Interpreting Voids in Atom Probe Tomography Data via Experiment and Theory

Jonathan D. Poplawsky,1,* Xing Wang,1 Constantinos Hatzouglo,2 Wei Guo,1 Ke Jin,3 Hongbin Bei,3 Yongqiang Wang,4 William J. Weber,3,5 Yanwen Zhang,3,5 Francois Vurpillot,2 and Karren L. More1

1. Center for Nanophase Materials Sciences, Oak Ridge National Laboratory, Oak Ridge, TN, USA.
2. Normandie Université, UNIROUEN, INSA Rouen, CNRS, GPM, Rouen, France.
3. Materials Science and Technology Division, Oak Ridge National Laboratory, Oak Ridge, TN, USA.
4. Materials Science and Technology Division, Los Alamos National Laboratory, Los Alamos, NM, USA.
5. Department of Materials Science and Engineering, University of Tennessee, Knoxville, TN, USA.
* Corresponding author: poplawskyjd@ornl.gov

Nanosized voids and/or pores are features that occur in many important materials for energy applications such as zeolites and nuclear materials [1-4]. Quantifying elemental segregation associated with voids at the nm-scale is important for understanding atomic-level material functionality as well as degradation mechanisms. For instance, He bubbles that form in irradiated structural materials cause unwanted swelling and embrittlement and can ultimately lead to failure of these materials [2,4]. Understanding elemental segregation effects associated with He bubbles in irradiated alloys is important for determining defect (vacancies and interstitials) diffusion/migration mechanisms, which is important for designing radiation-tolerant alloys. A commonly used method for nanoscale material characterization is electron microscopy; however, the small volume of individual voids with respect to typical scanning/transmission electron microscopy (STEM) lamella thicknesses make compositional quantification of the void shell difficult. The specified local electrode atom probe (LEAP) resolution ranges from 0.1 nm in z and 0.3 nm in x and y, making this an ideal technique for measuring the local compositions surrounding voids; however, the erratic ion flight paths from the void opening during field evaporation severely compromises the LEAP resolution. Understanding the field evaporation behavior of voids via atom probe tomography (APT) will enable improved methods for interpreting the reconstructed data and for comparing data acquired for different materials.

APT data for several single-phase concentrated solid-solution alloys (SP-CSAs), including NiCo, NiFe, NiCoCr, NiCoFe, and NiCoCrFe, was acquired and then analyzed using the reconstruction method currently available in IVAS that assumes a constant tip diameter for a hemispherically shaped needle. SP-CSAs are ideal specimens for understanding void evaporation behavior because the localized elemental segregation surrounding a small void acts as a marker for identifying the detector hit location of the void shell during field evaporation. For NiCoCr and NiCoCrFe, a correlative STEM/APT approach using the same needle-shaped sample was used to confirm the location of He bubbles in the APT datasets and to determine if He bubble sizes could be extracted from the APT data. Results of the correlative STEM and APT experiment for NiCoCr are shown in Figures 1a and 1b, respectively. Voids were consistently identified as high-density regions for NiCoCr and low-density regions for NiCoCrFe in the APT data.

Simulated APT experiments of needles containing voids were performed to explain the observed density phenomena, in which simulated detector data was reconstructed using the same method as the experimental APT data. The simulated APT needles contained voids with a 2 nm elemental segregation shell around the voids. The void size to needle radius was kept constant for the simulations while the evaporation field for the segregation layer around the void was varied with respect to the matrix. Three voids were included in the simulations and included segregation layer evaporation fields that were lower, higher, and the same as the matrix. In the APT reconstruction, the density and concentration profiles of the voids were different for the three different cases studied. The simulation for the high evaporation field case indicated a high-density region for the void, while the simulation for the low evaporation field case contained a low-density region for the void in the APT reconstruction. These high and low evaporation field simulations match the experimental data obtained for NiCoCr (shown in Figure 1) and NiCoCrFe, respectively. The Cr⁺/Cr⁻ charge state ratio within the void segregation region was compared to that of the matrix in the experimental data to determine the field difference between the segregation layer and the matrix.
The void segregation layer in NiCoCr showed a smaller Cr\(^{3+}/\)Cr\(^{2+}\) ratio than the matrix, whereas the segregation layer around the void in NiCoCrFe showed a larger Cr\(^{3+}/\)Cr\(^{2+}\) ratio than the matrix, indicating that the segregation layer in NiCoCr had a lower evaporation field than the matrix and that in NiCoCrFe had a higher evaporation field than the matrix. The simulated and experimental APT data are in excellent agreement, such that the simulations correctly predicted how the segregation layer around the voids evaporates and can be used to correctly interpret APT data and improve void/bubble reconstructions.

A qualitative model for void evaporation as well as best practices to capture, identify, and measure void sizes, and to quantify and compare segregation behavior around voids in APT reconstructions will be presented. A fundamental understanding of void evaporation processes in multi-element SP-CSAs will lead to more robust analyses of nano-void containing materials by APT [5].

References:

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Figure 1. a) HAADF-STEM image and b) APT atom map with iso-density surfaces highlighting high density regions in a He-irradiated NiCoCr alloy. (c) 1D line profile in z-direction of red-circled void in (b) showing Cr depletion in void and high-density region on top of void.