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# Full Length Article

# Influence of solidification induced composition gradients on carbide precipitation in FeNiCr heat resistant steels



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# ABSTRACT

Secondary precipitation of Cr-rich carbides in heat resistant austenitic stainless steels has been investigated both experimentally and using finite element simulations. The microstructural evolutions in two commercial grades were characterized using electron microscopy. A special emphasis was given on the peculiar spatial distribution of  $M_{23}C_6$  secondary carbides exhibiting precipitate free zones surrounding primary carbides, and high density precipitate zones extending with longer aging times. Solidification induced chemical composition gradients were clearly exhibited in the as-cast alloys with significant chromium depletion in the vicinity of primary  $M_7C_3$  carbides. It is then proposed that these gradients play a major role in the secondary precipitation mechanism. Classical nucleation and growth theories have been adapted to account for (i) the flux of solutes with large difference in diffusion coefficients, (ii) the initial composition gradients in the matrix and, (iii) the chemical driving force for nucleation and growth of  $M_{23}C_6$  carbides. Within this framework, the whole kinetics has been reproduced. It clearly shows that the spatial distribution of secondary carbides that play a key role in the creep resistance of these alloys is the result of a complex interaction between initial composition gradients in as-cast alloys and solute flux resulting from phase transformation during aging.

# 1. Introduction

Heat resistant steels used in cracking or reforming furnaces have to face harsh service conditions including high temperatures (from 750 °C up to 1100 °C), oxidizing environment on the external surface of pipes and in some cases carburizing on the inside [1-4]. Steels with high chromium contents (> 25 wt%) have been designed to resist against oxidation but, since a significant creep resistance is also required, a significant amount of nickel has also been introduced to avoid the ferritic structure and to obtain fully austenitic steels [3]. Pipes of such FeNiCr steels (known as HK or HP grades) are typically produced using centrifugal casting. During solidification, large columnar grains with a dendritic structure are typically created. Dendrites from the fcc austenitic phase are surrounded by micrometer scale eutectic carbides. These carbides, called primary carbides, are MC (Nb or Ti rich) or M7C3 (Cr rich) carbides [5]. It is important to note that these materials are often directly used in as-cast conditions but M7C3 primary carbides are not stable in the temperature range of applications and transform into the stable M<sub>23</sub>C<sub>6</sub> [6]. This transformation is accompanied by the precipitation of secondary carbides which typically nucleate and grow within the austenitic matrix [7]. It has been demonstrated that secondary carbides play a very important role since they significantly improve the creep resistance by hindering dislocations [8–12]. Therefore controlling the number density, the volume fraction and the size distribution of these carbides is a key issue to improve the creep resistance.

The transformation of  $M_7C_3$  in  $M_{23}C_6$  and the subsequent secondary carbide precipitation within the austenitic matrix have been studied in many systems like high Cr steels [13–16], Ni superalloys [17,18] or austenitic stainless steels [5,7,19–21]. It is important to note that when the primary  $M_7C_3$  carbides transform into  $M_{23}C_6$ , their carbon content is significantly reduced. Even though after aging the original macrostructure looks relatively unchanged under the optical microscope where the micrometer scale primary  $M_7C_3$  are transformed into  $M_{23}C_6$ , the carbon excess is released in the austenitic matrix giving rise to a homogeneous nucleation and growth of submicrometer  $M_{23}C_6$ . These carbides exhibit a crystallographic structure with a super lattice very close to the fcc structure of the austenitic matrix (with an average mismatch of only 1.3% parallel to the {111} planes) and therefore grow with a cube on

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#### Table 1

Nominal composition (wt%) of the two steels investigated in the present work (Fe and other minor elements balance). Other minor elements are: Mn, Si, P, S, Ti, Zr, Pb, Sn, Cu and Mo, with concentrations very close in both alloys.

	Steel 1	Steel 2
С	0.45-0.5	0.4–0.5
Ni	33.0-36.0	39.0- 42.0
Cr	24.0-27.0	29.0-32.0
Та	-	0.0-2.0
Al	-	0.0-2.2
Nb	0.5-1.0	0.0-0.2

cube orientation. Due to the elastic distortions resulting from the misfit, it has been shown that the nucleation barrier could be reduced in the vicinity of already existing carbides, leading to some typical rows of carbides [7]. It has also been often reported that  $M_{23}C_6$  carbides easily nucleate along dislocations where the elastic energy associated with the lattice distortion is reduced [19].

There are however few interesting and intriguing features about the peculiar secondary carbide distribution and growth mechanisms that has not been fully clarified yet. Indeed, despite the great variety of heat resistant steels, aged microstructures often exhibit: (i) a precipitate free zone (PFZ) surrounding primary carbides, (ii) a high density precipitate zone (HPZ) few micrometers away from primary carbides, (iii) a low density precipitate zone (LPZ) in the austenitic grain interior [22–26]. The extension of these different zones depends on the alloy composition, casting conditions and ageing conditions but they are always observed. Thus, the aim of the present work is to carefully examine the structure of aged heat resistant steels to provide some input in a model that takes into account chemical gradients and diffusion fluxes (especially the flux of carbon atoms released by the transformation of primary carbides) to account for the complex microstructural evolution resulting from secondary precipitation in heat resistant steels.

In this paper, we investigate the precipitation of  $M_{23}C_6$  secondary carbides in the vicinity of existing primary carbides by Scanning Electron Microscopy (SEM) and Transmission Electron Microscopy (TEM). A simple precipitation model based on classical nucleation and growth theories is then proposed to analyze the precipitation kinetics of secondary carbides and to explore the role of solidification induced chemical gradients revealed around primary carbides.

# 2. Experimental details

Two different austenitic heat resistant steels produced by Manoir Industries were investigated in the present study. Centrifugally cast tubes made of these materials are typically used in cracking furnaces [2]. The nominal composition of these two commercial steels is given in Table 1. They have been selected with similar carbon content but a significant difference in Ni/Cr ratio. It is also interesting to note that steel 1 contains a small amount of Nb while steel 2 contains some Al and Ta. Nb and Ta are known to form MC carbides (M being Nb or Ta) and thus slightly affect the volume fraction of M7C3 and M23C6 carbides, however the aim of the present study was to show that the detailed mechanisms of the secondary precipitation during aging could be captured for most commercial grades and that it does not significantly depend on small variations of the volume fraction of carbides. Artificial aging experiments were carried out in furnaces with a natural air environment at 750 °C. This temperature is lower than most of industrial applications, but it has been selected to slow down the kinetics and capture the different stages of the secondary precipitation mechanism (observations done after 16, 64 and 112 h of aging have been selected for the present manuscript). Equilibrium phases and their temperature range of stability have been computed with Thermocalc software using the TCFe-6 data



**Fig. 1.** Thermocalc prediction of equilibrium molar fractions of phases in steel 1 (a) and steel 2 (b) as a function of temperature.  $M_7C_3$  corresponds to the Crrich primary carbides.  $M_{23}C_6$  corresponds to the Cr-rich secondary carbides. NbC corresponds to the Nb-rich primary carbides.  $\sigma$  corresponds to an Fe- and Cr-rich intermetallic phase. G corresponds to the G-phase, an intermetallic silicide.

base (Fig. 1). In both alloys, the  $M_7C_3$  phase is stable only just below the liquidus temperature, but at 750 °C the only stable Cr-rich carbide is  $M_{23}C_6$ . It is important to note that below 800 °C a significant equilibrium volume fraction of G phase and  $\sigma$  phase is exhibited for steel 1 and steel 2, respectively. However, these phases nucleate only after very long aging time [20] and thus does not significantly affect the secondary precipitation mechanisms.

To follow the transformation of primary M7C3 carbides and the resulting precipitation of M23C6, microstructures were characterized by SEM and TEM. SEM observations were carried out with a ZEISS LEO 1530 XB operating at 15 kV. Images were recorded using a secondary electron detector and a 60 µm aperture. Prior to SEM investigation samples were mechanically polished with  $\frac{1}{4}$  µm diamond paste. In order to image primary and secondary carbides, the samples were chemically etched using either an oxalic acid solution (preferential etching of the austenitic matrix) or an A2 solution from Struers (preferential etching of Cr-rich carbides). TEM observations were carried out with a probe corrected JEOL ARM-200F operated at 200 kV. Analytical data were collected by scanning TEM (STEM) with a probe size of 0.2 nm and a convergence angle of 34 mrad. Elemental mapping was carried out using Energy Dispersive X-ray Spectroscopy (EDS) with an Oxford Instruments X-max detector, which has a solid angle of 0.7 sr. TEM samples were prepared from 3 mm disks thinned down to electron transparency with a twin jet electropolisher (Tenupol 5 from Struers) using a solution of 10% perchloric acid - 90% acetic acid at +14 °C and a voltage of 20 V. To obtain better quality samples, a final polishing was carried out



**Fig. 2.** SEM images (secondary electron detector) showing the microstructure of steel 1 in the as-cast state (a) and after aging at 750 °C for 16 h (b), 64 h ((c) and (e)) and 112 h (d). Precipitation of  $M_{23}C_6$  secondary carbides in the vicinity of primary carbides is clearly exhibited for the early stage of aging, and it progressively extends in the matrix. The higher magnification image recorded in the austenitic matrix (e) reveals smaller secondary carbides and show that some of them nucleated along a linear defect (supposedly a dislocation). For images (a) and b) the samples were etched using A2 solution from Struers to brightly image the matrix and for images (c)–(e) the samples were etched using oxalic acid solution to brightly image the carbides). In Fig. 2(a) the inset is a SAED pattern collected on a primary carbide and that demonstrates that these carbides exhibit a  $Cr_7C_3$  structure.

by ion milling (Gatan PIPS II operated at 3 keV during 5 min with  $\pm 3^{\circ}$  inclination angle).

# 3. Experimental results

Microstructures of steel 1 and steel 2 after aging at high temperature are typical for such heat resistant steels with large primary eutectic carbides (several micrometers in size) located between austenitic dendrite arms (width ranging from 30 to 60  $\mu$ m). The evolution of the microstructure near primary carbides is shown in Fig. 2 for aging times up to 112 h at 750 °C. In the as-cast state, large and lamellar eutectic M<sub>7</sub>C<sub>3</sub> carbides are exhibited (darkly imaged on Fig. 2(a) and crystallographic structure confirmed using selected area electron diffraction (SAED) in the TEM) while the matrix is free of any precipitate. Then, after only 16 h at 750 °C (Fig. 2(b)), numerous small carbides nucleate in the matrix within a distance of only a few microns from primary carbides. The structure and the orientation relationship of these carbides has been checked using TEM (Fig. 3), they are M<sub>23</sub>C<sub>6</sub> carbides as expected. Further annealing, up to 64 h leads to an apparent growth of secondary carbides (brightly imaged in Fig. 2(c)) that have nucleated earlier while new carbides seem to nucleate in the matrix (top left of Fig. 2(c)). A closer look at higher magnification in such a region clearly reveals that some of the secondary  $M_{23}C_6$  carbides have nucleated on linear defects, supposedly dislocations (Fig. 2(e)) as reported by other authors for similar steels [19]. Then, further annealing (Fig. 2(d)) leads to a significant extension of the secondary precipitation zone and also to further growth of  $M_{23}C_6$  carbides. It is interesting to note that, as already reported by other authors [3], secondary carbides do not nucleate on primary carbides (which are carbon reservoirs) but in the matrix at a short distance of a few micrometers, leaving a PFZ.

A very similar microstructure evolution was observed for steel 2 (Fig. 4) with also a progressive growth of the secondary precipitation zone and a PFZ near primary carbides. The microstructural differences of the two investigated steels as observed by SEM are quantitatively plotted as a function of the annealing time in Fig. 5. The mean thickness of the PFZ is less than a micrometer in both alloys, while the size ( $\lambda$ ) of



Fig. 3. TEM images (Bright field, High resolution and SAED pattern in (001) zone axis) showing secondary  $M_{23}C_6$  carbides that have nucleated and grown during aging at 750 °C for 112 h in the austenitic matrix of steel 1 with a cube on cube orientation relationship.



**Fig. 4.** SEM images (secondary electron detector) showing the microstructure of steel 2 (etched using oxalic acid solution to brightly image carbides) after aging at 750 °C during 16 h (a), 64 h (b) and 112 h (c). Precipitation of  $M_{23}C_6$  secondary carbides in the vicinity of primary carbides is clearly exhibited for the early stage of aging, and it progressively extends in the matrix.

the observable secondary precipitation extension (or High Precipitation Zone (HPZ)) toward the austenitic matrix is almost five times larger in steel 2. This feature is also connected with an apparent volume fraction of secondary carbides significantly higher in steel 2 comparing to steel 1 after 112 h at 750 °C (Figs. 2(d) and 4(d)). Such a difference in volume fraction was not predicted by the TCFe-6 database with thermocalc (Fig. 1) and is supposedly due to a faster growth and coalescence in

steel 2. Indeed, as shown in Fig. 2(e) there is also a large density of secondary  $M_{23}C_6$  carbides away from the HPZ in steel 1 but not exceeding 100 nm in size and thus hardly observable and quantifiable from low magnification SEM (in the following it will be called the Low Precipitation Zone (LPZ)). This fastest growth and coalescence in steel 2 is also leading the formation of carbides with an elongated shape (Fig. 4(c)). It has indeed been reported by other authors that  $M_{23}C_6$  carbides may



**Fig. 5.** Mean thickness of the precipitate free zone (PFZ) around primary carbides and mean distance ( $\lambda$ ) of the observable secondary precipitation extension toward the austenitic matrix as a function of aging time at 750 °C for steel 1 and steel 2 (measured from SEM images).

nucleate aligned along some specific crystallographic directions due to the lattice distortions resulting from the small misfit between their crystal structure and the fcc matrix [7]. Then, in SEM images, rows of precipitates appear as needles when they coarsen or eventually coalesce to form very elongated shape carbides.

The transformation of primary M7C3 carbides into M23C6 with a lower molar fraction of carbon triggers the secondary precipitation by releasing some carbon into the austenitic matrix. Then, if the solubility product  $K_s = [M]^{23}[C]^6$  is reached, secondary carbides may nucleate if their radius is larger than the critical radius [3]. SEM observations clearly show that they do not nucleate at primary carbide / austenitic matrix interfaces and that a PFZ appears instead. This means that the solubility product is not reached in the PFZ. Since this solubility product is a function of the local composition, this might indicate that there is a composition gradient across primary carbide/matrix interfaces. To measure the local composition of the matrix near primary carbides in the as-cast material, analytical STEM analyses using EDS have been carried out in both steels. As shown in Fig. 6, a significant Cr gradient appears at such interfaces with a significant Cr depletion in the matrix. This depletion is about 5 and 10 at% in steel 1 and steel 2, respectively, (from 30 at% down to 25 at% Cr in steel 1 (Fig. 6(a)) and from 32 at% down to 20 at%Cr in steel 2 (Fig. 6(b))). These Cr gradients in the matrix spread over about one and two micrometers in steel 1 and steel 2, respectively.

Then, if such experimental data give some qualitative explanation of the observed PFZ, it does explain why some secondary  $M_{23}C_6$  carbides continuously grow further and further away from primary carbides. This is obviously linked to matrix composition changes during the carbide growth which affects the solubility product. Besides, the diffusivity of carbon at 750 °C in the austenitic matrix is relatively high. The equivalent diffusion distance estimated as  $\approx \sqrt{6D_c t}$  where  $D_c$  is the carbon diffusion coefficient given in Table 2 and t the time, is indeed about 1 mm in only 16 h. Then, it is hard to understand why  $M_{23}C_6$  does not homogeneously nucleate in the austenitic matrix since the carbon concentration should be quite homogenous in the austenite. Therefore, to account for these complex phenomena (initial composition gradient, local solubility product evolution and carbon flux), the coupled diffusion/precipitation model described below has been developed.

#### Table 2

Thermodynamic and kinetic parameters	used for modeling of the precipitation
with nodePreciSo.	

Parameter	Value	References
$\begin{array}{c} S^{12},S^{23},S^{34},\ldots\\ S^{01}\\ X^1,x^2,x^3,\ldots\\ V^1,V^2,V^3,\ldots\\ M_{Fe} \end{array}$	1m <sup>2</sup> 1.5×10 <sup>-6</sup> m <sup>2</sup> 1 μm 10 <sup>-6</sup> m <sup>3</sup> 55.845×10 <sup>-3</sup> kg/mol	[30]
$M_{Cr} \\ M_C \\ D_{Cr}(750 °C) \\ D_C(750 °C) \\ \gamma_{matrix} \\ v_{Cr_{23}C_6} \\ \gamma_{Cr_{23}C_6} \\ K_S (750 °C) \\ \end{cases}$	51.996 × 10 <sup>-3</sup> kg/mol 12 × 10 <sup>-3</sup> kg/mol 1.4 × 10 <sup>-17</sup> m <sup>2</sup> /s 9.26 × 10 <sup>-12</sup> m <sup>2</sup> /s 1.664 × 10 <sup>-29</sup> m <sup>3</sup> 5.2 × 10 <sup>-30</sup> m <sup>3</sup> 0.5 J/m <sup>2</sup> 6.77 × 10 <sup>-18</sup>	[30] [30] [31] Thermo-Calc Software TCFE8 Thermo-Calc Software TCFE8 Adjusted Parameter Thermo-Calc Software TCFE8

# 4. Modeling of the secondary precipitation of $M_{23}C_6$ carbides

Classical nucleation and growth theories have been extensively used to model precipitation in isothermal and/or non-isothermal condition for steels [32], aluminium alloys [33] and superalloys [34]. From the pioneering contribution of Böhm et al. [35], H.P. Van Landeghem [37] modeled the precipitation of CrN in an evolving N chemical gradient and Rougier et al. [36] reported precipitation of  $\gamma$ ' precipitates in a Ni-based superalloy. However, modeling precipitation kinetics in a concentration gradient has been seldom reported in the literature (see the recent contribution of [36]).

## 4.1. Fundamental basis of the modeling approach

The purpose is here to model the precipitation of the secondary carbides, supposedly driven by the carbon release from the primary  $M_7C_3$  to  $M_{23}C_6$  transformation, and within an initial chromium gradient resulting from the precipitation of former  $M_7C_3$ . Therefore, a model coupling precipitation and one-dimensional diffusion is designed.

The model developed for the present study is based on the software PreciSo developed by Perez et al., which is described in [27]. PreciSo uses classical nucleation theory, growth and coarsening in order to predict the size distribution of precipitates in multicomponent systems. For the sake of simplicity, complex  $M_{23}C_6$  carbides are assumed to be spherical with  $Cr_{23}C_6$  chemistry. The description for a multi-component system can be found in reference [28]. At each time step of the simulation, when nucleation occurs, a new precipitate class is created. Its radius corresponds to the critical radius  $R^*_{k_BT}$  calculated as follow:

$$R^*_{k_BT} = R^* + \frac{1}{2}\sqrt{\frac{k_BT}{\pi\gamma}}$$
(1)

where  $k_B$  is the Boltzmann constant,  $\gamma$  is the precipitate/matrix interfacial energy.  $R^*$  is the classical critical radius above which nuclei are stable. It is the solution of  $d\Delta G(R^*)/dR = 0$ , where the energy change  $\Delta G(R)$  is given by:

$$\Delta G(R) = \frac{4}{3}\pi R^3 \Delta g + 4\pi R^2 \gamma \tag{2}$$

where  $\Delta g$  is the driving force for precipitation. The number of precipitates in the newly created class is determined by the nucleation rate:

$$\frac{dN}{dt} = N_0 \beta^* Z \exp\left[-\frac{\Delta G(R^*)}{k_B T}\right] \left[1 - \exp\left(-\frac{t}{\tau}\right)\right]$$
(3)

where  $N_0$  is the number of nucleation sites per unit volume,  $\beta^*$  is the condensation rate of solute atoms in a critical cluster and *Z* the Zeldovich factor.



**Fig. 6.** (a) STEM-EDS line profile analysis measured perpendicular to the interface of a primary carbide towards the matrix in steel 1 in the as-cast state. It clearly exhibits a Cr depletion and a Ni enrichment at the carbide interface. (b) STEM-EDS line profile recorded in similar way in steel 2 in the as-cast state. A chemical gradient is also exhibited at the interface between primary carbides and the matrix. One should note that the carbon quantification is not accurate using EDS, especially because of contamination, however the level being relatively constant in the matrix, it does not affect the Cr gradients.

At each time step, the radius of each class is updated. The new radius is calculated using a classical growth equation:

$$\frac{dR}{dt} = \frac{D_C}{R} \frac{X_C^0 - X_C^i(R)}{\alpha X_C^p - X_C^i(R)} = \frac{D_{Cr}}{R} \frac{X_{Cr}^0 - X_{Cr}^i(R)}{\alpha X_{Cr}^p - X_{Cr}^i(R)}$$
(4)

where  $D_j$  are the diffusion coefficients (*j* stands for either Cr or C),  $X_j^0$  are the initial compositions of the matrix (updated according to solute atoms diffusion, see next paragraph),  $X_j^p$  are the compositions of the precipitate and  $\alpha = v_{at}^M / v_{at}^P$  is the matrix to precipitate atomic volume ratio.

Eq. (4) implicitly takes into account the Gibbs-Thomson effect and therefore Ostwald Ripening since the composition of the matrix at the precipitate interface  $X_j^p(R)$  depends on its radius. In a  $\operatorname{Cr}_{23}C_6$  phase, the composition is modified as follows:

$$X_{Cr}^{i}(R)^{23}X_{C}^{i}(R)^{6} = K_{S} \exp\left(\frac{58\gamma v_{at}^{P}}{Rk_{B}T}\right)$$
(5)

where  $K_S$  is the solubility product of  $Cr_{23}C_6$  in austenite.

In its most recent version called nodePreciso [29], the software is able to simulate precipitation in different "nodes" experiencing different thermal history. The precipitation software has been improved here to be able to model the precipitation within a chemical gradient of diffusing solute atoms. At the beginning of the simulation a mesh is defined. Each node *i* of the mesh has a given volume  $V^i$ , a given solute content  $X_j^{0^i}$  composition and is connected to neighboring nodes *k* via surfaces  $S^{ik}$ . Each time step of the simulation starts with a diffusion step during which a node can exchange atoms with its neighbors. Fluxes of atoms through the defined surfaces are given by Fick's law. At each time step, the solute contents  $X_j^{0^i}$  of each node *i* and for each atom species *j* are updated according to:

$$X_j^{0i} \leftarrow X_j^{0i} + \frac{D_j}{V^i} \Delta t \sum_{k(neigh of i)} \frac{X_j^k - X_j^i}{|x^k - x^i|} S^{ik}$$

$$\tag{6}$$

where  $V^i$  is the volume of node *i*,  $\Delta t$  is the time step,  $x^{i,k}$  is the position of node *i*, *k* and  $S^{ik}$  is the surface of *ik* interface. The summation is performed on all nodes *k* neighbors of node *i*.



**Fig. 7.** Result of the modeling for a steel having a Cr content representative of steel 2 and exhibiting chemical gradients across the interface between primary carbides and matrix. The volume fraction of precipitates (a) and the Cr content of the matrix (b) are plotted as a function of time in nodes 1–20.

This diffusion step is followed by performing the mass balance of all solute atoms in each node i according to:

$$X_{j}^{i} = \frac{X_{j}^{0^{l}} - X_{j}^{p^{l}} \alpha f^{i}}{1 - \alpha f^{i}}$$
(7)



**Fig. 8.** Visualization of the secondary carbide distribution at 750 °C after  $10^4$  s (a),  $10^5$  s (b) and  $10^6$  s (c). Both the precipitate free zone and the evolution of the size distribution as a function of time is clearly exhibited. Images were produced on the basis of the modelling results with a random distribution of carbides in each node.

where  $f^i$  is the precipitate volume fraction in node *i* and  $\alpha \sum_i X_j^i$  is the net balance of solute atoms that entered node *i* since the beginning of the simulation.

Finally, in each node, the nucleation and growth stages (Eqs. (3)–(5)) are calculated.

This new implementation will be used in order to simulate the precipitation of  $M_{23}C_6$  secondary carbides in a chromium composition gradient. The chosen thermodynamic and kinetic parameters for such simulations are reported in Table 2.

The simulations focus on understanding how a composition gradient can influence nucleation, growth and coarsening. Therefore, the studied system is simplified with respect to the rather complex alloys studied experimentally and the following model was build:

## i) the $M_{23}C_6$ phase is the only phase precipitating;

- ii) each simulation consists of 21 nodes and the name of a node corresponds to its position along the x axis;
- iii) node 0 represents the primary carbide that releases C from its transformation from  $M_7C_3$  to  $M_{23}C_6$  (the transformation itself is not modeled – precipitation is inhibited in this node);
- iv) node 0 initially contains 18 wt% Cr and 12 wt% C (in order to take into account the fact that the transformation of the primary carbide is driven by Cr diffusion, the exchange surface between node 0 and node 1 has been reduced to  $1.5 \times 10^{-6}$  m<sup>2</sup> which corresponds to the ratio  $D_{\rm Cr} / D_{\rm C}$ );
- v) nodes 1-20 represent the surrounding austenitic matrix;
- vi) each node is 1 µm thick and is connected to its two neighbors by a unit surface of 1 m<sup>2</sup> (the problem is 1D);
- vii) node 20 is 20 μm away from the primary carbide interface and represents the center of a dendrite (the boundary condition applied at node 20 outer interface is that there is no flux);
- viii) the initial Cr and C compositions of all nodes are fixed at the beginning of a simulation (initial C concentrations have been set to zero for all nodes, whereas various initial Cr gradients have been tested).

In short, this model represents a situation where C is released from the transformation of  $M_7C_3$  into  $M_{23}C_6$ , within a pre-existing Cr gradient. In this paper we will present three different cases: (i) no Cr gradient; (ii) sharp Cr gradient; (iii) smooth Cr gradient.

## 4.2. Modeling results

Fig. 7 shows the evolution of the precipitate volume fraction and the Cr content in nodes 1–20 in the case of the smooth composition gradient (it corresponds to gradient 3 in Fig. 9). The initial Cr content can be seen at the origin of the diagram. Shortly after 2000 s, a small amount of  $M_{23}C_6$  precipitates starts to appear in nodes 9–20 quickly followed by nodes 8 and 7, and later on by nodes 6, 5 and 4. At the beginning of the simulation, the volume fraction of precipitates is higher in nodes further away from the primary carbide. After  $5 \times 10^4$  s though, nodes 7 and 8 become the more populated nodes. Nodes 1–3 remain precipitate-free for the whole simulation time.

Even if it seems surprising at first, the evolution of the volume fraction can be understood as follows. First, it is important to consider all parameters of the simulation: not only the volume fraction, but also the size distribution of the precipitates, their number density and the matrix composition. Since it is hard to follow all these parameters at once, typical simulated volumes have been represented in Fig. 8. These volumes have been generated by taking into account the aforementioned parameters. A cross section has been randomly cut through these volumes in order to provide a view of the microstructure that could be compared qualitatively with SEM observations.

Thanks to this representation, the precipitation sequence becomes clear. In the first steps of the simulation, the Cr content is highest in nodes 9-20. Since carbon diffuses much faster than chromium, all nodes see their C content increasing before nucleation even starts. Carbon flows out from the primary carbide (node 0) and meets nodes in which the Cr content is low. In these nodes, the Cr content is not high enough to reach the solubility limit for nucleation. Hence, nodes 1-3 correspond to the experimentally observed PFZ. In the following nodes (4-20), where the product [Cr]<sup>23</sup>[C]<sup>6</sup> is high enough, nucleation starts. Nodes 9-20 contain a lot of small precipitates. Nodes 4-8 contain much less precipitates, but these precipitates are larger. During the following time steps, these larger precipitates coarsen faster at the expense of the other ones due to the Gibbs-Thomson effect. With time, the difference between the large precipitates and the smaller distant ones is amplified. This explains the observation of a HPZ and the presence of small secondary carbides far from the transforming  $M_7C_3$  carbides as can be seen in Fig. 2. Even if the Cr content increases through time in nodes 1-3, the conditions for nucleation in these nodes is never met.



**Fig. 9.** Result of the modeling  $(750 \,^{\circ}\text{C} \text{ for } 10^6 \text{ s})$  showing the influence of the chemical gradient at the primary carbide / austenitic matrix interface. (a) Representation of the three different initial gradients (labelled 1: no gradient, 2: sharp gradient, 3: smooth gradient) used in the modeling, (b) corresponding visualization of the secondary carbide distribution showing a PFZ and a larger size distribution for the largest gradient.

These simulations reproduce features observed during SEM investigation qualitatively. The first nodes are precipitate-free. This PFZ is followed by a zone where the volume fraction of precipitates is the highest and contains large precipitates. The more distant nodes, corresponding to the center of a dendrite, contain a lot of smaller precipitates. In addition, it helps understanding the reasons for such a peculiar distribution of precipitates and how exactly a composition gradient can influence precipitation. The results show how thermodynamic and kinetics considerations have to be taken into account in order to understand the precipitation sequence, and that the interplay between composition gradients, the solubility product and diffusion coefficients is the key controlling the whole phenomena.

In the literature, typical features such as the size distribution of precipitates, the PFZ, HPZ and LPZ vary a lot depending on aging conditions and alloy composition. This variability can be explained using our diffusion/precipitation model. For instance, simulations have been used in order to predict the precipitate distribution in the case of three different composition gradients. These gradients and the associated simulated volumes are represented in Fig. 9. In case 1 (no initial composition gradient), precipitation occurs everywhere. Even if the volume fraction is slightly higher in the nodes closer to the primary carbide / matrix interface (not shown here for the sake of simplicity), there is no striking difference between node 1 and 20. This shows clearly that an initial composition gradient is the necessary condition leading to a PFZ, HPZ and LPZ. Moreover, by comparing cases 2 (sharp gradient) and 3 (smooth gradient) it clearly appears that the spatial extent of the gradient significantly affects the distribution of precipitates. In the case of a sharper gradient, the size of the PFZ is reduced. In addition, a smaller difference in precipitate size is observed between the nodes closer to the primary carbides and these ate the dendrite centre.

## 5. Conclusions

- i) Heat resistant austenitic stainless steels exhibit secondary carbide precipitation as a result of the transformation of  $M_7C_3$  primary carbides into  $M_{23}C_6$  leading to some carbon release into the austenitic matrix.
- ii) Secondary  $M_{23}C_6$  carbides start nucleating at a short distance of primary carbides, leaving a PFZ of one micrometer or more.
- iii) Nanoscaled secondary  $M_{23}C_6$  carbides nucleate in the whole austenitic matrix (excepted in the PFZ), but those near the carbon source (primary carbides) grow quickly and are the only ones visible by SEM for the early stage of aging (HPZ).
- iv) In as-cast alloys, some chemical gradients exist within the structure, and especially a Cr depletion in the vicinity of M<sub>7</sub>C<sub>3</sub> primary carbides has been observed.
- v) The Cr gradients near the carbon source for secondary precipitation affect the local driving force for nucleation and give rise to a PFZ. This has been confirmed using coupled diffusion/precipitation simulations. Theses simulations take into account the flux of solutes with large difference in diffusion coefficients, the initial composition gradients in the matrix and the chemical driving force for nucleation and growth of  $M_{23}C_6$  carbides.
- vi) The simulations also predict that the extent and the amplitude of the Cr depletion zone affect the final spatial distribution of secondary carbides. This might explain the tremendous variations of the PFZ, and HPZ reported in the literature for commercial heat resistant steels.
- vii) The secondary precipitation of  $M_{23}C_6$  carbides can also be affected by crystalline defects like dislocations or the strain field near MC carbides. Such features were beyond the scope of the present work but should be addressed in the future to capture the whole mechanisms.

# **Declarations of interest**

None.

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