OXIDATION AND THE EFFECTS OF HIGH TEMPERATURE EXPOSURES ON NOTCHED FATIGUE LIFE OF AN ADVANCED POWDER METALLURGY DISK SUPERALLOY

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Abstract

Oxidation and the effects of high temperature exposures on notched fatigue life were considered for a powder metallurgy processed supersolus heat-treated ME3 disk superalloy. The isothermal static oxidation response at 704 °C, 760 °C, and 815 °C was consistent with other chromia forming nickel-based superalloys: a TiO₂-Cr₂O₃ external oxide formed with a branched Al₂O₃ internal subscale that extended into a recrystallized γ'-dissolution layer. These surface changes can potentially impact disk durability, making layer growth rates important. Growth of the external scales and γ'-dissolution layers followed a cubic rate law, while Al₂O₃ subcales followed a parabolic rate law. Cr-rich M₂₃C₆ carbides at the grain boundaries dissolved to help sustain Cr₂O₃ growth to depths about 12 times thicker than the scale.

The effect of prior exposures was examined through notched low cycle fatigue tests performed to failure in air at 704 °C. Prior exposures led to pronounced dehibs of up to 99 % in fatigue life, where fatigue life decreased inversely with exposure time. Exposures that produced roughly equivalent 1 µm thick external scales at the various isotherms showed statistically equivalent fatigue lives, establishing that surface damage drives fatigue debit, not exposure temperature. Fractographic evaluation indicated the failure mode for the pre-exposed specimens involved surface crack initiations that shifted with exposure from predominately single intergranular initiations with transgranular propagation to multi-initiations from the cracked external oxide with intergranular propagation. Weakened grain boundaries at the surface resulting from the M₂₃C₆ carbide dissolution are partially responsible for the intergranular cracking. Removing the scale and subscale while leaving a layer where M₂₃C₆ carbides were dissolved did not lead to a significant fatigue life improvement, however, also removing the M₂₃C₆ carbide dissolution layer led to nearly full recovery of life, with a transgranular initiation typical to that observed in unexposed specimens.

Introduction

Turbin disks are fabricated from polycrystalline γfcc/γ(L1₂) phase nickel-based superalloys for strength and fatigue resistance, with grain boundary carbides and borides to improve creep rupture strength. Advances in powder metallurgy alloys have produced disks with improved properties, allowing jet engines to run hotter for increased fuel efficiency and reduced emissions. However, increased operating temperatures desired for future, more efficient engines make these alloys prone to localized hot corrosion and oxidation attack [1-3], which has the potential to limit disk durability. Service conditions for disks in some aerospace and land-based gas turbine engines can produce major cycle periods extending from minutes to hours and days. Over a disk component’s life, this can produce total service times exceeding 1,000 hours for aerospace applications and 100,000 hours for land-based applications. Some of the effects of service environments can be captured by extended exposures at elevated temperatures prior to conventional LCF [4, 5].

The objective of this study was to examine oxidation at potential maximum service temperatures over extended periods, and its effects on notched fatigue life, for an advanced powder metallurgy disk superalloy ME3. Environmental attack by static oxidation of coarse-grained supersolus heat treated ME3 was examined thoroughly at temperatures of 704 °C (1300 °F) to 815 °C (1500 °F) for extended times. Notched fatigue specimens were exposed in air at several of these conditions. Specimens were then tested using conventional fatigue tests at 704 °C. The effects of these exposures on notched fatigue life, crack initiation and propagation are discussed.

Materials and Experimental Procedure

Advanced powder metallurgy disk superalloy, ME3, also sometimes referred to as Rene 104 [6, 7], was argon atomized, consolidated, hot compacted, extruded and isothermally forged into production scale disks [7, 8], approximately 61 cm diameter with a 10 cm maximum bore thickness and 5 cm rim thickness. ME3 has a nominal composition of Ni-20.6 Co-13 Cr-3.7 Ti-3.8 Mo-3.4 Al-2.4 Ta-2.1 W- 0.9 Nb wt %, bal. Ni and minor elements [9]. These disks were then supersolus solution heat treated, and given conventional stress relief and aging heat treatments [7, 8]. Low stress ground (RMS roughness = ~1 µm) coupons and square-cross section blanks, 15 mm x 15 mm x 54 mm in dimension, were extracted from the disk rims with comparable cooling rates.

The disk rim microstructure [7, 8] used to fabricate specimens consisted of a mean grain diameter that ranged at ASTM 7.1-6.8 (27.1-30.6 µm) with standard deviation of ASTM 0.2-0.4 (2-4 µm) and an ALA grain diameter of ASTM 105-125 µm; Fig. 1. The γ'-phase consisted of cuboidal secondary precipitates (0.1-0.6 µm diameter) and spherical tertiary precipitates (10-40 nm diameter), where measured area fractions of γ'-phase were about 0.5. Electron diffraction and qualitative energy dispersive x-ray ray established that (Ti,Ta,Nb,Mo)C carbides (0.15-0.70 µm) and (Mo,Cr,W)₂B₂ borides (0.4-1.0 µm) reside mostly within the grains, while smaller (Cr,Mo,W)₂C₆ carbides (<0.15 µm) ornament the grain boundaries. Also, a small fraction of ZrO₂ oxides and Al₂O₃ oxides were observed within the grains.
Cylindrical notched low cycle fatigue (NLCF) specimens with a circumferential notch were fabricated by an external vendor to an elastic stress concentration of K<sub>c=2</sub>. NLCF specimen notches were polished longitudinally to less than 0.41 µm RMS roughness. Fully machined NLCF specimens and two blanks were exposed at high temperature under ambient air in a conventional resistance furnace by: i) equilibrating at 400 °C for 10 h or longer, ii) ramp heating in 0.5 h to 25 °C below the exposure temperatures, iii) ramp heating in 0.5 h to the exposure temperatures, iv) holding isothermally up to 2,020 h. Specimens and blanks were then removed from the furnace and placed on a ceramic brick for air cool. For exposures in vacuum, specimens were wrapped with two layers of Ta foil to getter oxidation during exposure. A similar temperature profile to the air exposures, except specimens were furnace cooled. Several specimens pre-exposed at 815 °C for 2,020 h were repolished to remove 19-23 µm and 48-52 µm from the notch radius by the same external vendor. Notch surfaces of fatigue specimens were imaged with SEM prior to NLCF testing.

Notched fatigue tests at 704 °C were performed to failure using closed-loop servo-hydraulic testing machines. The load control testing used a triangular waveform having a maximum net section stress of 855 MPa (124 ksi), a stress ratio (R(s)=σ<sub>min</sub>/σ<sub>max</sub>) of 0.05, and a frequency of 0.333 Hz. Fracture surfaces of fatigue specimens were evaluated by SEM. Crack initiation origins and crack propagation failure modes were identified and characterized. Longitudinal metallographic cross sections oriented parallel to the loading axis were taken from the gage sections in the plane of the primary crack front, ± 20 µm, to investigate secondary cracking and other damage.

A goodness of fit parameter Q based on the incomplete gamma function was used to estimate the quality of the data fitting. All fits presented had values of Q > 0.1, indicating the models used had a strong goodness of fit. The error in data fitting is the standard deviation of Gaussian distribution, determined by the diagonal element of the covariance matrix.

**Results and Discussion**

ME3 static oxidation of flat coupons

Our oxidation studies of ME3 coupons focused on the static isothermal response at 704 °C, 760 °C and 815 °C. The oxidation behavior was found for the most part to be consistent with other high Cr content Ni-based disk alloys, Waspaloy, Astroloy, Udiment 720, RR1000 [1-4], where the external scale consisted of continuous Cr<sub>2</sub>O<sub>3</sub> (a.k.a. eskolite) and facetted TiO<sub>2</sub> (a.k.a. rutile) grains (Fig. 2). The TiO<sub>2</sub> grains, which may spall, were primarily located at the exposed surface and become more abundant with exposure time.

Cross section evaluation uncovered that complex near-surface damage results from these exposures. An internal subscale of branched Al<sub>2</sub>O<sub>3</sub> fingers was evident beneath the Cr<sub>2</sub>O<sub>3</sub>-TiO<sub>2</sub> scale in ME3, even at the shortest exposures. These Al<sub>2</sub>O<sub>3</sub> fingers extended into a volume of the alloy where γ' precipitates have been dissolved by Al depletion. As shown for the 815 °C for 440 h exposure in Fig. 2, the precipitate-free volume is recrystallized and contains isolated, spherical voids that were about 1 µm in diameter at the near-surface grain boundaries. These voids have been attributed by others to result from an accumulation of vacancies left by Cr and Ti diffusion [2]. Light contrast grains, present in the oxide, contained no O and were Ni and Co rich, suggesting these are recrystallized grains from the alloy which were captured during oxide growth. A sub-micron thick sub-layer of (Ta,Cr)O<sub>2</sub>, which images brightly and has a rutile compound structure, was found to line the oxide/metal interface. (Ta,Cr)O<sub>2</sub> also sometimes wets the Al<sub>2</sub>O<sub>3</sub> fingers.

Microprobe evaluation revealed that the oxidation process leads to chemical changes that extend beyond the near surface attack, Fig. 3. In the bulk, the M<sub>2</sub>C<sub>3</sub> carbides in the grain boundaries appear to be enriched in Cr, Mo, Co, which is consistent with observations that Co is a major component in M<sub>2</sub>C<sub>3</sub> carbides in a similar disk alloy, LSHR [11]. Towards the oxidized surface, the ME3 grain boundaries are depleted in Cr, Mo, Co and enriched in Ni relative to bulk grain boundaries as shown in Fig. 3a for a
accumulated underneath the external scale, within γ'-dissolution layer. Furthermore, the recrystallized γ'-dissolution layer is depleted in Al, Cr, Ti, and Ta due to the formation of the major oxides.

A comparison of the resulting oxides for several of the coupon exposures is shown in Fig. 4. The γ'-dissolution layer was best imaged with differential interference contrast (DIC) optical microscopy, which would resolve layer thicknesses to 1 μm. Even at the shortest times the external scale appeared to be continuous and fairly uniform in thickness, and Al₂O₃ fingers were present.

Table I compares the measurements for the extension of carbide dissolution with those for the oxide scale thickness, alumina finger penetration and thickness of γ'-dissolution layer at 2,020 h exposures. The M₂₃C₆ dissolution layers were 3 times greater than the thicknesses of the γ'-dissolution layer (where Al₂O₃ fingers reside), which were roughly 3 times greater than the external oxide scale thicknesses. These layer dimensions are additive. For example, for the most aggressive 815 °C at 2,020 h exposure imaged in Fig. 3b, a ~20 μm depth removal would ensure that both the external oxide and subscale would be fully removed, while a ~50 μm depth removal would remove material past the M₂₃C₆ carbide dissolution layer into the bulk alloy.

Table I. Oxide scale thickness, alumina finger penetration, γ'-dissolution layer thickness and the extension of carbide dissolution past γ'-dissolution at 2,020 hours with temperature.

<table>
<thead>
<tr>
<th>Temperature</th>
<th>Avg. Scale Thickness ± σ</th>
<th>Avg. Al₂O₃ Penetration ± σ</th>
<th>Avg. γ'-ppts. Dissolution ± σ</th>
<th>Mean M₂₃C₆ Dissolution ± SE</th>
<th>Technique (# values)</th>
</tr>
</thead>
<tbody>
<tr>
<td>704 °C</td>
<td>1.31 ± 0.41</td>
<td>2.97 ± 0.78</td>
<td>3.11 ± 1.03</td>
<td>9.6 ± 2.3</td>
<td>SEM (3840)</td>
</tr>
<tr>
<td>760 °C</td>
<td>1.96 ± 0.41</td>
<td>5.17 ± 0.74</td>
<td>5.80 ± 1.27</td>
<td>16.4 ± 2.4</td>
<td>SEM (24)</td>
</tr>
<tr>
<td>815 °C</td>
<td>3.73 ± 0.83</td>
<td>9.97 ± 1.23</td>
<td>10.58 ± 0.89</td>
<td>32.6 ± 4.2</td>
<td>Optical DIC (24)</td>
</tr>
<tr>
<td>815 °C</td>
<td>3.73 ± 0.83</td>
<td>9.97 ± 1.23</td>
<td>10.58 ± 0.89</td>
<td>32.6 ± 4.2</td>
<td>WDS EPMA (4)</td>
</tr>
</tbody>
</table>

1. σ is the standard deviation for the fitted distribution
2. M₂₃C₆ dissolution is measured to extend from the bottom of γ'-dissolution layer to the far-field saturation where the Cr gradient subsides.
3. SE is the standard error of the mean value.

Fig. 4. Surfaces of oxidized ME3 coupons, exposed at a) 704 °C for 100 h, b) 704 °C for 2,020 h, c) 815 °C for 100 h and d) 815 °C for 2,020 h.

specimen exposed at 815 °C at 2,020 h. Fig. 3b shows that the volume adjacent to the oxide is depleted in carbides/borides, which image brightly in back-scattered SEM due to their high refractory content. This M₂₃C₆ dissolution is accompanied by gradients in both Ni and Cr, Fig. 3c. The M₂₃C₆ carbide liquidation temperature for ME3 in equilibrium was predicted to be 880.7 °C, which is near the exposure temperatures utilized in this study, using microstructural modeling package JMatPro-Ni 6.0. Between 25 μm and 50 μm depths, there is still a good number of carbides/borides present. Further in, at roughly 50 μm depth, there is a clear transition from no/limited M₂₃C₆ coverage to full M₂₃C₆ coverage at the grain boundaries. Chemical mapping, Fig. 3a, shows Mo rich areas (i.e. Mo-rich minor phases) within the grains that are not rich in Cr, which suggests that the carbides/borides present are mostly the more stable MC carbides and M₃B₂ borides. For stable M₂₃C₆ carbides in equilibrium, the Cr content is nearly 2.5 times greater than the bulk concentration in at.% [11]. Near the surface, Cr from M₂₃C₆ carbides has diffused out to help sustain the Cr₂O₃ scale growth, while Mo and Co
The thickness of external scale, the depth of Al$_2$O$_3$ finger penetration, and the thickness of $\gamma'$-dissolution layer were consistently measured on the left-side, middle and right-side of Ni-plated cross sections for all exposures. The standard errors were determined by the variation in average values of the three areas. Oxide scale measurements were log-normally distributed, while the finger depths and $\gamma'$-dissolution layer thicknesses were normally distributed.

The isothermal oxidation reaction kinetics for the measured averages of all three quantities is displayed in Fig. 5. The trends were determined by fitting the data ($y$) with time ($t$) to a simple power law of $y = (k \cdot t)^n$, where $k$ is the growth rate constant and $n$ is the temporal exponent. The kinetics for external oxide scale thickness (OST) deviated from classic parabolic growth, $t^{1/2}$ and followed a cubic rate law, $t^{1/3}$, which may be attributed to significant grain growth within the Cr$_2$O$_3$ oxide [12]. In contrast, a parabolic rate law was determined for the average planar extension of the Al$_2$O$_3$ fingers (AFD) from the oxide/metal interface into the metal. The inward growth suggests that oxygen diffuses inward from the oxide/metal interface [13]. For 704 °C for 10 h and 100 h exposures, Fig. 5, the finger depths measured were noticeably smaller than the continuation of the trend from longer exposure times, likely a deviation associated with the earliest stages of Al$_2$O$_3$ growth. For these short exposures, $\gamma'$-precipitates in the near vicinity of the forming Al$_2$O$_3$ fingers are dissolved, yet the $\gamma'$-dissolution layer has not reached a critical thickness to recrystallize. Al$_2$O$_3$ appears to wet the oxide/metal interface as a very thin, semi-continuous layer, which is not observed for the more aggressive exposures. The $\gamma'$-dissolution layer (GPD) kinetics mimicked the cubic rate law of the oxide scale, Fig. 5. Average thickness of the $\gamma'$-dissolution layer was observed to be roughly 2-3 times greater than the oxide scale thicknesses.

Assuming either a cubic or parabolic rate holds, further analysis of the data in Fig. 5 is possible using an Arrhenius relationship of $k = k_0 \exp(-Q/RT)$, where $Q$ is an activation energy for growth, $R$ is the gas constant, $T$ is the exposure temperature (in Kelvin) and $k_0$ is the pre-exponential factor. Plotting $\ln k$ versus $1/T$ produced linear relationship with $Q/R$ that gives a slope $-Q/R$, where $Q$ is the activation energy for growth and $R$ is the gas constant.

Fig. 5. The average measured growth for the external scale thickness (top), Al$_2$O$_3$ finger length (middle) and $\gamma'$-dissolution layer thickness (bottom) for static oxidation of ME3 exposed in air at 704 °C, 760 °C and 815 °C up to 2,020 hours. The solid lines denote the power law fit, $y = (k \cdot t)^n$, while the dashed lines mark the continuation of the fit to shorter exposure times. The standard deviation of the geometric averages of the three areas is used for the measurement error.

Fig. 6. The Arrhenius plots of the natural logarithm of the experimentally determined rate constants for the growth of the oxide scale thickness (OST), $\gamma'$-dissolution layer (GPD), and Al$_2$O$_3$ finger depths (AFD) shows a linear relationship with (temperature)$^{-1}$ that gives a slope $-Q/R$, where $Q$ is the activation energy for growth and $R$ is the gas constant.
straight lines with a slope of $-Q/R$. Fig. 6. The activation energies for growth were determined to be $Q_{(OST)} = 262 \pm 33$ kJ/mol, $Q_{(AFD)} = 218 \pm 18$ kJ/mol, and $Q_{(GPD)} = 285 \pm 35$ kJ/mol. There is good agreement for $Q_{(OST)}$ with the literature value for Cr$_2$O$_3$ growth of 256 kJ/mol [14], even though TiO$_2$ grains have been included in the thickness determination.

Effect of Prior Exposure on ME3 Notched Fatigue Life at 704 °C
Static oxidation ME3 tests were used as a basis for the pre-exposure conditions used for the NLCF testing. Short (100 h), mid (440 h) and long (2,020 h) exposure times at the low (704 °C) and high (815 °C) isotherms were selected, as well as 760 °C for 440 h. To isolate the contributions of environmental attack and heat treatment on the notched fatigue behavior, specimens pre-exposed in vacuum at 815 °C for 440 h and blanks machined after pre-exposure in air at 815 °C for 440 h and 815 °C for 2,020 h were tested. Lastly, ~20 µm and ~50 µm in depth was polished away from the notch radii of specimens pre-exposed at 815 °C for 2,020 h to fully remove material to depths past the Al$_2$O$_3$ fingers and past the M$_2$3C$_6$ carbide dissolution layer.

Notch surfaces for at least one specimen for each condition were imaged with SEM prior to testing Fig. 7. As expected, no environmental attack was evident on notch surfaces of the unexposed, Fig. 7a, pre-exposed in vacuum, and machined pre-exposed blanks. This was also the case for pre-exposed specimens with material removed from the notch, Fig. 7g-h. A check of the longitudinal gage cross sections of these conditions confirmed that the repolish procedure was indeed successful in removing the targeted damage layers. With increasing temperature and time, the external oxide grains appear to grow, supporting the cubic rate law observed, and coverage becomes more uniform, Fig. 7c-f. For the 815 °C h for 2,020 h exposure, Fig. 7f, the number of faceted TiO$_2$ grains is significant.

A comparison of the effects of the various pre-exposure conditions on the mean notched fatigue life ($N_f$) of ME3 specimens tested at 704 °C is displayed in Fig. 8. Each test is represented by a diamond and $N_f$ by the bar chart. The majority of the conditions tested, including the unexposed, exhibited a scatter of about 2x–4x between the lowest and highest $N_f$. However, specimens exposed at 704 °C for 440 h, 704 °C for 2,020 h and at 760 °C for 440 h showed larger scatter.

Environmental exposures led to debits in fatigue life, which became more pronounced with increasing time and temperature, Fig. 8a. The unexposed samples had a mean fatigue life $N_f$ of 44,949 cycles compared to only 489 cycles for samples exposed to 815 °C for 2020 h, a nearly 99% debit. The 704 °C for 100 h exposure resulted in minor damage to the specimen surface, Fig. 8a, not sufficient to affect the fatigue life as there was no debit in $N_f$. Three of the four specimens exposed at 704 °C for 440 h showed no significant debit, while the fourth specimen exhibited 90% lower life. As shown for coupon oxidation, Fig. 5, 704 °C for 2,020 h, 760 °C for 440 h, and 815 °C for 100 h, produce roughly equivalent external scale thicknesses, Al$_2$O$_3$ finger depths and γ'-dissolution layer thicknesses. These samples had statistically equivalent mean fatigue lives, showing the depth of the surface damage is the major driver, not exposure temperature.
Specimens pre-exposed in vacuum at 815 °C for 440 h and machined blanks that were pre-exposed at 815 °C showed no debit in life, Fig. 8b. This confirms that the debits associated with prior exposures are due to environmental attack and any overaging of the microstructure by the pre-exposures does not significantly affect $N_f$. Removing the external oxide and oxide subscale from the notch (i.e., ~20 µm) only showed marginal improvement in $N_f$, 1,305 vs. 489 cycles, for the 815 °C for 2,020 h pre-exposure. However, material removal past the carbide dissolution layer yielded almost a full recovery in life, with $N_f$ of 32,980 cycles (i.e., ~50 µm).

The damage to the ME3 surfaces by the growth of an oxide layer likely influences the initial crack size, shape and distribution that govern the NLCF lives. The cumulative $N_f$ data for the various pre-exposures in air were normalized to the unexposed $N_f$ and plotted against the predicted external oxide scale thicknesses, Fig. 9. A log-log plot revealed a linear relationship that fit well to a simply power law of $n = -3$, which gave a $k_2 = 0.316 ± 0.024$ (µm %)$^3$. Coupling the cubic rate dependence of the oxide scale thickness, it follows that the percentage of fatigue life is proportional to (time)$^3$. This derived relationship predicts debits in fatigue life for oxide scale thicknesses that are greater than 0.68 ± 0.06 µm, which is roughly near the oxide scale thickness for the 704 °C for 440 h exposure.

**Figure 9.** Normalized mean fatigue lives for pre-exposed specimens versus external oxide scale thickness, predicted by measured rate dependencies for ME3 static oxidation. The solid line denotes the power law fit, $y = (k_2 \cdot t)^n$.

**Effect of Prior Exposure on ME3 NLCF Failure Modes**

To discuss the failure modes with some clarity, it is helpful to group the air exposure conditions by: modest (704 °C for 100 h, 704 °C for 440 h), moderate (704 °C for 2,020 h, 760 °C for 440 h, 815 °C for 100 h), and aggressive (815 °C for 440 h, 815 °C for 2,020 h) exposures. No debit in NLCF life was observed for the majority of the modest exposures, while there were measurable debits for both the moderate and aggressive exposures.

The number of major initiations sites observed for these NLCF tests is related to the degree of environmental damage. All failures initiated from the notch surfaces. Unexposed ME3 specimens failed from a primary crack initiating and propagating transgranularly normal to the loading axis. For the representative fracture surfaces in Fig. 10, the major initiations for the modest and moderate exposures are marked with white arrows, while they are unmarked for the aggressive exposure, since there are many initiations (>15). For the modest exposures, Fig. 10a, specimens failed with either one or two primary initiations. The number of initiations sites clearly increases with exposure, Fig. 10a-d, which likely results from the increased thickness of the brittle oxide at the surface. For the moderate and aggressive exposures, Fig. 11 shows that notch surface oxide has multiple cracks normal to loading, which are located near the major crack initiation sites from the failed specimens. With thicker oxides, the loading conditions will produce a higher number of cracks within the oxide. For the various moderate exposures, the cracks varied in length from 10 µm to 50 µm, while for the aggressive exposures the cracking is more extensive, with some cracks reaching 300 µm in length due to crack propagation and coalescence during testing. More developed oxide cracks are ripe for major initiations, which is consistent with the observed increase in initiations with exposure.

**Figure 10.** Fracture surfaces for ME3 NLCF tests at 704 °C of specimens pre-exposed in air at: a) 704 °C for 100 h, b) 704 °C for 2,020 h, c) 815 °C for 100 h and d) 815 °C for 2,020 h. Initiation sites are marked with white arrows.

**Figure 11.** From failed ME3 NLCF specimens exposed at a) 815 °C for 100 h and b) 815 °C for 2,020 h, these notch surfaces underneath a major crack initiation area show cracks within the oxide normal to the loading direction (approximately vertical).
The surface failure initiation for the moderate exposures is ambiguous. Flat areas were often observed along the initiation ledge, which are consistent with the oxide cracks, Fig. 11a. It is plausible that these cracks coalesce and initiate failure. The surface failure initiation for an aggressively exposed specimen clearly follows the shape of the typical well-developed oxide crack, Fig. 12, indicating these specimens failed by cracks that initiated in the oxide scale. Furthermore, the short lives (<1000 cycles) as well as fractographic evaluation, Fig. 10d, suggests that the onset of the multi-initiations likely occurs within the first few cycles, and then fatigue life is primary consumed by propagation, and initiated coalescence of oxide cracks.

The fracture surface views of the typical surface failure initiation sites for the moderate and aggressive exposures are also shown in Fig. 13. Complimentary lower magnification images of the near surface crack propagation are shown in Fig. 14, including the failure modes often observed for the modest exposures: transgranular initiation and propagation normal to loading. For the failure initiation site for the moderate exposure, Fig. 13a, the
darker contrast volume at the surface, observed in back-scattered SEM image of Fig. 13a, is the external oxide and recrystallized $\gamma'$-dissolved layer (GPD). Together these layers are approximately 4 $\mu$m thick. Environmental exposures cause the near surface grain boundaries to be degraded leading to early intergranular propagation that extends about one or two grains into the depth before transitioning to transgranular propagation, Fig. 13a and Fig. 14b. For the failure initiation site for an aggressive exposure displayed in Fig. 13b, the M$_2$X$_6$ carbides image as bright spots. It is clear that M$_2$X$_6$ carbides are dissolved through the first ~50 $\mu$m into depth. Despite this, intergranular propagation normal to the load for these specimens extended several hundred microns, reaching a full 1 mm for some cracks, Fig. 14c and Fig. 15. This indicates the ME3 grain boundaries beyond the carbide dissolution layer were also weakened by the exposures.

The most aggressive exposure condition of 815 °C for 2,020 h was used for layer removal experiments. As noted, the specimens that were aggressively exposed failed from cracks initiating at the environmental-affected surface layer with multiple initiations that propagated intergranularly normal to the loading axis. These exposed specimens had a mean NLCF life of 489 cycles. By removing the external and subscale oxides from the notch, yet leaving the carbide dissolution layer, the fatigue life only showed marginal improvement of 1,305 cycles. Fractographic evaluation revealed that the specimen failed by multiple intergranular initiations, where the cracks continued to propagate intergranularly normal to loading, Fig. 16. Intergranular propagation extended over 1 mm in distance before transitioning to transgranular overload, Fig. 16c, consistent with the specimens where no material was removed, Fig. 15. Surface initiations

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**Fig. 15.** White marker indicates the 810 $\mu$m transition from intergranular propagation to transgranular overload for a notched fatigue specimen that received an 815 °C for 440 h exposure prior to testing.

**Fig. 16.** Pre-exposed at 815 °C for 2,020 h, a specimen with the external oxide and oxide subscale removed (~20 $\mu$m from notch radius) shows: a) an intergranular initiation, b) multiple initiations and c) an intergranular propagation that extended 1490 $\mu$m.

**Fig. 17.** Pre-exposed at 815 °C for 2,020 h, a specimen with material past the carbide dissolution layer removed (~50 $\mu$m from notch radius) shows: a) a transgranular initiation, b) a single initiation and c) an intergranular propagation that extended 1570 $\mu$m.
clearly fail by the weakened ME3 grain boundaries, where the M₂₃C₆ carbides had dissolved from the environmental exposure.

By removing material to below the M₂₃C₆ carbide dissolution layer, the weakest grain boundaries at the surface are removed. This removal condition gave a significant recovery in fatigue life of 35,998 cycles, which is nearly comparable to the fatigue life of 44,949 cycles for the unexposed specimens. Furthermore, the initiation observed is a single, transgranular initiation, Fig 17, which is typical of unexposed failures. However, not surprisingly the intergranular propagation extended over 1 mm in distance before transitioning to transgranular overload, Fig. 17c, which is consistent with all specimens exposed at 815 °C for 2,020 h.

**Summary and Conclusions**

Oxidation at potential maximum service temperatures over extended periods, and its effects on notched fatigue life for an advanced power metallurgy disk superalloy ME3 were investigated.

Supersolvus ME3 oxidizes like other high Cr containing disk superalloys. The near surface damage that results is complex. A continuous Cr₂O₃ scale forms, with faceted, superficial TiO₂ grains at the exposed surface. Beneath the external scale, an internal subscale of branched Al₂O₃ extends into a layer where γ'-precipitates have been dissolved by Al depletion. This γ'-dissolution layer is recrystallized with finer grains and contains micron-sized voids at the grain boundaries. A semi-continuous (Ta₃Cr)O₅ wets the interface between the Cr₂O₃-TiO₂ scale and Al₂O₃ subscale. Throughout these layers and beyond, the M₂₃C₆ carbides have been dissolved from the original grain boundaries via grain boundary diffusion of Cr that helps support the Cr₂O₃ scale growth. The Al₂O₃ subscale growth follows the classic parabolic growth, however, the Cr₂O₃-TiO₂ oxide scale deviates. The cubic rate dependence is attributed to grain growth in the external scale. The underlying diffusion processes also lead to a cubic rate dependence for the γ'-dissolution layer.

Prior exposure in air at 704°C, 760 °C and 815 °C significantly affected mean notched low cycle fatigue (NLCF) life. Mean NLCF fatigue life at 704 °C was measured from the average of 2-4 tests for each condition, where the majority of the conditions exhibited a scatter of about 2-4 times between low and high values. Reductions in NLCF fatigue life showed a power law decay with oxide scale thickness, suggesting that initial crack size, shape and distribution is correlated to the scale thickness. Experiments on aged material and vacuum exposures showed that environmental damage, not microstructural overaging, was driving the reduction in life. In addition, experiments on specimens with equivalent scale thicknesses from pre-exposures at the three isotherms had statistically equivalent fatigue lives, which demonstrates that fatigue debit is caused by surface damage not exposure temperature. No significant improvement in NLCF fatigue life was observed with the removal of the oxide scale and subscale, however a near full recovery was observed for specimens where material was removed past the M₂₃C₆ carbide dissolution layer. This indicated that grain boundary strength from the M₂₃C₆ carbides is important to the crack initiation for these tests. Despite material removal from the surface of these aggressively exposed specimens, the primary cracks propagated intergranularly, in contrast to the transgranular propagation typical for the unexposed specimens. This suggests the ME3 grain boundaries beyond the carbide dissolution layer were also weakened by the exposures. Further work is planned to determine the oxygen content at the grain boundaries.

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