Al}}^{**} \). Table III lists the values of the parameters in equation (9). Nesbitt, et al. reported that the critical Al concentration in the cyclic oxidation of NiAl at 1200 to 1400 °C was ~20 wt% (ref. 2). As will be shown in the following discussion, the critical Al concentration in the cyclic oxidation of VPA coating in this study was less than 14 wt%, indicating the dependence of the critical Al concentration on alloy composition. Using equation (9), the oxidation life, \( t_{\text{life}} \), can be plotted as a function of the critical Al concentration, \( f_{\text{Al}}^{**} \) (fig. 16). If the critical Al concentration were 14 wt%, the coating would fail at \( t = 10000 \) h. This indicates that the aluminum concentration in the coating at \( t = 10000 \) h was 14 wt%, which is a drop of ~18 wt% from the initial concentration (~32 wt%). (Fig. 16 can be interpreted as a plot of oxidation time versus Al concentration in the coating.) Since about 11.5 wt% was lost through diffusion (fig. 10), the remainder (~6.5 wt%) was lost through oxidation, or about half of what was lost through diffusion. This demonstrates the importance of Al loss through diffusion in the consideration of the oxidation lifetime of coated alloys. The critical Al concentration of the VPA coating used in this study needs to be determined to project the cyclic oxidation life using equation (9). One way is to perform an accelerated oxidation at higher temperature until the coated coupon fails then analyze the Al content in the coating using EMPA.

### 4. CONCLUSIONS

- Re108 and In939 need a protective coating for cyclic oxidation life longer than thousands of hours at \( T > 980 \) and \( 870 \) °C, respectively.
- VPA and CODEP coatings on Re108 and VPA, CODEP, and SERMALOY coatings on In939 showed excellent oxidation resistance out to 10000 h at 980 and \( 870 \) °C, respectively.
- A substantial amount of Al was lost from the coating into the substrate through diffusion. In the case of VPA-coated Re108, the Al loss through diffusion was twice as great as the Al loss through oxidation after 10000 h cyclic exposure at 980 °C.
- The critical Al concentration of VPA coating needs to be determined to project the oxidation life of VPA-coated Re108.
REFERENCES

5. Jon Schaeffer: GE Aircraft Engines, unpublished research.

**TABLE I.** — NOMINAL COMPOSITION OF Re108 and In939 (wt%)

<table>
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<tr>
<th>Alloy</th>
<th>Ni</th>
<th>Al</th>
<th>Cr</th>
<th>Co</th>
<th>W</th>
<th>Ta</th>
<th>Ti</th>
<th>Hf</th>
<th>Mo</th>
<th>Fe</th>
<th>Si</th>
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<td>Re108</td>
<td>Balance</td>
<td>57</td>
<td>85</td>
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<td>85</td>
<td>3.88</td>
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<td>0.55</td>
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<td>0.03</td>
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<td>In939</td>
<td>Balance</td>
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<td>22</td>
<td>19</td>
<td>2</td>
<td>1.5</td>
<td>3.6</td>
<td>--</td>
<td>--</td>
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**TABLE II.** — COMPOSITION OF VPA, CODEP AND SERMALOY COATINGS DETERMINED IN THIS STUDY BY EMPA (WT%)

<table>
<thead>
<tr>
<th>Coating</th>
<th>Ni</th>
<th>Al</th>
<th>Cr</th>
<th>Co</th>
<th>Si</th>
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</thead>
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<tr>
<td>VPA</td>
<td>Balance</td>
<td>31.29</td>
<td>1.79</td>
<td>6.41</td>
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<tr>
<td>CODEP</td>
<td>Balance</td>
<td>35.45</td>
<td>1.47</td>
<td>5.67</td>
<td>--</td>
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<tr>
<td>SERMALOY J</td>
<td>Balance</td>
<td>27.43</td>
<td>15.74</td>
<td>13.54</td>
<td>7.04</td>
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**TABLE III.** — THE VALUES OF THE PARAMETERS IN EQUATION 9

<table>
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<tr>
<th>$f_a$</th>
<th>$f_{a}^{**}$</th>
<th>p ($mg/cm^2$)</th>
<th>l (μm)</th>
<th>t (hr)</th>
<th>$k_1$ ($mg/cm^2/hr$)</th>
<th>$k_2$ ($mg/cm^2/hr$)</th>
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<tbody>
<tr>
<td>0.32</td>
<td>0.20</td>
<td>6000</td>
<td>40</td>
<td>4000</td>
<td>0.00011</td>
<td>0.00012</td>
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</table>

NASA/TM—1999-209072
Figure 1.—Oxidation kinetics of Re108 at 980 °C with 1 hr cycle.

Figure 2.—Cross-section of Re 108 after 4500 hr at 980 °C with 1 hr cycle.
Figure 3.—Oxidation kinetics of In939 at 870 °C with 1 hr cycle.

Figure 4.—Cross-section of In939 after 4500 hr at 870 °C with 1 hr cycle.
Figure 5.—Oxidation kinetics of coated Re108 at 980 °C with 1 hr cycle.

Figure 6.—Linear weight loss rate of VPA-coated Re108 at 980 °C with 1 hr cycle.
Figure 7.—Cross-section of VPA-coated Re108 after 1000 hr at 980 °C with 1 hr cycle. (a) Low magnification, showing the coating and the diffusion zone. (b) High magnification showing the oxide scale.
Figure 8.—Oxidation kinetics of coated In939 at 870 °C with 1 hr cycle.
Figure 9.—Concentration profile of elements in VPA-coated Re108. (a) As-processed. (b) 9000 hr in Ar-5% H₂.
Figure 10.—Plot of Al concentration in the coating vs. the annealing time. (VPA: 980 °C, Sermaloy J: 870 °C)

Figure 11.—Cross-section of as-processed VPA-coated Re108.
Figure 12.—Cross-section of VPA-coated Re108 after 9000 hr in Ar-5% H₂ at 980 °C. (a) Low magnification, showing the coating and the diffusion zone. (b) High magnification, showing the diffusion zone.
Figure 13.—Concentration profile of elements in SERMALOY J-coated In939.
(a) As-processed. (b) 9000 hr in Ar-5% H₂.
Figure 14.—Cross-section of as-processed SERMALOY J-coated In939.

Figure 15.—Cross-section of SERMALOY J-coated In939 after 9000 hr in Ar-5% H₂ at 870 °C.
Figure 16.—Plot of oxidative life vs. critical Al content for VPA-coated Re108 at 980 °C with 1 hr cycle.
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