Structure, Morphology and Coarsening Behavior of MX (NbC) Nanoprecipitates in Fe-Ni-Cr Based Alloys

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The precipitates of MX-type (where M = Ti, Nb, V, Mo, Zr, … and X = C, N) are considered as the most important strengthening phase in Fe-Ni-Cr based alloys designed for high temperature applications. Despite extensive effort focused on the study of this phase, many questions related to the structure, nucleation, coarsening behavior, as well as the mechanisms of interaction with dislocations, remain unanswered. This work is focused on Sanicro 25, a newly designed alloy with superior high temperature properties (e.g. the highest reported creep strength at 700°C among all commercial heat-resistant austenitic steels) which are strongly determined by the precipitates of MX-type, in particular NbC, nucleating during high temperatures [1]. Various thermo-mechanical experiments combined with state-of-the art electron microscopy and modelling tools were used to study structure, morphology, interface character and coarsening behavior of MX nanoprecipitates.

A set of specimens where the thermal exposure at 700°C was varied, combined without and with symmetrical cyclic tension-compression loading (see Figs. 1a and 1b), were cut and electron backscatter diffraction mapping was performed on polished cross-sections. Site- and orientation-specific thin foils were extracted using the focused ion beam. Regions with NbC precipitates were analyzed along different zone axis orientations using combined atomic-resolution high-angle annular dark-field (HAADF) scanning transmission electron microscopy (STEM) Z-contrast imaging, Super-X energy-dispersive X-ray spectroscopy, electron energy loss spectroscopy and nanobeam diffraction. NbC precipitates have a face-centered cubic crystal structure and are oriented in a cube-on-cube relationship with the austenite matrix (see Figs. 1c and 1d). They have a lattice parameter of 4.47 Å, i.e. a positive lattice misfit of about 24% relative to the matrix. Structural characteristics were further validated by HAADF-STEM image simulations of Moiré-like contrast using the quantum excitation of phonons model (Fig. 1e) [1, 2]. Several authors describe MX precipitates as coherent, semi-coherent or even incoherent; however, taking in account both lattice misfit and particle size, coherency with the matrix is very unlikely. Howe’s [3] physical limit for misfit dislocation-based interfaces is at misfits of approximately 25% for which the spacing of misfit dislocations is close to 4b and their cores start to overlap. The large misfit measured for NbC in Sanicro 25 is very close to the Howe’s limit. In the case of TiC, a variant of MX phase with lower lattice misfit than NbC, octahedra with {111} facets and an interface consisting of a triangular network of misfit dislocations have been reported [4]. Our experimental data and structural characteristics were used as input in embedded atom model and phase field modelling to quantitatively confirm that the NbC precipitates indeed have the proposed shape and interface character with the matrix. Because of the large lattice misfit between the two crystal structures, precipitation of NbC in the defect-free matrix is extremely difficult. However, a large nucleation barrier can be overcome by nucleation at lattice defects, where part of the strain energy can be released by replacement of the dislocation line length and strain field with the precipitate nucleus. As the mode of
NbC precipitation is profoundly affected by plastic deformation, the overall characteristics of nanoprecipitates in terms of their size and distribution can differ significantly depending on loading conditions. An example is shown in Figs. 1a and 1b. After exposure at 700°C for 153 h, large particles of 50 nm in diameter are found nucleated only on dislocations. However, if cyclic loading (total strain amplitude 0.2%) is also present during aging, arrays of much smaller NbC carbides approximately 8 nm in diameter are found heterogeneously dispersed with locally varying volume density. Our experimental data confirm that at the early stages of nucleation, it is possible for dislocations to break away from the NbC, leaving behind a non-uniform distribution of nanoparticles. Moreover, once detached from dislocations, precipitates do not coarsen significantly. This conclusion is in agreement with Dutta et al. [5] who found that the coarsening mechanism of Nb(C,N) is governed by accelerated pipe diffusion of Nb atoms along dislocation cores. If the dislocation on which the precipitate nucleates is detached under cyclic stress, accelerated pipe diffusion is no longer in operation and the precipitates do not grow significantly. The dispersoid-like arrangement of nanoscale NbC particles can be then maintained with further thermal exposure.

In summary, Sanicro 25 acts as a “smart” system at nanoscale as NbC precipitates are able to heterogeneously pin dislocations under high temperature fatigue loading, but once broken free of dislocations, they do not coarsen rapidly. Fundamental understanding of structure, morphology and coarsening behavior of MX precipitates for different thermo-mechanical treatment conditions will contribute to the design of new fine-precipitation strengthened materials [6].

References:


Figure 1. HAADF-STEM image of Sanicro 25 alloy (a) aged only and (b) aged & cyclically loaded with total strain amplitude of 0.2% at 700°C for 153 h. (c) NbC precipitate overlapping with matrix and forming Moiré-like contrast in HAADF-STEM [001] zone axis imaging. (d) FFT pattern & schematic corresponding to image (c). (e) Simulated Moiré-like contrast. HAADF-STEM image simulated along [001] zone axis is compared with structural model of overlapping NbC and austenitic matrix.