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State of the Sub-surface Microstructure of Carbides strengthened cast Superalloys after High Temperature Oxidation – Use of Thermodynamic Modelling for a better Understanding

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Abstract. Several cast strengthened superalloys, Ni base and Co base, were exposed to high temperature oxidation for long times and metallographically examined. Different phenomena occurred in the sub-surface microstructure, depending on both alloy and temperature. Thermodynamic modelling was used to know what it happened for carbon during oxidation, then to explain the observed microstructural changes. It appears that carbon atoms either quit the alloy probably after its oxidation into gases, or on the contrary go deeper into the bulk where they promote the precipitation of new carbides by solid state transformation. Thereafter, thermodynamic modelling allowed to know the new local refractoriness of the zones affected by oxidation, then to appreciate the new mechanical properties in the sub-surface.

Introduction

When exposed to high temperature oxidation most of superalloys become covered by alumina or chromia oxide scales. They are often affected by internal oxidation too, in the matrix or in grain boundaries [1]. But other phenomena can also occur, such as microstructural modifications in the neighbourhood of the oxidation front. For carbides strengthened superalloys after high temperature oxidation, it is often seen that there is an external zone, all along the alloy surface, in which microstructure has been damaged (Fig. 1). For example, this sub-surface microstructure can encounter a disappearance of secondary carbides, a lost of the intergranular carbides with formation of holes or not, precipitation of new phases, … These modifications of the sub-surface microstructural state are not always easy to understand, notably since some elements, like carbon, cannot be analysed because of their too light atomic weight and too small weight content in the bulk. Thermodynamic modelling can be in such cases of a good help. First, when applied in an inverted sense with regard to how it is usually used, one can think that it allows to know the local content of carbon, and then what happened for this element during oxidation. Second, knowing the whole chemical composition, it leads to appreciate the consequences, on the local refractoriness, of the microstructural and chemical changes due to oxidation.

Experimental

Synthesis of the studied alloys. The alloys studied here are of two types. In order to begin this study with a simple example, a ternary Ni-Cr-C nickel alloy was first synthesised.
Fig. 1: Example of deterioration of the sub-surface microstructure by oxidation

Working on such an alloy simplifies the problem by allowing first to analyse in the best conditions the local chemical compositions (except carbon) without any interaction between two elements, and second to perform thermodynamic calculations using a well known database. It was prepared in small quantity (ingots of nearly 50g) by high frequency induction melting under argon gas in a copper cold crucible in which it also solidified. Its microstructure is constituted of a nickel austenitic matrix (including a part of the Cr and C atoms in solid solution) and of chromium carbides at grain boundaries.

Second, two cobalt base superalloys were obtained using another induction furnace (capacity of several kg) under argon gas too. The molten alloys were poured and solidified in a sand mould. Thereafter these two cobalt base superalloys underwent a homogenisation treatment at 1200°C for 2 hours, followed by a precipitation stage at 1000°C for 10 hours, in order to get a precipitation of thin carbides in matrix. These Co base superalloys contain mainly cobalt, nickel and chromium but also tantalum and carbon, the latter elements promoting the formation of refractory Cr$_{23}$C$_6$ and/or TaC carbides which are known to be of a great importance for mechanical properties at high temperature [2,3].

![Co base superalloy oxidised at 1100°C for 100h](image)

<table>
<thead>
<tr>
<th>Alloys</th>
<th>Co</th>
<th>Ni</th>
<th>Cr</th>
<th>C</th>
<th>Ta</th>
</tr>
</thead>
<tbody>
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<td>/</td>
<td>bal</td>
<td>30</td>
<td>0.2</td>
<td>/</td>
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<tr>
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<td>30</td>
<td>0.4</td>
<td>6</td>
</tr>
<tr>
<td>Alloy3</td>
<td>bal</td>
<td>8.5</td>
<td>30</td>
<td>1.0</td>
<td>6</td>
</tr>
</tbody>
</table>

Table 1: Typical chemical compositions of the studied alloys (in weight %)

High temperature oxidation runs and microstructure examinations. Oxidation tests were performed at 1000°C, 1100°C and 1200°C for 100 hours, using a Setaram thermobalance under a flow of industrial air (80%N$_2$-20%O$_2$) of about 1.5L/min. Samples were parallelepipedes, the dimensions of which were approximately 10mm x 10mm x 2mm. After tests, oxidised samples were covered with a gold layer by cathodic plasma pulverisation which allowed the external surface...
to become electrically conductive. Then a sufficiently thick electrolytic nickel coating was deposited in order to protect the external chromia layer before cutting. Thereafter samples were cut, embedded in epoxy resin and polished up to 1200 grid paper. A Scanning Electron Microscope (SEM) Philips XL30 was used in Back Scattered Electrons (BSE) mode for the surface and bulk examinations. The constitutive phases can be clearly evidenced since their average atomic numbers are sensibly different. In addition, some of them were identified by Energy Dispersion Spectrometry (EDS), or by Wavelength Dispersion Spectrometry (WDS) using a Cameca SX100 microprobe. EDS enabled also to analyse the mean chemical composition of a selected area over the local microstructure while microprobe was also used to get concentration profiles through the external zone damaged by oxidation.

**Image analysis and thermodynamic calculations.** Surface fractions of the different phases were estimated on SEM photographs (BSE mode), using the AnalySIS software of Soft Imaging System GMBH. Making the hypothesis that surface fraction and volume fraction have similar values, volume fractions were assessed. Using the equation (Eq. 1) with room temperature densities of the different phases, it was possible to estimate the mass fraction of each phase. The values used for the specific weights (in g cm$^{-3}$) were 8.90 for the matrix, 6.94 for the Cr$_7$C$_3$ carbide, 6.95 for the Cr$_{23}$C$_6$ carbide and 14.50 for the TaC carbide [4].

$$f_{\text{mass}}(\varphi_j) = \sum_i (\varphi_i \cdot \rho_i \cdot f_{\text{vol}}(\varphi_i))$$

Calculations on the simple nickel alloy were carried out using Windows version Thermo-Cal software with the SSOL database (SGTE). For the cobalt alloys, N Thermo-Calc software was used in association with a new database. This last one is the SSOL database completed with binary and ternary systems involving the Ta element. It had been already used to study analogous alloys in other studies [5,6,7].

**Results obtained on the simple nickel alloy (Alloy1)**

**Surface microstructure after oxidation at 1000°C and 1100°C for 100 hours.** At 1000°C, the initial sub-surface microstructure of Alloy1 is transformed into two separated zones (Fig. 2). The outer one, thereafter called "carbideless zone", has completely lost its carbides. The inner zone still contains some carbides but with a rounder shape. One also can see a population of new phases with an acicular shape, different sizes and precise orientations. Spot microanalysis showed that both acicular particles and compact particles are Cr$_7$C$_3$ carbides. WDS chromium profiles, were compared to the different microstructural zones. The carbideless zone corresponds to a high Cr gradient and a chromium range of [21%wt , 26%wt] while the zone containing acicular carbides exhibits a lower Cr gradient and higher Cr contents. SEM photographs were taken on several areas (of about 80µm x 120µm) in the acicular precipitation zone, on which image analysis were performed. EDS analysis of the chemical composition was realised on the same areas. The six results lead to an average of 4.6%mass of Cr$_7$C$_3$ carbides (mass fraction) in the microstructure and to a Cr content of 29.8%wt Cr.

At 1100°C, the damaged zone of Alloy1 seems to be only composed of the carbideless zone, which is three times deeper than the one resulting from oxidation at 1000°C. From the bulk, the WDS Cr content profile shows that chromium content decreases from 31%wt Cr to about 25%wt Cr.

**Thermodynamic calculations.** An isothermal ternary Ni-Cr-C diagram was computed for T=1000°C (Fig. 3). Considering the nickel-rich corner and the Cr contents measured with WDS
Fig. 2: Sub-surface microstructural changes for Alloy1 and correspondence between the different zones and the Cr profile

profiles over this zone, a range of possible values for %wt C was assessed. For each limit of chromium (21 and 26%wt Cr), Thermo-Calc Windows gave the two upper limits for the carbon content allowing the existence of the matrix alone. Thus, the carbideless zone must contain less than 0.037%wt C close to the surface and less than 0.019%wt C on the frontier <carbideless zone - acicular carbides zone>. Secondly, for each microphotograph obtained on the zone with acicular carbides, Thermo-Calc Windows calculations were performed, taking into account the chromium content measured by EDS. Successive values of the carbon content were tested until the mass fraction of the Cr7C3 carbides reached the experimental value given by image analysis (without appearance of any new phases different than matrix and Cr7C3 of course). On the six carbon contents which were obtained on the six studied areas (all contained between 0.40 and 0.55%wt C), the average value was 0.435%wt C.

Then, the initial microstructure near the surface, which contained the same chromium and carbon contents (30%wt Cr, 0.20%wt C) and the same phases (matrix, primary Cr carbides) as the bulk before oxidation test, has evolved because of the oxidation phenomena. It has been replaced by two distinct zones: a carbideless zone strongly impoverished in carbon (21 to 26%wt Cr and less than 0.04 to 0.02%wt C) and another zone containing new acicular carbides and a higher carbon content (over the double of the initial content). These two zones have almost the same depth of about 50µm.

Fig. 3: Isothermal ternary Ni-Cr-C diagrams for T=1000°C and for T=1100°C (in the Ni-rich corner)
On the isothermal diagram obtained for 1100°C, the absence of any other phases than matrix indicates that the carbon content must be lower than 0.054%wt C at the extreme surface (where %wtCr=25) and lower than 0.025%wt C at the junction of the carbideless zone and the bulk (where %wtCr=31). In this way, this zone of the alloy has lost a great part of its initial carbon content. Everywhere else in the alloy, the carbide surface fraction seems to be the same as in the bulk, then there is no region that became enriched with a part of carbon lost by the damaged zone.

Results obtained on the two cobalt base superalloys (Alloy2 and Alloy3)

Surface microstructure after oxidation at 1000°C and 1200°C for 100 hours. Initial bulk microstructure is composed first of an austenitic cobalt matrix with other elements in solid solution. In the interdendritic spaces of Alloy2 there are tantalum carbides belonging to an eutectic with the matrix, whereas Alloy3 contains both eutectic TaC carbides and eutectic Cr$_7$C$_3$ in grain boundaries. These carbides strengthen these superalloys which are very resistant mechanically at high temperature. Apart the external (Cr$_2$O$_3$) and internal (Ta$_2$O$_5$ or CrTaO$_4$) oxides, new acicular oriented phases appeared in the alloys not far from the oxidation front. They are visible in a zone separating a thin carbideless zone and the not damaged zone (Fig. 4).

EDS and WDS spot analysis identified these acicular phases as being Cr$_7$C$_3$ carbides. The two alloys also present a carbideless zone between the extreme surface and the acicular precipitation
zone, which is of about 15µm for Alloy2 and of 10µm for Alloy3. EDS profiles were performed through the carbideless zone. They showed for Alloy2 that the chromium and tantalum contents decrease respectively from 20%wt Cr to 16%wt Cr and from 1.0%wt Ta to 0.0%wt Ta between the frontier of the zone which still contains carbides and the extreme surface. For Alloy3, these contents decrease from 21%wt Cr to 18%wt Cr and from 1.6%wt Ta to 1%wt Ta.

Several high magnification SEM photographs were taken on delimited areas of about 500µm² in the acicular precipitation zone (Fig. 4 right) to be thereafter subjects to image analysis measurements. On the same areas global chemical compositions were analysed by EDS. For each alloy three results were obtained and they showed similar values. On these areas Alloy2 contains 3.3%surf Cr₃C₃ and 1.0%surf TaC (surface fractions), then 2.6%mass of Cr₃C₃ and 1.6%mass of TaC (mass fractions), for a local chemical composition of 27.7Cr-9.8Ni-2.1Ta (bal. Co) in weight percent. Alloy3 contains 10.1%surf Cr₃C₃ and 0.7%surf TaC, then 8.0%mass of Cr₃C₃ and 1.2%mass of TaC, for a local chemical composition measured of 28.9Cr-9.4Ni-1.6Ta (bal Co).

Alloy3 was also exposed to oxidation at 1200°C for 100 hours. Apart an external chromia layer three times thicker and an internal oxidation of Ta more developed than for 1000°C, it exhibits a deeper carbideless zone (Fig. 5). Like for Alloy1 at 1100°C, no precipitation of new phases is visible. Here too EDS spot analysis were performed across the carbideless zone. From the frontier of the not damaged zone towards the extreme surface, Cr and Ta contents decrease from 22%wt Cr and 1.5%wt Ta to 17%wt Cr and 0.3%wt Ta, the nickel content staying at about 10%wt Ni.

Fig. 5: Surface and sub-surface microstructural state of Alloy2 oxidised at 1200°C for 100 hours

**Thermodynamic calculations.** For Alloy2 oxidised at 1000°C and for the EDS results obtained, Thermo-Calc N indicates that the carbon weight content must be lower than 0.076%wt C at the extreme surface and lower than 0.050%wt C at the other side of the carbideless zone. These values are about ten times smaller than the initial C content of Alloy2. For Alloy3 at the same locations, calculations show that carbon content is smaller than 0.061%wt C and 0.019%wt C respectively, as is to say between ten and twenty times smaller than the initial C content of this alloy. Calculations were also performed on the compositions of the zones containing acicular carbides. For Alloy2, calculations converged to the experimental mass fractions of phases resulting from image analysis for 0.27%wt C. The final calculation results gave 2.6%mass of Cr₂₃C₆ and 1.6%mass of TaC. For Alloy3, calculations allowed to come close to the real mass fractions for 0.55%wt C. For Alloy2 oxidised at 1200°C, from the extreme surface towards the beginning of the bulk the maximal possible carbon content decreases from 0.23%wt C down to 0.08%wt C. Like Alloy1 at 1100°C, no zone enriched with carbon can be seen here all over the bulk.

6
Consequences on the thermal properties of the alloys

In the carbideless zone and in the acicular carbides zone, the new local microstructure is due to an evolution of the chemical composition. It can induce changes for the local refractoriness of each zone in the alloy, and there is a melt hazard if working temperature suddenly raises quickly, as it can be encountered for hot industrial pieces. Using ThermoCalc on the local new chemical compositions allows to know what are locally the new solidus temperature. Table 2 displays these calculated values with two cases for the carbideless zones: carbon being equal to its lowest possible value (0\%) or to its highest one ($C_{\text{max}}$). In fact it appears generally that these new microstructural areas are more refractory than the initial alloy, except for the acicular carbide zone (the solidus of which is sensibly equal to the bulk one).

<table>
<thead>
<tr>
<th>Alloys</th>
<th>Locations</th>
<th>external limit</th>
<th>middle</th>
<th>internal limit</th>
<th>acicular $\text{Cr}_7\text{C}_3$ carbide zone</th>
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<tbody>
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<td>/</td>
<td>1380</td>
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<td>1350</td>
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<tr>
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<td>/</td>
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<td>1391</td>
<td>/</td>
<td>1382</td>
<td>/</td>
</tr>
</tbody>
</table>

Table 2: Values of carbon content limits and of resulting local solidus temperatures

Interpretation and discussion

The existence of the carbideless zone is possible only if carbon quantity is smaller than very low values. There, carbon content can be considered as negligible compared to carbon contents of initial alloys and of neighbour zones containing old or new carbides. This zone has almost lost all its carbon atoms whereas the zone in which acicular carbides appeared exhibits a higher carbon content that the initial one. Since bulk presents the initial carbon content, the additional carbon necessary comes from the zone which has lost its carbides. It is possible to simplify the carbon content profile through the damaged zone (Fig. 6). Taking the example of the Alloy1 oxidised at 1000°C, it can be considered that the carbon content is zero across the carbideless zone. It increases quickly and stays at 0.435\%$_{\text{wt}}$C across the acicular precipitation zone, and falls quickly to the bulk content 0.20\% and thereafter stays at this value.
The mass of carbon in the damaged zone is equal to \((S\rho)(e_1%\text{wtC}_1 + e_2%\text{wtC}_2)\). \(S\) is the considered surface, \(\rho\) is the alloy density (about 8.9g.cm\(^{-3}\)). \(e_i\) and %\text{wtC}_i\) are the depths and carbon contents of the two parts of the damaged zone. For 1cm\(^2\) of external surface, we find about 0.30mg of C in the damaged zone, which is sensibly the initial carbon mass contained in the same volume (0.27mg). Then, almost all the carbon lost by the carbideless zone came into the matrix more deeper in the alloy. This carbon left the grain boundary chromium carbides, the decomposition of which was promoted by the diffusion of Cr towards the oxidation front. Thereafter, they diffuse backwards and entered into the matrix and met chromium atoms present in solid solution. There carbon formed with chromium new carbides which precipitated in matrix with an acicular shape. In a same grain, these new carbides exhibit a same orientation. They may coexist with the initial grain boundary carbides which have not yet disappeared and which have a more compact shape.

At 1100°C, the mechanisms would not be the same, since no carbon enrichment can be noted elsewhere in the alloy. Carbon lost by decomposition of the initial grain boundary carbides disappears, without any doubt being oxidised into CO or CO\(_2\) gases. For 1cm\(^2\) of Alloy1 external surface and a carbideless zone of about 100µm in which carbon content is negligible, the same type of calculation as before leads to a carbon mass lost of 180mg that became gas.

Unfortunately, such calculations are not really possible for the two cobalt alloys. Indeed, zones with acicular precipitated carbides often are partially mixed in their inner part with remaining grain boundary carbides and it is difficult to be precise. But new acicular Cr\(_7\)C\(_3\) carbides also appear among the initial TaC secondary carbides for Alloy2, and among the initial secondary TaC and Cr\(_7\)C\(_3\) for Alloy3. For the two alloys it is a consequence of an increase of the total carbon contained in the matrix alone. This clearly shows that the mechanism of integration of carbon into matrix also acts here. Like Alloy1 at 1100°C, it was seen for Alloy2 that a raising of the temperature unable this phenomenon which is probably also replaced by a disappearance of carbon from the alloy (becoming oxidised into gases). But this difference of behaviour remains to explain.

Finally thermodynamic modelling allowed us to verify that these new compositions induced by oxidation are not prejudicial for the alloys sub-surface refractoriness.

**Conclusion**

It can be underlined that thermodynamic modelling, associated to microstructure phase quantitative analysis and microanalysis, was of a good help to better characterise and understand what it happened to our specific alloys and superalloys. This method allowed to supply the lack of microanalysis techniques for what quantitative measurements of contents for light and rare elements is a real problem. With this particular procedure it became possible to localise carbon, which is an element of a great importance in our alloys. Consequently curious phenomena have been identified.

**References**