PROPERTIES OF A HFC-REINFORCED NICKEL-BASED SUPERALLOY IN CREEP AND OXIDATION AT 1100°C

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A cast Ni-based superalloy strengthened by hafnium carbides was cast and characterized at 1100°C in creep and in hot oxidation. The as-cast microstructure, composed of a dendritic matrix and of eutectic script-like hafnium carbides (HfC) imbricated with matrix in the interdendritic spaces, is very favorable for high mechanic properties at high temperature. This was confirmed by a three-point flexural creep test carried out at 1100°C under 20 MPa. The slow parabolic mass gain kinetic showed that this alloy is very resistant against oxidation at 1100°C in air. The HfC carbides remained stable during these tests. The oxidation behavior was chromia-forming. This alloy appeared ready to be used for structural applications at a temperature reachable by only $\gamma \gamma'$ Ni-based single-crystals or ODS superalloys.

Keywords: Ni alloy, HfC carbides, creep resistance, high temperature oxidation.

In front of the demand of superalloys for applications at increasing temperatures the microstructure solutions begin lacking to offer a complete set of good properties at high-temperature (HT): toughness, mechanical strength, good behavior in hot oxidation and corrosion, and furthermore workability with complex shapes [1]. Obtaining coarse grains for HT strength by keeping complex shape feasibility leads to a casting process. The obtained microstructures are necessarily polycrystalline. The presence of grain boundaries and, consequently of intergranular reinforcing particles (e.g. carbides), limit the fusion start temperatures. This does not allow applying optimized heat treatments to achieve high mechanical strength at very high temperature such as the high volume fractions of Ni₃Al precipitates in Ni-based γ/γ single-crystals.

Metal carbide (MC) carbides are among the most efficient particles for strengthening grain boundaries. The most common MC carbides are the tantalum carbides (TaC). The script-like shape of TaC in cast Co-based alloys are very useful for resisting creep at HT. Unfortunately they are not so stable in Ni-based alloys [2]. Hafnium carbides are other possible MC which may precipitate in Co-based alloys with the same morphology as TaC [3]. This second type of MC carbide is also script-like and very stable at HT in Co-based alloys [4].

Combining a Ni-based matrix rich in Cr and HfC carbides in a cast alloy may be an interesting solution for constituting complex-shaped pieces resisting both chemical aggressiveness from gases (for example, hydrogen at high temperature and pressure [5, 6]) and melts (intrinsic property of a Ni–Cr base) and mechanical solicitations (austenitic network of Ni matrix and interdendritic reinforcement by HfC), at HT. This is what was attempted in this work with a Ni–25Cr–0.25C–3.7Hf alloy.

Method of investigation. 40g of this alloy, named "NCCHf", were cast from pure elements (Alfa Aesar, > 99.9 wt.%), in the water-cooled copper crucible of a High Frequency Induction furnace, under 300 mbars of pure Ar. A Ni(bal.)–25Cr–0.25C alloy, named "NCC", was added to the study in order to specify the effect of Hf on the oxidation behavior. This second alloy was elaborated in the same conditions as NCCHf.

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Three types of samples were prepared. The first compact one was kept for the metallographic study of the as-cast microstructure of NCCHf. The second sample, an elongated parallelepiped, was subjected to the creep bending tests. The third one, a square-based parallelepiped, was destined to oxidation test in a thermobalance. A sample for oxidation test was prepared from NCC, as a base of comparison.

The metallographic sample was embedded then ground with 250 to 1200-grit SiC papers. Thereafter it was polished with 1 μ m hard particles for obtaining a mirror-like state. The creep sample 1×2×15 mm, was cut then polished with papers up to 1200-grit with final strips in the length direction. This parallelepiped, with homogeneous thickness and width (tolerance: ±0.01 mm), was thoroughly examined by microscopy to be sure of the absence of any defects able to weaken it. The dimensions of the samples for HT oxidation tests were 8×8×3 mm. They were polished with 1200-grit SiC papers with smoothing of edges and corners.

The metallographic samples were observed using a scanning electron microscope (SEM, JEOL JSM6010LA). Observations were done under 20 kV in the back scattered electrons (BSE) mode, with the \times 125; 250; 500 and 1000 magnifications. The obtained chemical composition was measured using the energy dispersive spectrometry (EDS) device attached to the SEM. The different phases were specified by EDS spot analyses. Indentations were carried out for the two alloys using a Testwell Wolpert apparatus, to measure their Vickers hardness under a 30 kg load. In each case three indentations were performed; average value and standard deviation values were calculated.

NCCHf was submitted to bending tests at 1100°C under constant load, according to the symmetrical three-point flexural method, using a dilatometer SETARAM TMA92-16.18 modified to allow 3-points flexural testing. The space between the two bottom supports was 12 mm and the maximal possible deformation was about 1.4 mm (height of these supports). The load was applied to the sample by the upper support, inducing a 20 MPa tensile stress in the middle of the bottom side of the sample. Its value was calculated for each sample by taking into account the exact values of their thickness and width. The load was progressively applied at room temperature, until the vised value. The heating was then done at a constant rate of 20°C min⁻¹. The isothermal stage was more or less long, depending on the behavior of the tested sample. The downward vertical displacement of the upper support was recorded every 60 s. At the end of the experiment a cooling was applied at a rate of -20° C min⁻¹.

NCCHf and NCC were both tested in oxidation at 1100°C during 46 h, using a thermo-balance (SETARAM TG-92), in dry synthetic air. The heating was achieved at 20° C min⁻¹, and the post-isothermal stage cooling was done at -5° C min⁻¹. X-ray diffraction (XRD) was carried out to characterize the oxides scales. The oxidized samples were thereafter covered by a thin gold coating then by a thick electrolytic nickel coating to protect the oxide scales during cutting. The coated oxidized samples were cut, embedded and polished. They were examined using the SEM in BSE mode. The sub-surfaces and the oxide scales were subjected to spot EDS analysis.

Results and discussion. EDS analyses showed that the vised chemical compositions were well obtained. The NCCHf microstructure is illustrated in Fig. 1. HfC is obviously the single carbide phase present. This demonstrates that Hf is a much more carbide-former element than Cr in nickel alloys while this was the contrary for Ta in the same base (Ta less carbide-former than Cr in nickel alloys [2]). All the Hf atoms are present as HfC since no Hf was detected in the matrix by EDS. Thanks to the atomic ratio Hf/C equal to 1 obtained by choosing 0.25 wt.% C and 3.7 wt.% Hf, all the C atoms were also involved in HfC. HfC are script-like shaped and located in the interdendritic areas where they form an eutectic with the matrix, which let us think of high mechanical properties at HT. The Vickers indentations led to 194 ± 9 Hv_{30kg} for

NCCHf, i.e. higher than NCC (163 ± 3 Hv_{30kg}). This hardness close to 200 allows expecting the respectable strength at high temperature.



Fig. 1. As-cast microstructure of NCCHf (SEM/BSE mode); visualization of matrix ("M") and the HfC carbides ("HfC"): *a* – general view; *b* – details.

The deformation curve (displacement of the upper support) was obtained for NCCHf during the flexural 3-point creep test (Fig. 2) together with a scanned view of the deformed sample. The deformation rate in the secondary creep stage is of only $2 \ \mu m \cdot h^{-1}$ which is very low for a nickel alloy at this temperature for a resulting tensile stress of 20 MPa.



Fig. 2. Flexural creep deformation of the NCCHf alloy at 1100°C under 20 MPa (σ_{max}).

The thermogravimetry test carried out at 1100°C in synthetic dry air led to a parabolic mass gain kinetic (Fig. 3). Compared with the mass gain curve obtained for the reference NCC alloy – the good behavior of which is well-known (among the best chromia-forming alloys in this field) – this result is very interesting. Indeed, NCCHf did not present any mass gain jump (better adherence of the oxide scale on the substrate due to Hf) and its mass gain rate is lower (see Table). The transitory linear constant K_1 (slope of the mass gain curve at the beginning of the isothermal stage) and the parabolic constant K_p classically determined (slope of the mass gain curve when plotted versus the square root of time) are effectively a little lower than for NCC (see Table): 14 against $18 \cdot 10^{-8}$ g·cm⁻²·s⁻¹, and 4 against $6 \cdot 10^{-12}$ g²·cm⁻⁴·s⁻¹, respectively. The comparison was also done by taking into account the mass loss by chromia volatilization (small loss of Cr₂O₃ by re-oxidation in volatile CrO₃). For that the mass gain files were plotted according to the { $m \cdot (dm/dt) = f(-m)$ }-method [7] which allows the determination of the volatilization constant K_v (K_v is the slope of the obtained straight line) and of the real parabolic constant K_p (ordinate at the origin of this same straight line) corrected from the minimization of mass gain induced by the volatilization of chromia. The volatilization-corrected K_p is again slightly lower for NCCHf than for NCC (12 against $14 \cdot 10^{-12} \text{ g}^2 \cdot \text{cm}^{-4} \cdot \text{s}^{-1}$). In contrast the volatilization rate seems a little higher for NCCHf (87 against $82 \cdot 10^{-10} \text{ g}^2 \cdot \text{cm}^{-4} \cdot \text{s}^{-1}$).



Fig. 3. The comparison of the mass gain curves of NCCHf (curve *1*) and the reference NCC (curve *2*) (*a*); test of the determined kinetic constants by comparison between the experimental mass gain curve (curve *1*) and the mathematical curves plotted from the K_p values either classically determined (curve *2*) or determined according to the K_p - K_v method (curve *3*) (*b*) [5].

Constants	$\underset{\times 10^{-8} \text{ g} \cdot \text{cm}^{-2} \cdot \text{s}^{-1}}{K_{\text{I}}}$	$\times 10^{-12} {\rm g}^2 \cdot {\rm cm}^{-4} \cdot {\rm s}^{-1}$		$\underset{\times 10^{-10}}{\overset{K_{\nu}}{\rm g} \cdot \rm cm^{-2} \cdot \rm s^{-1}}$
Alloys	Slope at stage start	Classic	$m \cdot (dm/dt) = f(-m)$	$m \cdot (dm/dt) = f(-m)$
NCCHf	14.1	3.87	12.0	86.6
NCC (for comparison)	17.8	5.80	14.3	71.7

Values of the oxidation kinetic constants

Before cross-section preparation the two oxidized samples were photographed and subjected to X-ray diffraction. The two types of results are given together in Fig. 4. NCC obviously suffered scale spallation during the -5° C min⁻¹ cooling after the isothermal oxidation, while NCCHf obviously kept all its external scale. The X-ray diffractograms show that even if Cr₂O₃ is the main oxide present in both cases, the scales formed over NCC contain significant parts of spinel NiCr₂O₄ and even NiO. In addition the substrate (Ni–Cr austenitic alloy) is also more detected in its case than for NCCHf consequently to its more extended oxide spallation. The external oxide scale formed around NCCHf is richer in chromia than NCC, but it contains a little spinel NiCr₂O₄ and also the hafnium oxide.

The cross-sectional examinations (Fig. 5, NCCHf surface and subsurface with two magnifications) confirm these observations. Additionally one can see that no carbide-free zone developed from the oxide-alloy interface, in contrast with what generally occurs for cast Ni alloys with chromium carbides or TaC carbides. Indeed, the HfC carbides seem being still present in alloy close to the external oxide. X-ray diffraction and cross-sectional EDS spot analysis show that the carbides close to the oxide/alloy interface are oxidized *in situ*: HfC are replaced by HfO₂ at the same location.



Fig. 4. NCC (*a*) and NCCHf (*b*) oxidized at 1100°C for 46 h: the X-ray diffractograms obtained on the oxidized surfaces ("c": Cr₂O₃; "cn": NiCr₂O₄; "n": NiO; "N": FCC Ni-based matrix); visualization of the oxide spallation (photographs in the top-left corners).



Fig. 5. Cross-sectional observations of the oxidized surface and subsurface of the NCCHf sample (a – general view; b – detailed view).

CONCLUSION

The studied alloy demonstrated very interesting properties: rather high level of hardness but still compatible with a sufficient machinability, good creep resistance at 1100°C for a resulting tensile stress of 20 MPa and very good oxidation behavior. Concerning the two first points the high hardness of the HfC carbides (harder than Cr_7C_3 and $Cr_{23}C_6$) and their stability at HT bring important advantages by comparison with other more classical carbides. Concerning the last point this is essentially the Nibase matrix which brings its intrinsic good high temperature behavior, compared to the Co-based alloys for example). But the second beneficial role of Hf here, which is usually the main one in many cases, was also observed: the improvement in oxide scale

adherence and thus resistance against spallation. Workable by foundry and offering so high levels of mechanical and chemical properties at HT, this alloy ought to meet needs in structural metallic alloys for many high temperature applications.

РЕЗЮМЕ. Литий надміцний сплав на основі Ni, зміцнений карбідами гафнію, виливали та витримували при 1100°С в умовах повзучості та гарячого оксидування. Мікроструктура виливок містить дендритну матрицю, а між дендритами виявлено евтектичну суміш карбідів гафнію, що поліпшує механічні властивості за високих температур. Це підтверджено дослідженнями на триточковий згин в умовах повзучості (за максимальних напружень 20 MPa) та температури 1100°С. Повільна параболічна кінетика втрати маси свідчить про тривкість сплаву до окиснення при 1100°С у повітрі. Карбіди HfC під час цих випробувань залишалися стабільними. Сплав придатний для використання при температурі, яку можуть витримати тільки монокристали $\gamma \gamma \gamma'$ на основі Ni чи суперсплави типу ODS.

РЕЗЮМЕ. Литий сверхпрочный сплав на основе Ni, упрочнен карбидами гафния, выливали и выдерживали при 1100°C в условиях ползучести и горячего оксидирования. Микроструктура отливок содержит дендритную матрицу, а меж дендритами обнаружено эвтектическую смесь карбидов гафния, что улучшает механические свойства при высоких температурах. Это подтверждено исследованиями на трехточечный изгиб в условиях ползучести (при максимальных напряжений 20 MPa) и температуры 1100°C. Медленная параболическая кинетика потери массы свидетельствует об устойчивости сплава к окислению при 1100°C в воздухе. Карбиды HfC оставались стабильными во время этих испытаний. Сплав пригоден для использования при температуре, которую могут выдержать только монокристаллы γ/γ' на основе Ni или суперсплавы типа ODS.

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