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What is the Role of Rhenium in Single Crystal Superalloys?

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Abstract. Rhenium plays a critical role in single-crystal superalloys –its addition to first generation alloys improves creep life by a factor of at least two, with further benefits for fatigue performance. Its use in alloys such as PWA1484, CMSX-4 and Rene N5 is now widespread, and many in this community regard Re as the ‘magic dust’. In this paper, the latest thinking concerning the origins of the ‘rhenium-effect’ is presented. We start by reviewing the hypothesis that rhenium clusters represent barriers to dislocation motion. Recent atom probe tomography experiments have shown that Re may instead form a solid solution with Ni at low concentrations (< 7 at.%). Density functional theory calculations indicate that, in the solid solution, short range ordering of Re may be expected. Finally, Re has been shown to diffuse slowly in the γ -Ni phase. Calculations using a semi-analytical dislocation climb/glide model based upon the work of McLean and Dyson have been used to rationalise the composition-dependence of creep deformation in these materials. All evidence points to two important factors: (i) the preferred partitioning of Re to the γ phase, where dislocation activity preferentially occurs during the tertiary creep regime and (ii) a retardation effect on dislocation segments at γ/γ' interfaces, which require non-conservative climb and thus an associated vacancy flux.

1 Introduction

Rhenium additions are essential for the high-temperature properties of single-crystal superalloys. The higher performance of superalloys have resulted in longer-lasting components and higher fuel efficiencies. Re has been so important to the industry that single-crystal superalloys are commonly grouped into generations based on their Re content: first-generation Re-free alloys gave way to second-generation alloys in the 1990s, containing 2-3 wt.% Re, superseded by third-generation alloys in the 2000s, containing 5-6 wt.% Re.

Since the advent of third-generation alloys, engine manufacturers have opted to reduce Re content in their alloys, primarily due to cost concerns: Re is one of the most expensive transition metals. Its average price between 2000 and 2010 has been US\$ 3000/kg, and the price is likely to increase in future. The high price means that Re alone can be, depending on the alloy, responsible for over half of the raw material cost needed to produce turbine blades. In addition, the use of Re also poses a risk due to the fact that the majority of Re reserves are clustered in a limited number of countries.

The high price of Re, and its importance as a strategic element, underline the need for the community to understand the role of Re on the high-temperature properties of Ni-based superalloys, with the objective of designing new grades of alloys with lower Re content or completely

Re-free. In this paper, after describing the beneficial effects of Re additions to the high-temperature properties of Ni-based superalloys, we examine all the available experimental evidence and discuss its significance in conjunction to recent modelling efforts.

2 The rhenium-effect

The earliest evidence of Re additions to the Ni-based superalloys can be found in a 1975 patent submitted by Smashey [1]. The objective of Smashey’s work was to improve the high-temperature properties of a unidirectionally solidified eutectic Ni-based superalloys by precipitating fibrous carbide phases aligned with the growth direction [1]. In these high-C superalloys, Re (contrary to W) is not a carbide former and can therefore be used to strengthen the matrix phase and promote the partitioning of elements such as Ta and V to the γ' phase [1]. Research efforts on investigating the effect of Re additions on carbide-strengthened directionally-solidified Ni-based superalloys continued through the beginning of 1980s [2], when the advent of single-crystal Ni-based superalloys changed the field dramatically.

The first attempts to add Re to single-crystal specimens were made by Giamei and reported in [3]. In this work, Giamei modified MAR-M200 (Ni-5Al-2Ti-1Nb-9Cr-12W wt.%) by adding Re instead of W, keeping the W+Re weight concentration fixed. From the base alloy, hereafter called alloy 0Re, three other compositions were obtained:

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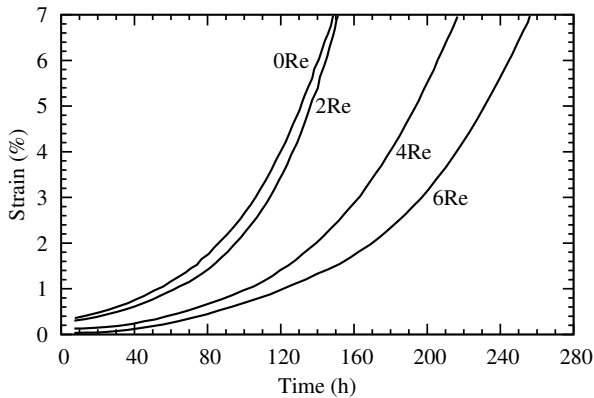


Figure 1: Creep curves for alloy 1444 and Re-containing derivatives at 899 °C under a load of 380 MPa [3]

2Re, 4Re and 6Re, containing 2, 4 and 6 wt.% Re respectively. These compositions were grown as single-crystal along the [001] direction, using a modified Bridgman technique. The resulting specimens were tested in creep at 899 °C under a load of 380 MPa, revealing the creep behaviour that can be observed in Figure 1. Since this initial report, the dramatic beneficial effects of Re additions on tertiary creep deformation has been confirmed in various single-crystal alloys [4–6], and is commonly referred to as the rhenium-effect.

Beside the improved tertiary creep lives of Re-containing alloys, Re additions have been observed to considerably reduce the coarsening kinetics of γ' precipitates [7], primarily due to the fact that Re strongly favours partitioning to the γ phase. The strong partitioning of Re to the γ phase has been observed in most single-crystal Ni-based superalloys with high γ' volume content [4, 8–10], and it is supported by first-principles calculations showing Re substitutional defects are unstable in the γ' phase [11]. As the γ' precipitates coarsen, Re has to diffuse ahead of the γ/γ' interface [7]. The strong partitioning of Re to the γ phase, coupled with the fact that Re is the slowest-diffusing transition metal in Ni [12–14], means that γ' coarsening rates are severely limited by the presence of Re. This is also beneficial, as slower γ' coarsening kinetics result in longer rupture lives at elevated temperatures, when rafting (i.e. directional coarsening) becomes problematic [7]. This behaviour was also later confirmed and discussed in great detail by the work of Seidman *et al* [15–18].

Whereas the effect of Re on the γ' coarsening kinetics can be clearly attributed to the partitioning characteristic of Re and its diffusion coefficient in Ni, the strengthening effect of Re additions on tertiary creep has been the subject of wider discussion, which we hope to summarise here.

3 Summary of Evidence

3.1 One-dimensional atom probe

Blavette and co-workers pioneered the use of atom probe analysis to study the behaviour of Re in Ni-based superalloys [4, 19]. In this work, the one-dimensional version of the tool (1DAP) was used to examine the solute distribution in PWA 1480, CMSX-2, PWA 1480-Re (with a

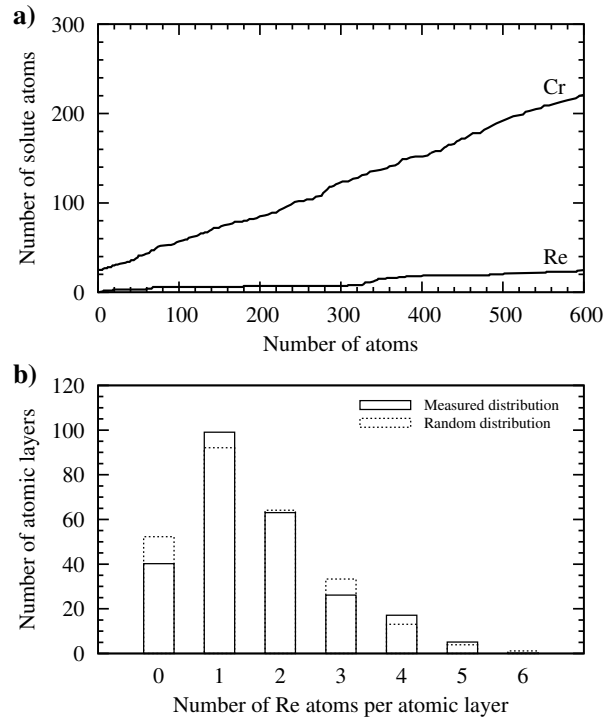


Figure 2: a) cumulative profile of Re showing spatial irregularities and b) histogram of the number of atomic layers containing a given number of Re atoms [4]

nominal 4 wt.% Re and no W) and CMSX-2-4Re (containing 4 wt.% Re and 4 wt.% W). Creep tests carried out at 850 °C and 500 MPa revealed a 30–40 % increase in rupture life associated with the addition of Re. The atom probe analysis confirmed that the Re atoms strongly partition to the γ matrix phase, leaving the γ' precipitates depleted of Re content. The Re content in the γ phase ranged from 2.87 at.% to 3.71 at.% for CMSX-2-4Re and PWA 1480-Re respectively.

The atom probe also revealed irregularities in the spatial distribution of Re: cumulative profiles (see Figure 2a) of Re desorption indicated the presence of areas with higher Re content [4]. In order to quantify the spatial distribution of Re, Blavette and co-workers plotted a histogram of the number of atomic layers containing a set number of Re atoms, and compared that to a Poisson's random distribution (see Figure 2b). A larger number of atomic layers contained 1, 4 or 5 Re atoms than expected from a random distribution, while fewer atomic layers contained 0, 2, 3 or 6 Re atoms than expected from a random distribution [4]. It is arguable whether this is evidence for *either* short range ordering *or* clustering of Re atoms in the γ phase of Ni-based superalloys. The authors proposed the evidence collected proved the presence of Re clusters, 1–1.5 nm in size, that contained up to 90 at.% Re [4, 19], suggesting this would be expected when considering the large miscibility gap in the established Ni-Re binary phase diagram [20, 21]. It follows that these clusters would act as powerful barriers against dislocation glide when compared to distributed Re atoms in solid solution with Ni.

Following the initial atom probe work by Blavette *et al*, Wanderka and Glatzel also used the 1DAP to study the distribution of Re in CMSX-4 [22]. Also in this case, cu-

mulative profiles of Re revealed fluctuations in Re distribution, and a comparison of the Re distribution of a Poisson's distribution revealed a lower than expected number of atomic layers containing 0 and 3 Re atoms, while a higher than expected number of atomic layers contained 1 or 2 Re atoms [22]. The authors indicated this as evidence of clusters, suggesting these are likely to be composed by 5 Re atoms, 3 in one atomic layer, with an additional atom in the upper and lower layer [22]. The pioneering work in these early studies was limited by the capabilities of the 1DAP.

3.2 Three-dimensional atom probe

The advent of the three-dimensional atom probe (3DAP) allowed for a more complete investigation of the distribution of Re in the superalloys. Warren *et al* used a 3DAP to study the Re distribution in the γ phase and at the γ/γ' interfaces of RR3000, a Re-containing third-generation Ni-based superalloy [8]. Along with the strong partitioning of Re to the γ phase, the authors found Re-enriched areas near the interfaces of the secondary and tertiary γ' precipitates, on the γ phase side [8]. These would be expected since, upon cooling from higher temperatures, the secondary γ' precipitates grow and tertiary γ' precipitates are expected to form in the middle of the γ channels, ejecting Re ahead of their interfaces. The authors found no evidence of Re clusters in their samples [8].

Rüsing *et al* argued for the presence of Re clusters in their samples of Re31, a model quaternary alloy [23]. In their work, a long and thin volume ($1.5 \times 1.5 \times 90$ nm) was divided into slices 0.5 nm thick in order to obtain a one-dimensional concentration profile, revealing slices where the local Re concentration exceeded 2σ (see Figure 3a) [23]. As this approach would not conclusively demonstrate the presence of clusters, the authors adopted the wavelength dependent filtering method. This approach, based on Fourier analysis, can be used to investigate the heterogeneity of the data, without providing any indication of the shape and size of these heterogeneities. This analysis indicated that Re is heterogeneously distributed with a principal wavelength of 20 nm [23] (see Figure 3b), but the heterogeneities observed may also be due to a non-homogeneous distribution of Re.

Recently, more 3DAP studies have been carried out to determine whether Re clusters may be present in the γ phase of Ni-based superalloys [24, 25]. A number of Ni-X binary alloys (where X is Ta, W, Re, Ir and Pt), with 10 wt.% nominal solute content, were analysed using a direct-flightpath 3DAP. A maximum-separation friends-of-friends algorithm was used to evaluate the presence of Re clusters in the data. The results were compared to simulated atom probe datasets [24] and randomised versions of themselves [25] (see Figure 4). No clusters were detected. The same approach was adopted to study the distribution of Re in the γ phase of CMSX-4 where, similarly to the binary alloys, clusters were not found [24, 25]. A sensitivity analysis of the approach on simulated 3DAP datasets revealed that sparse clusters ($3.5 \cdot 10^{24} \text{ m}^{-3}$) with

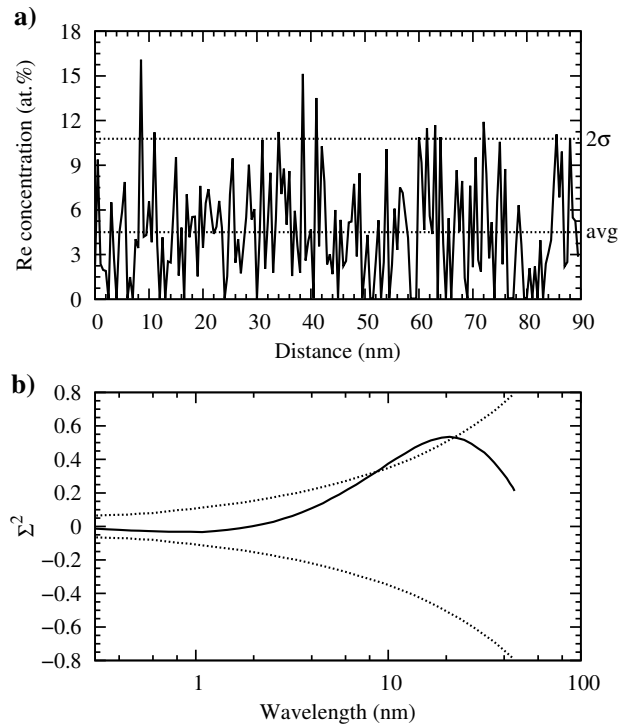


Figure 3: a) compositional profile of Re along $1.5 \times 1.5 \times 90$ nm volume and b) results of the wavelength dependent filtering method on a $1.5 \times 1.5 \times 90$ nm volume [23]

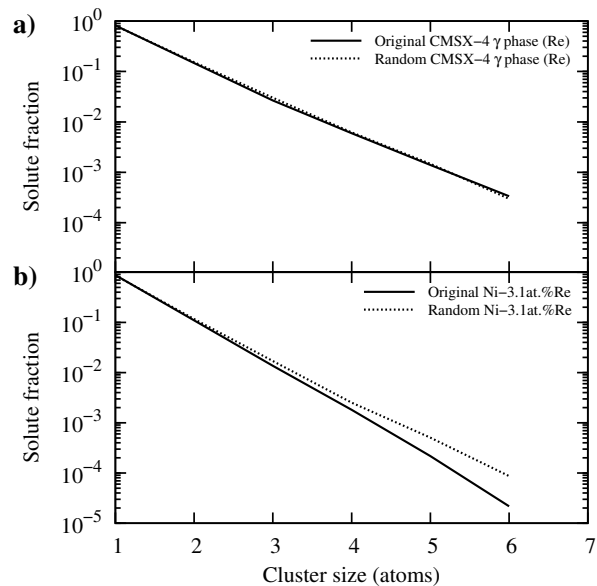


Figure 4: The results of the maximum-separation friends-of-friends algorithm on a) the γ phase of CMSX-4 and b) a Ni-Re binary alloy, showing Re distribution to be consistent with a solid solution [25]

1 nm diameter should be evident, even when considering the detection efficiency and spatial resolution of the tool [24, 25].

3.3 Other experimental evidence

Ge *et al* studied the distribution of Re in the 2nd-generation single-crystal superalloys DD6 using high angle annular dark field (HAADF) and energy dispersive spectroscopy (EDS) on a scanning transmission electron microscope

(STEM) [26]. Chemical analysis across the γ/γ' interface revealed no Re enrichment in the un-crept specimens, whereas Re enriched areas were present in the crept specimens, suggesting these form only after considerable γ' particle coarsening occurring during creep tests [26]. The authors also proposed that high-contrast areas observed near the γ/γ' interfaces after the creep tests could be evidence of Re/W clusters 1-2 nm in diameter [26]. However, these high-contrast areas could also be due to expected compositional fluctuations.

The presence of Re clusters in the Ni lattice was also investigated using extended x-ray absorption fine spectroscopy (EXAFS) [27, 28]. In EXAFS, the local atomic structure surrounding atoms of a specific element can be probed by examining the oscillations in x-ray absorption coefficient immediately above the absorption edge of that element. In this case, the L_3 edge of Re was used to detect the local environment surrounding the Re atoms in the specimens. Both studies clearly indicate that Re-Re first neighbour pairs are unlikely in the Ni lattice, as the data clearly indicates that Re is coordinated by 12 Ni atoms [27, 28].

Diffusion couple studies have conclusively shown that Re is one of the slowest-diffusing transition metals in Ni [12, 13]. When compared to other solute elements commonly added to the superalloys, Re is shown to diffuse almost an order-of-magnitude slower than W [12]. The general trends observed showed that elements sitting in the middle of the transition metal series diffuse slower than the elements at the edges of the transition metal series, despite being smaller in size [13].

4 The modelling contribution

4.1 Density functional theory

Ab initio density functional theory (DFT) proved instrumental in understanding and explaining why elements in the middle of the $4d$ and $5d$ transition metal series diffuse slower than elements at the edges of it [14, 29]. In contrast to the outer elements, such as Hf and Au, elements in the middle of the $5d$ transition metal series, such as Re, form directional d bonds with the surrounding Ni atoms. These directional d bonds are less compressible, and result in a higher energy barrier for the Re to exchange position with a neighbouring vacancy [14, 29]. This explains the higher activation energy for diffusion observed for the smallest atoms in the middle of the $4d$ and $5d$ transition metal series [14, 29].

Calculations of binding energies of Re-Re 1st and higher-order pairs using DFT reveal that Re clusters would be energetically unfavourable [30]. Investigation of the Ni-Re 0K formation energy hull indicates that Ni_4Re ($D1_a$) and $NiRe_3$ ($D0_{19}$) should be stable at absolute zero [31, 32]. High-throughput DFT calculations were also used to parameterise a cluster expansion to determine the effect of configurational entropy on the free energy of the relevant phases through a Monte Carlo approach [31, 33]. Results indicate that the Ni_4Re ($D1_a$) intermetallic would

not be stable above 600-700 °C due to the effects of configurational entropy. Including vibrational entropy would result in a further reduction of the Ni_4Re ($D1_a$) solvus temperature [33]. Considering the activation energy of Re diffusion in Ni, it is reasonable to expect this phase not to form except after very long heat treatments. Above the solvus temperature of the Ni_4Re ($D1_a$) intermetallic, the Monte Carlo simulations predict that relatively low concentrations of Re (0-7 at.%) can form a solid solution with Ni [33]. Simulated x-ray diffuse scattering data reveals remnant short-range ordering of Re in Ni, which gradually reduces as temperature is increased due to configurational entropy [33]. Clustering of Re atoms in the Ni lattice is predicted to be energetically unfavourable at all temperatures [33].

4.2 Microstructure evolution models

Yoon *et al* investigated the effect of Re additions on the temporal evolution of Re-free and Re-containing model Ni-Al-Cr superalloys by comparing experimental data, obtained using 3DAP and TEM, to calculations made with the Umantsev–Olson model and PrecipiCalcTM [16, 17]. In their alloys, Ni-8.5Cr-10Al, Ni-8.5Cr-10Al-2Re and Ni-8.5Cr-10Al-2W (at.%), the authors did not observe any bow-wave or monotonic Re-enrichment at the γ/γ' interfaces, at all ageing times [16, 17]. This is possibly due to the fact that specimens were water quenched, resulting in a smaller migration of the γ/γ' interface upon cooling. Other factors, such as a lower volume fraction of γ' phase compared to single-crystal commercial Ni-based superalloys, and a finer distribution of spherical precipitates may have also played a role on the absence of a monotonic Re-enrichment at the γ/γ' interfaces. Coarsening of the γ' precipitates during ageing did not cause the formation of a Re-enrichment at the γ/γ' interfaces, although both experiments and simulations showed that the addition of Re resulted in a considerable deceleration of coarsening kinetics [16, 17].

Warnken used phase field simulations to establish whether the Re enrichment observed close to the γ/γ' interfaces of uncrept CMSX-4 specimens [24, 25] should be attributed to the coarsening of the γ' secondary precipitates during heat treatment or to the growth of the γ' secondary precipitates when cooling from service temperature. Simulations were made using the MICRESSTM, coupling the Thermo-TechTM TTNI version 6 database and the NIST Ni superalloy mobility database to accurately represent the thermodynamics and kinetics of CMSX-4. The migration of the γ/γ' interface upon cooling from 950 °C to 650 °C was shown to be strongly dependent on cooling rates [25]. The migration of the γ/γ' interface upon cooling resulted in a bow-wave ahead of the interface, comparable to that observed in CMSX-4 using the 3DAP [25]. Subsequent heating simulations caused the γ/γ' interface to retreat and the Re composition close to the γ/γ' interfaces to re-homogenise [25]. It was concluded that no Re enrichment would be present at service temperature near the γ/γ' interfaces of CMSX-4 [25].

5 Discussion: a modern view of the role of rhenium in superalloys

A number of mechanisms contribute to the beneficial effects of Re additions on the high-temperature properties of Ni-based superalloys. Perhaps the most important of these is the partitioning behaviour of Re to the γ matrix phase. This has been observed in most Re-containing single-crystal commercial Ni-based superalloys. Whenever the partitioning of Re to the γ phase was not observed, the high-temperature properties were unaffected by small Re additions [34]. Although only relatively small amounts of Re (0-2 at.%) are sufficient to produce a two-fold increase in creep life, contributing to its fame as a ‘magic dust’, it is important to note that most of the Re will partition to the minority phase (γ), where the resultant concentration of Re may be as high as 7 at.%. Thanks to its partitioning behaviour and its low diffusion coefficient, small Re additions are very effective in retarding the γ' coarsening and rafting kinetics, thereby extending creep life at the higher temperatures ($> 1000^\circ\text{C}$). This, however, is not sufficient to explain the improved tertiary creep regime properties associated with the addition of Re to single-crystal Ni-based superalloys.

Therefore, Re must play a role on the deformation processes occurring in the γ phase during the tertiary creep regime. Under these conditions, dislocations glide in the γ phase and stop at the γ/γ' interfaces, since stresses are not high enough to allow penetration and cutting of the γ' precipitates. To continue on their path, these dislocations must move along the γ/γ' interface by a thermally-activated climb process [35, 36]. Re must act as an effective barrier to dislocation motion. This process cannot be dismissed as a simple solute-strengthening mechanism as it has been shown that Re is a poor solid-solution strengthener in Ni [37, 38]. Furthermore, single solute atoms do not pose a considerable barrier to dislocation glide at elevated temperatures. Small Re clusters could stop dislocation glide in the γ phase, but experimental evidence of their presence is not conclusive. Small inhomogeneities of the Re distribution have been observed in the γ phase of a variety of alloys, but these may be a natural occurrence in a multi-component solid-solution where composition is highly inhomogeneous due to the microstructure and manufacturing route. Examination of simpler Ni-Re binary specimens provided no evidence of the presence of Re clusters, despite state-of-the-art cluster-detection algorithms were used. Similar conclusions were drawn when examining the Re distribution in the γ of CMSX-4.

Density functional theory conclusively shows that Re-Re first-neighbour pairs and small first- and second-neighbour clusters would be unstable in Ni. Further studies of the formation energy convex hull reveal an intermetallic with chemical formula Ni_4Re and crystal structure D1_a may be stable at absolute zero, but configurational and vibrational entropy effects mean that this intermetallic would only form below $600\text{-}700^\circ\text{C}$, or not form at all due to the slow diffusion of Re. The fact that this intermetallic may only form under specific cooling con-

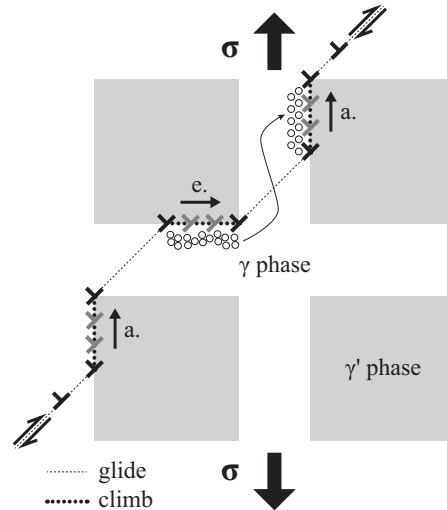


Figure 5: A 2D schematic of the climb-assisted glide process occurring in tertiary creep and the resulting vacancy flow from the horizontal to the vertical channels [35]

ditions may explain some of the conflicting atom probe results; regardless, calculations unequivocally show such a phase would not be present at service temperature, and cannot explain the anomalous strengthening effect caused by Re additions on tertiary creep.

Since atoms of Re cannot pose a sufficient barrier to dislocation glide, and Re clusters are not present at service temperature, the reason for the rhenium-effect must be sought elsewhere. In tertiary creep, the motion of dislocations is controlled by the climb rate at the γ/γ' interfaces. Climb is a thermally activated process that requires solute atoms to exchange position with vacancies at the dislocation core. In addition, depending on the climbing direction, a dislocation may act as a vacancy emitter or absorber (see Figure 5). Therefore, the surplus of vacancies in areas where the climb of dislocations results in the generation of vacancies have to diffuse to areas where the climb of dislocations results in the absorption of vacancies (see Figure 5). As a slow diffuser, Re would act to slow both of these processes, reducing the overall climb rate. A semi-analytical dislocation climb/glide model based upon the work of McLean and Dyson has been used to test this hypothesis [35]. The results indicate that the arithmetic average of the solvent and solute diffusion coefficients, weighted by their atomic concentrations, is a good first order approximation for the effective vacancy diffusion coefficient, and qualitatively captures the effects of γ phase composition on tertiary creep.

6 Conclusions

1. Small additions of Re have been shown to dramatically improve the high-temperature creep properties, specifically within the tertiary creep regime.
2. The mechanisms underlying the anomalous creep strengthening contribution of Re to the superalloys are still unclear.
3. It is well known that Re reduces the coarsening and rafting of the γ/γ' microstructure, but this alone is

not enough to explain the improved tertiary creep behaviour of 2nd- and 3rd-generation alloys.

4. Initial evidence suggested Re may form clusters in the γ phase of Ni-based superalloys. These clusters could act as barriers to dislocation motion.
5. The theory that Re forms clusters is not supported by the latest 3DAP and EXAFS result which, instead, suggest that Re forms a solid solution with Ni.
6. Electronic structure calculations have shown that small Re clusters are unstable in Ni. An intermetallic compound with formula Ni₄Re may be stable from absolute zero up to intermediate temperatures, but Re is expected to form a solid solution with Ni above 600–700 °C. The Ni-Re solid solution may be characterised by some remnant short-range ordering.
7. The consolidated hypothesis is that creep deformation at elevated temperature is controlled by climb-assisted glide, where the temperature-activated climb mechanism is rate-controlling.
8. Re additions must act to slow down the necessary flow of vacancies and other atomic-scale diffusion mechanisms required for climb to happen. This, together with the fact that most Re atoms partition to the γ phase, is enough to explain the beneficial effects of Re on creep life in the tertiary creep regime.
9. Further work is necessary to place this hypothesis on a firm quantitative basis, and explicitly link the chemical composition of the γ phase to the creep life in the tertiary regime.

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