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An overview of the oxidation of Ni-based superalloys for turbine disc applications: surface condition, applied load and mechanical performance.

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Abstract

In this review the role of oxidation of chromia-forming Ni-based superalloys used for rotor disc applications in gas turbines is examined. A detailed review of the oxidation performance of these alloys in air is undertaken with emphasis on oxide composition and kinetics. The actual surface condition of the component entering service is important but often overlooked during oxidation studies. The effect of shot-peening on the oxidation of Ni-based superalloys is here evaluated. The oxide growth rates and composition of these alloys was assessed with respect to their response to an applied load or stress. One area where a significant amount of research has been conducted is on the effect of oxidation on the mechanical performance, with oxidation having been shown to reduce time to crack initiation, increase fatigue crack growth rates and therefore reduce overall component fatigue life. It is argued that this enhancement in crack growth rates arises from the cracking of an oxide intrusion ahead of the crack tip, exacerbated by Stress-Aided Grain Boundary Oxidation (SAGBO).

Introduction

Ni-based superalloys have been optimised, compositionally and microstructurally, to operate in the highly stressed conditions occurring in the hot sections of gas turbine engines. Under such conditions degradation by oxidation occurs and plays a key role in determining the lifetimes of the components. For this reason it is important to understand the oxidation processes occurring and the mechanisms involved. Most oxidation studies, especially those on Ni-based superalloys, are performed on specimens having a ground or polished surface rather than the actual condition the component would have entering service, in particular shot-peening, and it is important to quantify the effect this may have. Additionally, studies have primarily focussed on the study of oxidation without the application of an external load. This may be an important omission since in the real life operating conditions of these alloys, and the components they will be used for, significant external stresses can be present. This overview will examine the basic oxidation properties of chromia-forming Ni-based superalloys, the influence of surface conditions on these and oxidation/mechanical properties interactions.

Oxidation of chromia-forming Ni-based superalloys in air

Typical chemical compositions of various examples of this important group of γ' (nominally, $\text{Ni}_3(\text{Al,Ti})$) strengthened alloys are given in Table 1. Of relevance to oxidation properties, they are usually characterised by Cr contents in the range 13-20 wt.%, Al of 0.5 to 3.5 wt.% and Ti of 0.75 to 5 wt.%. Peak operating temperatures are unlikely to exceed 800°C.

Composition (% wt)	Ni	Co	Cr	Mo	Ti	Al	Ta	Hf	Zr	C	B	W	Nb	Fe
Udimet 720 Li	Bal	15.0	16	3	5	2.5	-	-	0.05	0.025	0.018	1.25	-	-
Astroloy	Bal	17	15	5.3	3.5	4	-	-	-	0.06	0.03	-	-	-
Waspaloy	Bal	13.6	19.3	4.2	3.6	1.3	-	-	0.05	0.01	0.04	-	-	-
ME3	Bal	18.2	13.1	3.8	3.5	3.5	2.7	-	0.05	0.03	0.03	1.9	1.4	-
IN718	Bal	-	19	3	0.9	0.5	-	-	-	0.04	-	-	5.1	18.5
Rene 95	Bal	8.0	14.0	3.5	2.5	3.5	-	-	0.05	0.03	0.01	3.5	3.5	-
RR1000	Bal	18.5	15	5	3.6	3	2	0.5	0.06	0.03	0.02	-	-	-
IN718 plus	Bal	9.1	18	2.7	0.75	1.45	-	-	-	0.02	0.005	1.0	5.4	9.5

Table 1: Compositions of some representative alloys [1-4].

The oxidation behaviour of several of these alloys has been studied extensively in air, e.g. IN718, RR1000, Udimet720, ME3, etc. [5-19] using both isothermal and cyclic oxidation tests. In general, the oxide scale formed is usually duplex consisting of a dense chromia (Cr_2O_3) layer and an outer layer of rutile (TiO_2) and a sub-surface internal oxidation zone (IOZ) of alumina (Al_2O_3), often consisting of acicular intergranular penetrations [5, 16, 18]. The IOZ is associated with a γ' denuded zone, caused by Al depletion, that extends further into the alloy. Other surface oxides are often seen, e.g. NiO and CoO, and are often associated with early-stage transient oxidation. The newer more compositionally complex alloys, such as RR1000 and ME3, demonstrate most of these features as shown in Figure 1 but even in earlier, simpler alloys, e.g. Waspaloy, Astroloy and Udimet 720, internal oxidation of Al remains a dominant feature [5]. Both RR1000 and ME3 have each been reported to produce a new previously unseen oxide at the external oxide/alloy interface, with (Ti, Ta) O_2 being reported in RR1000 [17] and (Ta, Cr) O_2 in ME3 [15]. Of those alloys listed in Table 1, these are the only ones to contain Ta.

In addition to this new oxide, the external oxide/alloy interface becomes more complex with exposure time, developing protuberances and metallic regions within the oxide (e.g. Figure 1). This may be a

result of undercutting by oxide formation [20] or through outward alloy creep to accommodate the increase in volume from internal oxidation [21-25].

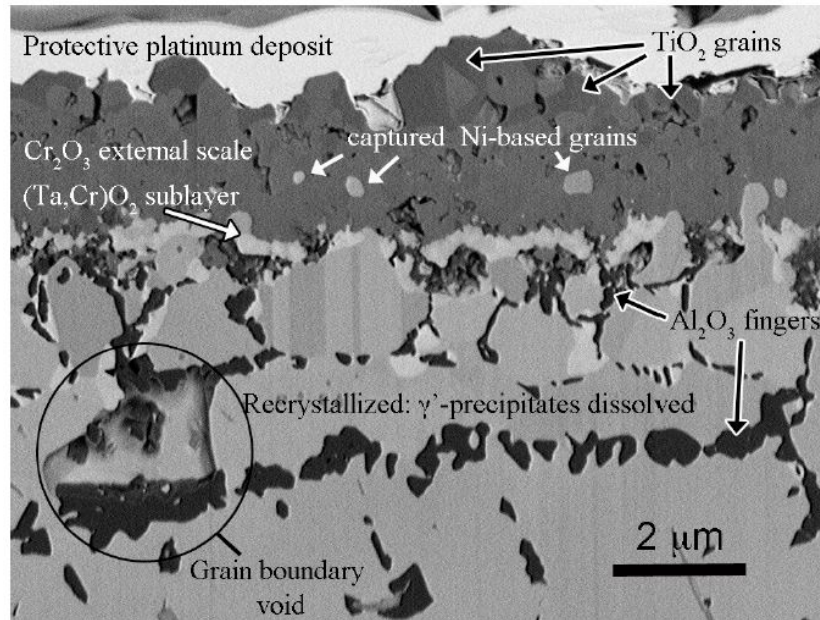


Figure 1: Oxidation of Ni-based superalloy, ME3, exposed to air at 815°C for 2020 hours, showing a surface layer of Cr₂O₃ with an internal oxidation zone of Al₂O₃ [15]. See text for fuller discussion.

Underneath this surface oxide, as mentioned previously, internal oxides and a γ' particle free zone develop. Within this region recrystallisation, or perhaps simply grain growth, has been reported in both ME3 (Figure 1) and RR1000, although this has only been found in the fine-grained not the coarse-grained variant of the latter [7, 15, 26]. To add further complication, at particularly high temperatures (e.g. $\geq 850^\circ\text{C}$), particles of TiN have been shown to form ahead of the IOZ [14].

Furthermore it has been reported that within and ahead of the internally oxidised region, due to the depletion of Cr to the external oxide scale, is a zone of Cr-rich phase dissolution [15, 18]. Beneath this layer, remote from the oxide scale the grain boundaries are decorated with $M_{23}C_6$ precipitates (nominally $(\text{Cr},\text{Mo})_{23}\text{C}_6$) [15, 18]. MC carbides reside on the majority of the grain boundaries in the as-received alloy and decompose during high temperature exposure to $M_{23}C_6$. It is possible that Cr and Mo rich sigma (σ) phase (nominally $(\text{Ni},\text{Co})_x(\text{Cr},\text{Mo})_y$) could also form notionally as [27]:



In principle, the dissolution of such precipitates could lead to void formation [28] and initial work suggested [7] that sub-surface voids were present in oxidised RR1000, albeit attributed to vacancy injection [e.g. 29]. Later studies [30] have confirmed that the apparent voids were indeed sub-surface oxide particles. Nevertheless, some voidage has been observed in these alloys, as shown in Figure 1, but the possibility cannot be neglected that the example shown results from preparation damage, i.e. the loss of a small grain.

Most oxidation studies of these alloys have assessed the reaction kinetics using mass change measurements. The oxidation kinetics tend to be well described as parabolic and, as Figure 2 demonstrates, broadly similar parabolic growth kinetics are found [5, 7, 11, 12, 14, 16, 17]. Here, k'_p is the parabolic rate constant evaluated from the mass gain, (Δm) after oxidation time 't' as:

$$k'_p = \frac{(\Delta m)^2}{t} \quad (3)$$

The best-fit line through all the data in Figure 2 gives an Arrhenius expression for k'_p of:

$$k'_p = 5.68 \times 10^6 \exp\left[-\frac{281329}{8.314T}\right], \text{ mg}^2 \cdot \text{cm}^{-4} \cdot \text{s}^{-1} \quad (4)$$

here T is temperature in K.

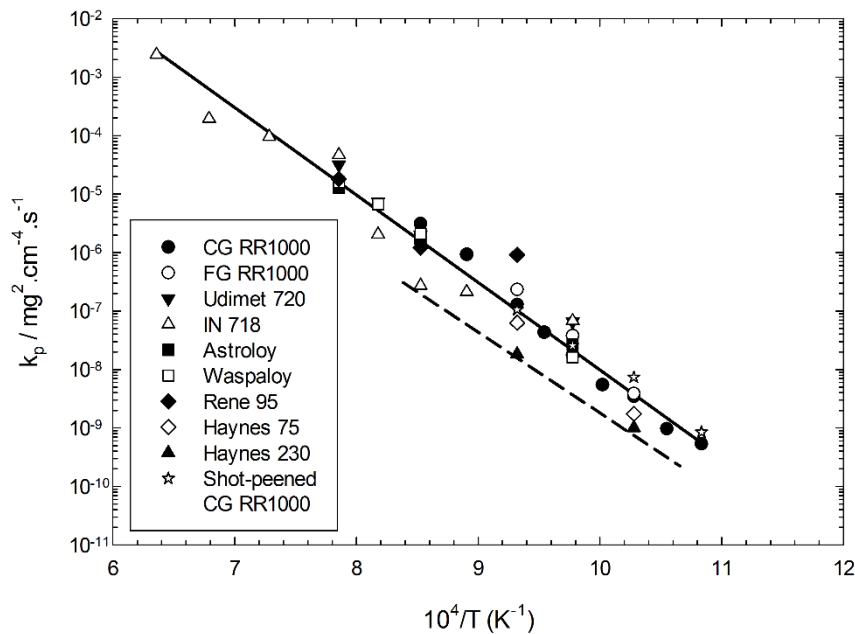


Figure 2: Arrhenius plot of k_p values from the literature of similar Ni-based superalloys used for rotor disc applications [5, 7, 11, 12, 31, 32] and typical expected behaviour for chromia formation on a high chromium containing austenitic stainless steel (dashed line) [29]. Image modified from Cruchley et al. [14] to include data taken from [17].

This value of activation energy seems appropriate for all the relevant alloys shown but it is clear that there exists some scatter of values of the parabolic rate constant shown in Figure 2. The obvious, and most robust of these, is that the temperature dependence of the oxidation rate of the alloys is broadly similar to that observed for pure chromium and chromia-forming simple austenitic steels [29, 33] (the dashed line in the figure). This line also forms a lower bound to the superalloy data. Within the spread of these data, there also appears to be a trend that alloys low in Ti and/or Al (e.g. IN718, Haynes 75 and Haynes 230, show improved behaviour than the other superalloys. This is not unexpected since reactive elements such as Ti and Al are also being oxidised [e.g. Figure 1] and will contribute to the mass increase. As will be shown below, however, for RR1000, the chromia thickening rate was also substantially increased and, it will be argued, that this is a direct influence of Ti doping of the chromia layer.

A more informative method of evaluating reaction kinetics is through extensive metallographic measurements of oxide thickness, morphology and type and this has been undertaken recently for RR1000 [14, 17]. In this work, particular attention was paid to the chromia layer and its thickening with exposure time. The kinetics of chromia growth on polished specimens at 750°C are shown in Figure 3 where a significant enhancement in the chromia growth rates over chromia formation on an austenitic steel and chromium was found. Sub-parabolic growth kinetics were found for the polished condition at 700 and 750°C and near parabolic growth at 800°C. The lower solid line in Figure 3 represents the best-fit through data obtained for chromia growth on a Ti-free austenitic steel (20Cr25Ni,Nb-stabilised) and for adherent layers on pure chromium [33] and given by:

$$k_p = 2.07 \times 10^{-6} \exp\left[-\frac{31020}{T}\right], \text{ m}^2\text{s}^{-1} \quad (5)$$

Here, T is temperature in Kelvin, the parabolic rate constant, k_p , is expressed in terms of thickness, ξ , of the chromia layer as:

$$k_p = \frac{\xi^2}{t} \quad (6)$$

where t is exposure time.

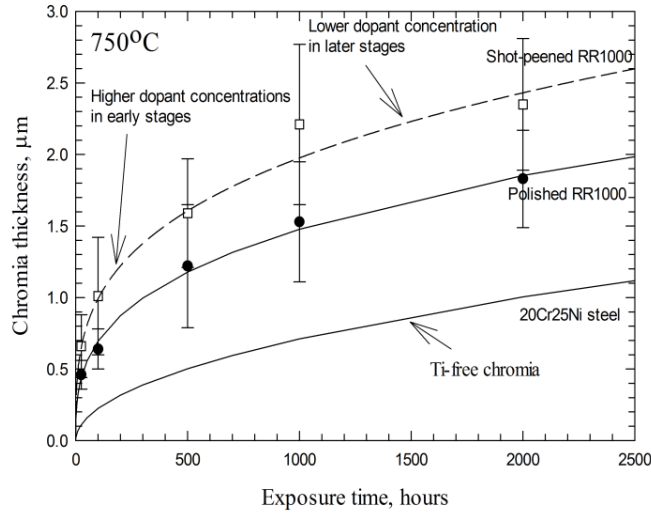


Figure 3: Plot of chromia thickness measured on RR1000, with and without shot-peening, at 750°C compared with Ti-free austenitic steel/pure chromium. Oxide measurements for both conditions were normally distributed and error bars are shown as ± 1 standard deviation. Hollow squares are measurements for shot-peened RR1000, filled circles are polished RR1000 [17].

SEM/EDX examination of the chromia layer in the above specimens has shown that substantial quantities of Ti are present in solution, not associated with TiO_2 . The ability of chromia to dissolve Ti, probably with a valency >3 , has been known for some time (e.g. [34, 35]). A possible defect reaction suggested by Atkinson and co-workers [35, 36] envisages a 4-valent Ti ion entering the chromia lattice according to:



here $3Ti_{Cr}^{\cdot}$ represents the excess positive charge associated with the titanium ion located on a chromium lattice site, $V_{Cr}^{\prime\prime\prime}$ is a triply charged Cr vacancy, O_o is the oxygen ion on the oxygen sublattice. The excess vacancies would increase the oxidation rates through the increased diffusion of chromium ions across the oxide layer. Sub-parabolic kinetics are thought to have developed as a result of the average quantity of Ti within the chromia layer decreasing as the layer thickens because of Ti depletion in the underlying alloy. As a consequence, the enhancement in growth rate reduces with exposure time and the kinetics tend to sub-parabolic rather than parabolic [14]. This trend of a decreasing enhancement in the growth rate of the doped oxide with exposure time is shown graphically in Figure 4(a) for polished RR1000 specimens and Figure 4(b) for shot-peened RR1000 specimens [17]. Here the enhancement factor is expressed by the ratio, r , of chromia growth rates in the superalloy (suffix S) to that of Ti-free chromia (suffix Cr) at a given oxide thickness in both cases:

$$r = \left[\frac{(d\xi/dt)_S}{(d\xi/dt)_{Cr}} \right]_{\xi} \quad (8)$$

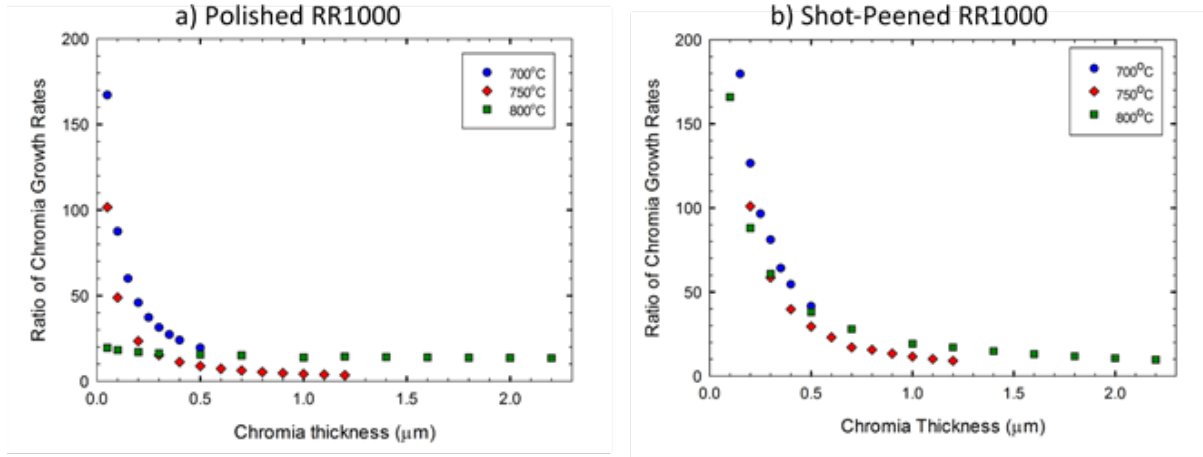


Figure 4: The variation of the enhancement ratio of the chromia growth rate due to Ti doping with oxide thickness for 700°C, 750°C and 800°C, in a) polished RR1000 and b) shot-peened RR1000 [17].

The enhancement ratio is high for thin oxides at 700 and 750°C on the polished specimens, indicating high dopant levels, but rapidly decreases with increasing oxide thickness. At 800°C, the enhancement is lower initially but remains reasonably constant with increasing oxide thickness. The reason for this is unclear but may be associated with the presence of a near-continuous layer of the (Ti,Ta)O₂ phase underlying the chromia scale. A consequence is that near-parabolic kinetics obtained at this temperature but not at the two lower temperatures shown.

A characteristic feature of the chromia-forming superalloys examined here is the presence of regions of sub-surface, oxidation-induced damage. The most striking of these is the formation of oxides of Al and Ti which can exist as acicular intrusions along alloy grain boundaries (Figures 1 and 5) as well as discrete particles within the alloy grains [18]. As will be discussed, the grain boundary oxides can influence the nucleation of surface cracks during mechanical testing. Internal oxidation of these alloys has been reported extensively [5, 9, 11, 13] but detailed measurements of kinetics are sparse. An extensive metallographic study has recently been performed by Cruchley et al. on RR1000 over the temperature range of 700-800°C for exposure periods up to 2000 hours in air [18]. This work showed that the intergranular and intragranular penetration rates were parabolic. The selective oxidation of Al and Ti results in the dissociation of the γ' (Ni₃(Al,Ti)) precipitates in the near-surface regions and the formation of a precipitate-free zone (PFZ). The depth of this zone also increases parabolically with time [18].

Effect of shot peening

Surface finish and residual stress underneath the surface are controlled and modified to improve the mechanical properties of in-service components. Various studies have established that the oxidation behaviour of stainless steels, Ni alloys and Fe-Cr alloys are dependent on the surface finish prior to oxidation [37-47]. Relatively little work has been conducted, however, on the effect of surface condition on the oxidation of Ni-based superalloys where most studies have examined the ideal highly polished or as-received machined condition [5-7, 15, 48].

Shot-peening has been widely employed to improve the fatigue resistance of components by inducing significant surface compressive residual stresses [49, 50]. For the current chromia-forming superalloys, the effect of shot-peening on oxidation has only been investigated in detail in two studies on coarse-grained (30-50 μm) Ni-based superalloy, RR1000 [17, 19]. Testing in both cases was performed in air between 700-800°C for up to 2000 hours. In this variant of RR1000 it has been reported that recrystallization of the near surface grains only occurs in the oxidised shot-peened condition [17], which is in contrast to the findings reported for the fine-grained variant of RR1000 (4-6 μm) and ME3 (27-30 μm) [7, 15, 26]. Such recrystallization can be expected to produce a surface region of different mechanical properties from that of the bulk alloy.

While no changes in oxide composition have been reported for shot-peened specimens, changes in morphology of the internal oxide and the γ' PFZ have been observed, together with the development of a more planar front of internal oxidation (Figure 5) [17, 19]. This morphology appears to result from the finer grain size associated with the recrystallized zone (Figure 5b). At 800°C, both the shot-peened and polished specimens developed (Ti, Ta) O_2 at the chromia/alloy interface although this occurred as isolated regions in the shot-peened condition compared with a near continuous layer in the polished condition [17].

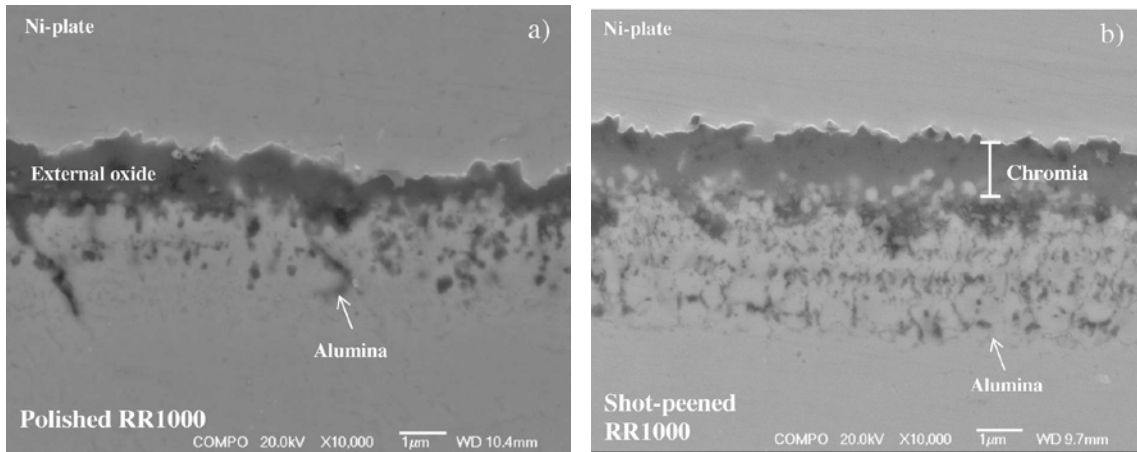


Figure 4. Cross-sectional SEM images of a) polished RR1000 and b) shot-peened RR1000 oxidised at 700°C [17].

The growth kinetics of the surface chromia layer as determined from the measurements [17]. As with the polished condition a significant rate enhancement over those expected for Ti-free chromia was reported, particularly in the early stages of oxidation. This is illustrated in Figure 4, using equations (5), (6) and (8). The behaviour is similar to that found for the polished condition (Figure 4a) except that the trend exists at 800°C also and this may be related to the non-continuous nature of the (Ti,Ta)O₂ phase at the chromia/alloy interface in the shot-peened specimens. As with the polished specimens, Ti doping of the chromia layer is thought to be responsible for the large rate enhancement. The decline of the enhancement ratio with increasing oxide thickness indicates that the chromia growth kinetics are sub-parabolic in the shot-peened specimens of RR1000 over the temperature range of 700-800°C.

An interesting study by Foss et al. [26] has investigated oxygen transport in fine-grained RR1000 at 800°C for 120 hours using a two stage isotopic exposure (¹⁶O₂ for 72 hours followed by ¹⁸O₂ for 48 hours) at 200 mbar. Both polished and shot-peened surface finishes were examined but the general oxide morphology was similar in both and also to the results reported by Cruchley et al. [14, 17] on the coarse-grained alloy variant. A significant observation was that the surface chromia layer grew both by cation diffusion (new oxide formed on the outer surface of the layer) and by anion diffusion (new oxide formed at the oxide/metal interface). The growth of chromia by such counter-current diffusion is well established for layer growth on simpler alloys [51] so its demonstration on a complex superalloy with Ti-doped chromia is a helpful contribution. It shows that the doping is not

accommodated by significant defect changes on the oxygen sub-lattice, an expected reduction in oxygen vacancy concentration, but by the cation sub-lattice as indicated by the defect reaction (7).

Application of an applied load on oxide composition and growth

Studies of high temperature oxidation of metals have focussed primarily on oxidation composition and growth kinetics in the absence of an external load. This is an important omission since in service these materials, and the components they will be used for, often experience significant external stresses. This is especially the case with Ni-based superalloys which are designed to operate in the highly stressed regions of gas turbine engines. While the literature is limited, several studies have investigated the application of an external load, either compressive or tensile, on the oxidation kinetics of a number of alloy systems [26, 32, 52-57].

Studies in this area have often focussed on Ni and simple Ni-based alloys as this is the base metal/system used in most superalloys. For example, application of tensile stresses of 10 and 20 MPa have been shown to increase the weight gain due to oxidation of Ni at 700°C whereas a stress of 6 MPa did not [52, 53]. This apparent critical stress could be associated with through-thickness cracking of the oxide scale which would be expected to increase oxidation rates. Its value can be estimated well enough from [58]:

$$\sigma_c = \left(\frac{K_{Ic}}{f(\pi a)^{0.5}} \right) \quad (9)$$

Here the defect half-length, a , is measured perpendicular to the in-plane critical stress, σ_c , in the oxide layer. f is a geometric factor, ~ 1 , and K_{Ic} is the critical stress intensity factor. Measurements on NiO scale grown on Ni give $K_{Ic}=0.4 \text{ MPa}\cdot\text{m}^{-1/2}$ at room temperature and $1.6 \text{ MPa}\cdot\text{m}^{-1/2}$ at 900°C [59]. The oxide layer thickness in the experiments of Zhou et al. [52] was $\sim 3 \mu\text{m}$ so that the maximum defect length present was unlikely to be larger than $1.5 \mu\text{m}$, i.e. $a=0.75 \mu\text{m}$. Using this value in equation (9) together with a mid-range value of $K_{Ic}=1 \text{ MPa}\cdot\text{m}^{-1/2}$ and $f=1$ gives the critical stress as $\sim 650 \text{ MPa}$, i.e. approximately 2 orders of magnitude larger than that reported by Zhou et al. [52]. This very large discrepancy was not addressed in the original papers but can be understood once it is appreciated that the stresses quoted by Zhou et al. seem to be those applied to the gross sectional area and not those necessarily present in the oxide layers. The specimens tested were composites consisting of an outer NiO layer and a Ni core. Prior to cracking of the oxide, i.e. up to a gross applied stress of $\sim 6 \text{ MPa}$, both of these components will deform to the same uniaxial strain in order to retain continuity. In order to achieve this, the applied load will redistribute across the cross section according to [60, 61]:

$$2\sigma_{ox} \xi w + 2\sigma_s h w = 2\sigma_g (h + \xi)w \quad (10)$$

Here, the specimen of rectangular cross-section is oxidised on all surfaces to produce an oxide thickness, ξ , on the major flat surfaces. The core width is $2h$ and the specimen has thickness w . σ_g is the gross applied stress, σ_{ox} is the average in-plane stress in the oxide layer and σ_s is the corresponding average value for the core alloy. From equation (10), the average stress in the oxide layer is:

$$\sigma_{ox} = \left[\frac{\sigma_g(h + \xi) - \sigma_s h}{\xi} \right] \quad (11)$$

Higher stresses will be generated in the stronger component, in this case, the oxide layer which is expected to deform elastically. The Ni core, by contrast, will deform by creep at much lower stresses. It is beyond the scope of the present paper to undertake a detailed evaluation but a rough estimate of the oxide stress can be made. The tensile fracture strain of the oxide will be around 0.5% [59] which requires an in-plane tensile stress of ~ 875 MPa for a Young's modulus of the oxide at 700°C of 175 GPa [62]. This value of stress is again very much larger than that applied to the entire cross-section but is consistent with that deduced from likely defect lengths through equation (9). Similar considerations apply to the tests undertaken in compression by Zhou et al. [52] where enhanced oxidation and the formation of fine-grained oxide was found for gross applied stresses of 20 MPa [52]. On the basis of the above arguments, actual in-plane compressive stresses in the oxide are likely to be of order 1 GPa and these may initiate oxide spallation. This was not reported by Zhou et al. but was found by Moulin et al. [54] under creep and creep/fatigue conditions.

It has been important to re-examine the results of Zhou et al. since their study provides a systematic investigation into the influence of stress on oxidation in the Ni/NiO system. Of concern was the possibility that significant deleterious effects could be produced at very low applied stresses which would be encountered readily in service. The above discussion, however, offers reassurance that, in fact, much larger stresses, of order 1 GPa, need to be applied to the oxide to produce the observed cracking and morphological changes. This issue of quantifying the magnitude of stress in the oxide layer is generic whenever testing is done using conventionally-shaped test specimens as used for uniaxial loading. As far as the present authors are aware, no allowance has been made in the available literature for the redistribution of load that occurs on testing the oxide/metal composite specimen. Nevertheless, when the core material is a strong superalloy of high creep strength and a high yield stress, e.g. ≥ 1 GPa, the load-bearing capacity of both the core and oxide layer will be similar. The load distribution across the section will then not be large but should still be evaluated in order to obtain quantitative insights.

The published work on the superalloys clearly shows qualitatively an influence of applied stress on oxide morphology and section loss. An example, taken from the work of O'Hanlon et al. on fine-grained RR1000 [56], is shown in Figure 6. In this study, the effect of a static load was investigated at both 650°C and 700°C for 200 hours under compressive and tensile gross stresses of 300 and 600 MPa. Figure 6 shows examples of the oxide morphology developed at 600 MPa together with that under unstressed conditions (Figure 6b). It is clear that under both tensile (Figure 6a) and compressive (Figure 6c) stress, the protective surface oxide has been compromised, probably by cracking or local spallation, and regions of enhanced damage develop. A related study [55] on fine-grained RR1000 tested under tensile fatigue at 700, 750 and 800°C did not demonstrate enhanced thickening of the surface oxide although, at 800°C only, there appeared to be an enhancement of sub-surface attack at the peak tensile stress of 670MPa, at least at mean behaviour.

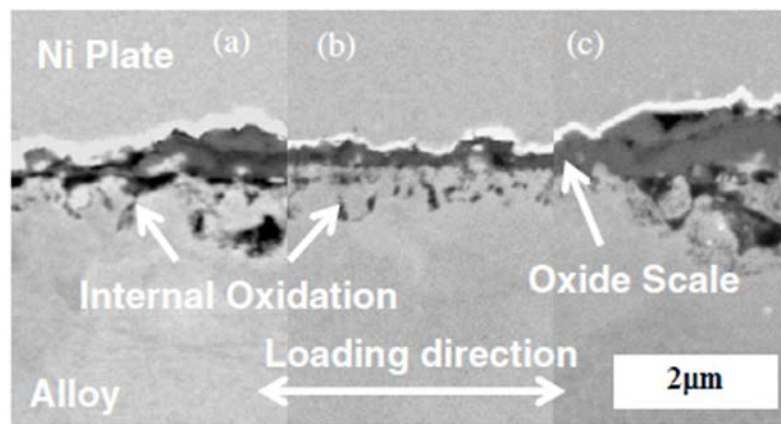


Figure 6: Backscatter electron images of oxidation damage on fine-grained RR1000 after 200 hours at 700 °C under hold stresses of (a) +600MPa, (b) 0MPa and (c) -600MPa [56].

Various other studies on chromia-forming superalloys such as Udimet 720, ME3 [63], IN100 [57] have demonstrated that applied stress can affect not just oxide morphology and thickness but also the depth of internal oxidation and the γ' depleted zone. These tend to show a deleterious effect at high stresses but not all results are consistent. In particular, a tensile stress (nominally, 133 MPa maximum) may promote the earlier beneficial formation of a chromia layer in Haynes 75 and 230 [32]. Interpretation of all these data is not straightforward because of the uncertainty in establishing actual oxide stress levels. The difficulty is compounded not just by the load redistribution discussed above but also by stresses developed by growth strains within the oxide layer. These can be negligible for the Ni/NiO system [64] but may be hundreds of MPa compressive for chromia scales [65].

The Influence of Oxidation on Crack Nucleation and Growth

Crack Nucleation

The alloys considered here usually experience systematic variations in stress during service and considerable research effort has been expended in attempting to understand their high-temperature fatigue behaviour under oxidising conditions. A recent overview of the topic has been provided by Pineau et al. [66]. It is not the intention to reprise this here but rather to focus on recent observations on oxidation effects on both crack nucleation and, in the next section, on crack growth.

An example of the oxide morphology that develops in these chromia-forming, Ni-based superalloys as a result of air exposure has been shown in Figure 1. As has been discussed, the oxides that form have low tensile fracture strains ($< 1.0\%$) and low fracture toughness ($0.4\text{-}2 \text{ MPa}\cdot\text{m}^{1/2}$) [67]. It would then not be unexpected for oxide cracking, either surface or intergranular, to occur at local net tensile stresses in the range $0.5\text{-}1.0 \text{ GPa}$ [68, 69] and an example of this is shown in Figure 7 [68]. This specimen of RR1000 had been exposed without stress in air for 2000h at 700°C and then fatigued (at $R=0.1$) under 4-point bend at room temperature at a maximum in-plane surface stress of 800 MPa . In the light of the earlier discussion, this calculated value of stress is probably reasonable since both the oxide and alloy will have deformed elastically and have similar load-bearing capacities. There remains uncertainty, unfortunately, in the oxide growth and thermal stresses that exist.

The micrograph of Figure 7 clearly shows the presence of cracking in both the surface chromia layer and the alumina sub-surface intergranular penetration. This crack has penetrated intergranularly into the alloy ahead of the oxidation front but within a presumably weak grain boundary zone depleted of γ' precipitates and Cr-rich carbides. This zone develops as the result of the selective oxidation of Cr, Ti and Al and its depth should be considered in assessments of oxidation-induced damage.

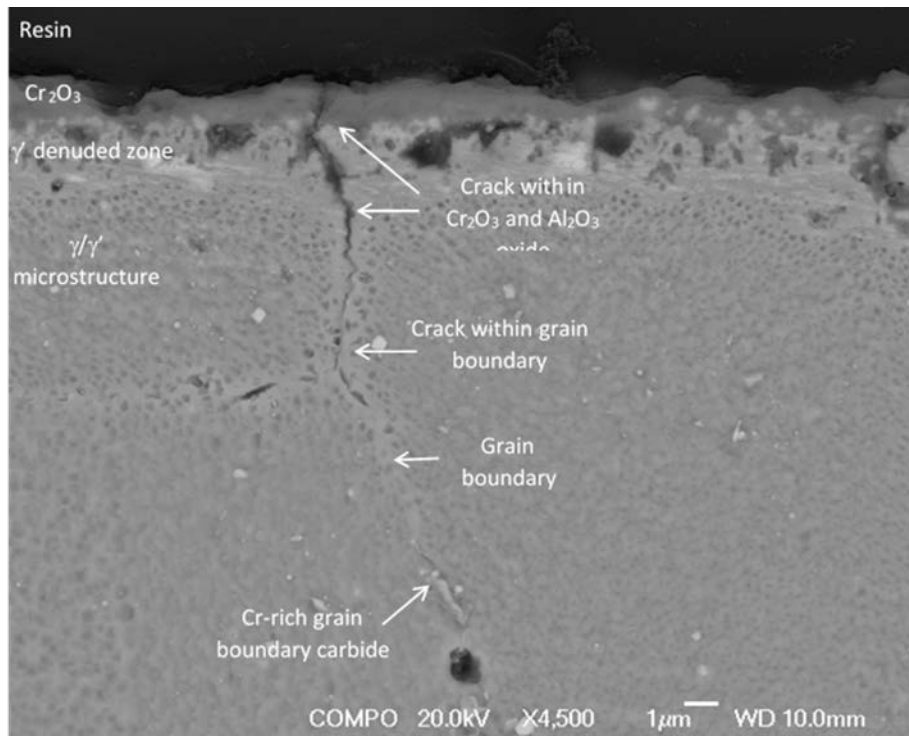


Figure 7: BSE image of an etched pre-oxidised (2000h, 700°C) specimen that failed after 1.9×10^6 cycles during room temperature 4-point bend at a maximum applied stress of 800 MPa, showing cracking of the external and intergranular internal oxide along with the presence of both a γ' denuded zone and carbide dissolution zone [68].

The importance of accounting for the presence of this zone is further demonstrated by the experiments of Sudbrack et al. [15] on ME3. Their results, summarised in Figure 8, show that prior high temperature exposures in an oxidising environment for exposure periods up to 2020 hours have a detrimental effect on the high temperature (704 and 815°C) notched fatigue life [15]. The thicker the external scale and the deeper the internal damage the more pronounced the reduction in life. The reduction in life in this case was driven by $M_{23}C_6$ carbide dissolution. Removal of the internally oxidised region did not lead to a complete recovery in high temperature fatigue life ($\sim 20\mu\text{m}$) but the removal of the carbide dissolution zone ($\sim 50\mu\text{m}$) did (Figure 8).

Another study using prior exposures on the Ni-based superalloys, ME3 (704°C for 439 hours) and Udimet 720 (650°C and 704°C for 100 or 1029 hours) found that the mean lives of pre-oxidised specimens had up to a 70% reduction in low-cycle fatigue (LCF) life. A change of crack initiation was also seen from sub-surface (as-received) to surface (pre-oxidised). Performing the prior exposures in vacuum led to no reduction in fatigue life, illustrating that oxidation damage is driving the reduction in life [63]. Pre-oxidation under an applied stress (97 MPa) has been shown to cause a further reduction in LCF life of the Ni-based superalloy, Rene 80, over both as-treated and unstressed prior oxidised (100 hours at 982°C) specimens. This is in addition to the reduction in life the

unstressed prior oxidised specimens recorded over the as-treated specimens [70]. Fatigue testing of the Ni-based superalloy, IN100, at 1000°C in an oxidising environment has been shown to drastically reduce the time to crack initiation compared with tests performed in vacuum. The crack initiation life was similar to the total number of cycles it took to fracture the oxide [57]. Similar observations of crack nucleation at oxidised grain boundaries have been reported recently on N18 [71] where it was also noted that the cycles to crack initiation formed the larger part of the total cycles to failure.

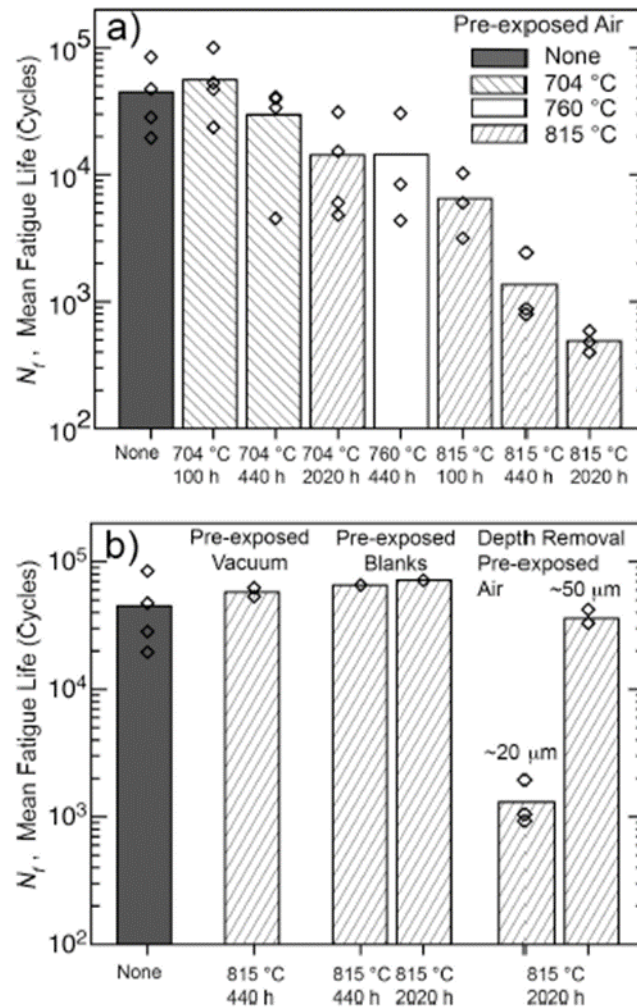


Figure 8: Comparison of the mean notched fatigue lives at 704 °C of the unexposed ME3 specimens and ME3 specimens with a) prior exposures in air and b) alternative conditions as marked. Diamonds are each test result, clearly showing that the removal of all of the environmental damage (50 mm removal) leads to a full restoration in fatigue life [15].

Further studies on RR1000 using prior exposure at 700°C have been shown to affect the high cycle fatigue (HCF) life significantly [68, 69]. Predominantly this was by reducing fatigue lives although exposed specimens were also observed to outperform the as-machined testpieces at stress levels around 825-900 MPa. This unexpected, beneficial effect of oxidation was attributed to local stress relaxation in the surface γ' -depleted zone associated with the selective oxidation of Al and Ti. The

yield stress of this zone was surmised to be around 825 MPa, i.e. substantially less than that (> 1 GPa) of the bulk alloy. At lower stresses, no relaxation took place and the cracking of surface oxides proved deleterious (Figure 7). At stresses > 900 MPa, crack propagation through the depleted zone occurred readily and again, prior oxidation proved deleterious. This is an intriguing result, in principle, but, as discussed above, further work is required to quantify the stress state in both the external oxide and the sub-surface internally oxidised region.

Crack Growth

Extensive evidence exists in the literature showing the deleterious effect of oxidising environments on high temperature dwell fatigue performance. Some of this is due to crack nucleation, as discussed above, but there is also a substantial increase in crack growth rates in the presence of oxygen [72-76] and a change in crack path from transgranular to intergranular, typically seen in low oxygen environments (vacuum/argon). An example of these features is shown in Figure 9 [76] for the LHSR alloy (12.5 Cr, 20.7 Co, 2.7 Mo, 4.3 W, 3.5 Ti, 3.5 Al, 1.6 Ta, 1.5 Nb, bal. Ni, wt.%) tested at 650 and 725°C in either air or vacuum with single notched specimens under 3-point bending. A general feature of such plots is that the crack growth rate decreases sharply as some notional threshold ΔK_{th} or K_{th} is approached. The threshold value is lower in air than in vacuum.

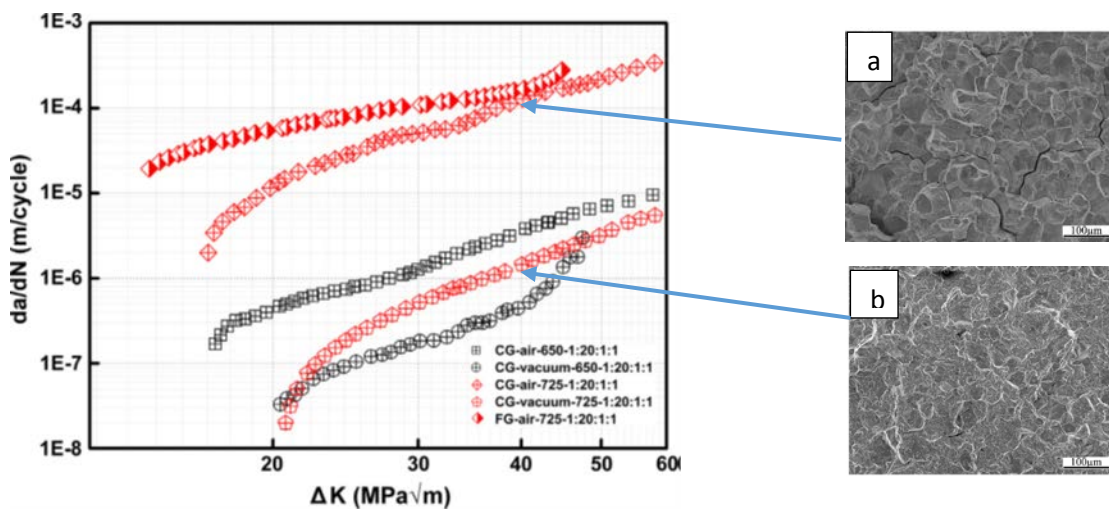


Figure 9: Main graph shows crack growth rates versus ΔK in the LHSR alloy tested under 3-point bend at 650°C (black symbols) and 725°C (red symbols) under air and vacuum. Increased rates were observed in air at both temperatures but were greater at the higher temperature. Micrograph (a) shows the intergranular fracture behaviour at 725°C in air and (b) the transgranular behaviour in vacuum both at $\Delta K \sim 40$ MPa.m^{-1/2} [76].

Much effort has been expended in attempts to understand the deleterious role of oxygen on crack growth but the process remains incomplete although two contending mechanisms currently dominate thinking. The dynamic embrittlement model envisages that elemental oxygen diffuses into the alloy

grain boundary from the crack tip and produces cohesive weakening [77-79] in a manner analogous to S embrittlement. The concept is influenced by the experiments of Bricknell and Woodford [80, 81] on Ni containing different amounts of carbon exposed to oxidising and non-oxidising environments at 1000°C. In impure Ni, oxidation of the C content occurred under oxidising conditions even though Ni oxidation was prevented and extensive bubbles of CO₂/CO were produced together with significant embrittlement at 800°C. In the purest grade of Ni examined, few if any bubbles were present but substantial embrittlement still occurred suggesting (but not by Bricknell and Woodford) that intergranular cohesive weakening arose in the presence of dissolved elemental oxygen. These experiments were well conducted but the extrapolation of their findings to complex Ni-based superalloys should be undertaken with much caution. The erroneous feature of the argument is the assumption that elemental O could exist in such alloys at sufficient concentration to cause intergranular embrittlement in the presence of elements such as Cr, Ti and Al that have a strong affinity for O. As an example, the reaction between O and Cr would reduce the fractional partial pressure of O in equilibrium with chromia to as low as $\sim 10^{-32}$ at 650°C [82]. Chemical reaction to form oxides of these elements is much more energetically favoured than O segregation to the alloy grain boundaries and this forms the basis of an alternative mechanism of enhanced crack growth under oxidising conditions.

In this second model, it is envisaged that oxides of low fracture toughness form at the crack tip and penetrate along the alloy grain boundary ahead of it. Crack advancement occurs by the subsequent repeated cracking and reforming of these brittle oxide intrusions [83, 84]. Recent atom probe tomography and TEM studies support the presence of oxides penetrating the alloy from the crack tip [82, 85, 86]. An example of these oxide wedges, observed with TEM [82], is shown in Figure 10 for RR1000 tested at 625°C. This particular image shows the tip of the intrusion and the thermodynamic sequencing of the oxide layers that exists within it. By this is meant that oxides of lower thermodynamic stability, in this case CoO and NiO, are present at the central regions of the inclusion whereas the most stable oxides such as alumina and rutile are present adjacent to the alloy surface. Similar observations have been reported by Viskari et al. [86]. This is an unexpected result that indicates that individual layers can reduce the partial pressure of oxygen beneath them to their equilibrium dissociation pressure.

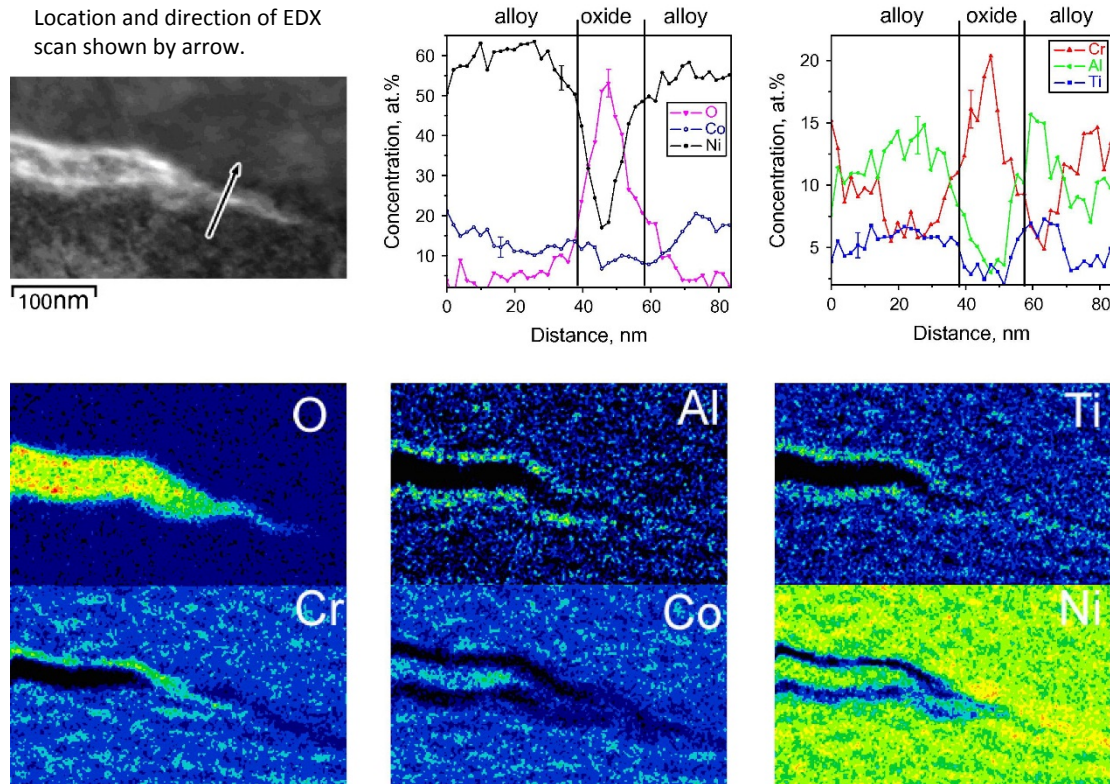


Figure 10: A bright field STEM image, EDX maps and line scans obtained ahead of the crack tip in RR1000, illustrating the presence and sequence of oxides. The relative concentration in the EDX maps increases from low to high in the sequence: black, blue, green, yellow, orange and red [82].

The principal oxides found by Kitaguchi et al. [82] that existed along the intrusion were NiO or Cr₂O₃ (chromia) depending on the detailed test conditions. In each case, the intrusion lengths were appreciably longer than would be expected, e.g. enhancement factors of ~17.6 for chromia and ~3.6 for NiO. This enhancement was considered to be a result of Stress Aided Grain Boundary Oxidation (SAGBO). The model of SAGBO that was developed [87] recognises that oxide formation will generally lead to an increase in volume and that this volumetric strain at the crack tip will fundamentally change the local stress distribution there. For loading conditions less than or of order that required for crack advance, i.e. near the threshold condition, K_{th} or ΔK_{th} , the normal stress at the crack tip will be compressive but will become less so with increasing distance from the crack tip and will be tensile at the tip of the wedge-shaped oxide intrusion [87]. This tensile stress leads to an increase in the outward flux of oxygen vacancies, and a corresponding increase in the inward diffusion of oxygen along the intrusion, producing an increase in oxide growth rate.

The ratio of growth rates of the intrusion of the stressed state to unstressed condition is given by the SAGBO enhancement factor, S_r , defined as [87]:

$$S_r = \left(\frac{\binom{\bullet}{l}_\sigma}{\binom{\bullet}{l}_o} \right) = \exp\left(\frac{\sigma_{it} \Omega_M (\Phi - 1)}{kT}\right) \quad (12)$$

Here, σ_{it} is the normal stress at the intrusion tip, k is Boltzmann's constant, T is absolute temperature, Ω_M is the volume of a metal atom in the alloy, calculated from molar volumes as $1.10 \times 10^{-29} \text{ m}^3$ for Ni and $1.22 \times 10^{-29} \text{ m}^3$ for Cr. The term $(\Phi - 1)$ represents the factor increase in metal volume on oxide formation. There is some uncertainty in the values to be used for Φ but taking those for the Pilling-Bedworth ratio is likely to introduce relatively little error for NiO and chromia. The calculated values of S_r at 650°C (923K) with $\Phi = 1.65$ for NiO and $= 2.07$ for chromia are shown in Figure 11 as a function of the normal stress at the intrusion tip.

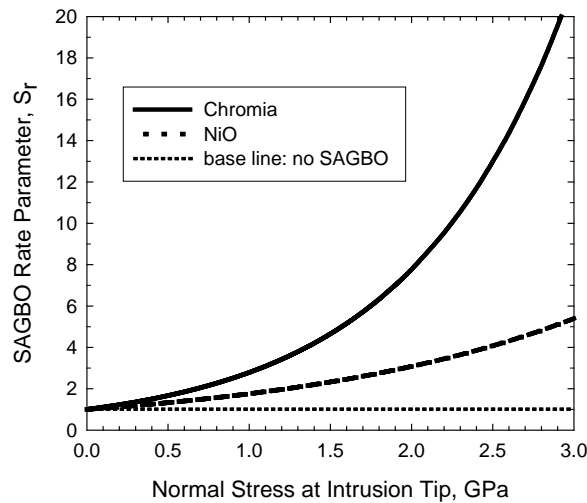


Figure 11: The dependence of the calculated SAGBO rate parameter, S_r , on tensile stress at the tip of the oxide intrusion for chromia and nickel oxide at 650°C [87].

The calculations show that the SAGBO parameter will be higher for chromia than for NiO and this is in agreement with the observations of Kitaguchi et al. [82] in terms of intrusion length. Furthermore, a value of ~ 2.5 GPa for the normal stress at the intrusion tip produces values of S_r similar to those found experimentally [82], again in terms of intrusion length (17.6 for chromia and 3.6 for NiO). This level of stress is not unreasonable for the strong RR1000 specimens examined by Taniguchi but it is also the case that microstructural changes or creep processes that weaken the alloy are likely to reduce the SAGBO enhancement factor [88] even though oxide intrusions form at the crack tip. Generally, it

can be appreciated from Figure 11, that in weaker alloys the experimental demonstration of the SAGBO phenomenon may prove elusive.

Conclusions

Chromia-forming Ni-based superalloys are widely used as structural components and, particularly, as turbine discs at temperatures up to $\sim 750^{\circ}\text{C}$. There is increasing demand for operation at even higher temperatures, say to 800°C , and this places emphasis on the oxidation resistance of the alloys and the interaction of oxidation with mechanical properties. This overview examines these aspects. It is shown that, in addition to the formation of a surface layer of chromia, typical alloys are also internally oxidised to form, principally, alumina and this results in the development of intergranular intrusions of alumina of acicular appearance, discrete intragranular precipitates and a surface zone depleted of γ' . Shot-peening the alloy surface does not affect these broad trends although detailed differences in the morphology of the internal oxides and recrystallization of the surface regions can occur.

Low alloy titanium content appears to have a beneficial effect on oxidation resistance whereas high contents result in the formation of discrete crystals of rutile above the surface chromia layer. This has been known for many years but more recent studies have shown that titanium also increases the growth rate of the chromia layer itself. This is attributed to the doping of the Cr sub-lattice with 4-valent Ti ions and the creation of extra vacancies that increase Cr diffusion rates within its oxide and, consequently, oxidation rates. This oxidation enhancement is highest in the early stages of oxidation but declines with time as the near-surface concentration of Ti becomes depleted.

There is currently a lack of clarity on the influence of applied loads on oxidation behaviour. This arises because of uncertainty in the stress levels, and their sign, present in the oxide layers. Allowances have not been made for intrinsic growth stresses, the redistribution of the applied load between oxide and alloy and, in some cases, cooling stresses. As a result, quantitative assessment is not possible although tensile cracking and localised oxide spallation are evident and both lead to enhanced oxidation rates. Similarly, the cracking of the surface oxides and/or the immediate sub-surface alumina penetrations provide nucleation sites for fatigue crack development.

The growth rate of fatigue cracks in air is very much larger than under inert atmospheres. Recent TEM observations indicate that this oxygen effect is associated with the development and cracking of oxide intrusions emanating from the crack tip and penetrating along the alloy grain boundary. The growth rate of these intrusions can be an order of magnitude larger than expected and this may be

evidence of stress aided grain boundary oxidation (SAGBO) although experimental data are limited. It is expected that the formation of typical oxides (NiO, chromia) at the crack tip will perturb the stress field such that out-of-plane normal stresses there can become compressive and then rise to tensile at the tip of the intrusion. This modified stress distribution has not yet been quantified but has been incorporated into a recent model of SAGBO.

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List of Figure Captions

Figure 1: Oxidation of Ni-based superalloy, ME3, exposed to air at 815°C for 2020 hours, showing an external layer of Cr₂O₃ externally with an internal oxidation zone of Al₂O₃ [15]. See text for fuller discussion.

Figure 2: Arrhenius plot of k_p values from the literature of similar Ni-based superalloys used for rotor disc applications and typical expected behaviour for chromia formation on a high chromium containing austenitic stainless steel (dashed line) [5, 7, 11, 12, 31, 32]. Image modified from Cruchley et al. (2013) to include data taken from [17].

Figure 3: Plot of chromia thickness measured on RR1000, with and without shot-peening, at 750°C compared with Ti-free austenitic steel/pure chromium. Oxide measurements for both conditions were normally distributed and error bars are shown as ± 1 standard deviation. Hollow squares are measurements for shot-peened RR1000, filled circles are polished RR1000 [17].

Figure 4: The variation of the enhancement ratio of the chromia growth rate due to Ti doping with oxide thickness for 700°C, 750°C and 800°C, in a) polished RR1000 and b) shot-peened RR1000 [17].

Figure 5: Comparison of the oxidation damage formed on a) polished RR1000 and b) shot-peened RR1000 oxidised for 2000 hours at 700°C [17].

Figure 6: Backscatter electron images of oxidation damage on fine-grained RR1000 after 200 hours at 700°C under hold stresses of (a) + 600 MPa, (b) 0 MPa and (c) -600 MPa [56].

Figure 7: BSE image of an etched pre-oxidised (2000h, 700°C) specimen that failed after 1.9×10^6 cycles during room temperature 4-point bend at a maximum applied stress of 800 MPa, showing cracking of the external and intergranular internal oxide along with the presence of both a γ' denuded zone and carbide dissolution zone [68].

Figure 8: Comparison of the mean notched fatigue lives at 704°C of the unexposed ME3 specimens and ME3 specimens with a) prior exposures in air and b) alternative conditions as marked. Diamonds are each test result, clearly showing that the removal of all of the environmental damage (50 µm removal) leads to a full restoration in fatigue life [15].

Figure 9: Main graph shows crack growth rates versus ΔK in the LHSR alloy tested under 3-point bend at 650°C (black symbols) and 725°C (red symbols) under air and vacuum. Increased rates were observed in air at both temperatures but were greater at the higher temperature. Micrograph (a) shows the intergranular fracture behaviour at 725°C in air and (b) the transgranular behaviour in vacuum both at $\Delta K \sim 40 \text{ MPa.m}^{-1/2}$ [76].

Figure 10: A bright field STEM image, EDX maps and linescans obtained ahead of the crack tip in RR1000, illustrating the presence and sequence of oxides. The relative concentration in the EDX maps increases from low to high in the sequence: black, blue, green, yellow, orange and red [82].

Figure 11: The dependence of the calculated SAGBO rate parameter, S_r , on tensile stress at the tip of the oxide intrusion for chromia and nickel oxide at 650°C [87].

List of Table Captions

Table 1: Compositions of some representative alloys [1-4].