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MICROSTRUCTURAL AND MECHANICAL CHARACTERISTICS OF W-2Ti AND W-1TiC PROCESSED BY HOT ISOSTATIC PRESSING

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Abstract

W-2Ti and W-1TiC alloys were produced by mechanical alloying and consolidation by hot isostatic pressing. The composition and microstructural characteristics of these alloys were studied by X-ray diffraction, energy dispersion spectroscopy and scanning electron microscopy. The mechanical behavior of the consolidated alloys was characterized by microhardness measurements and three point bending tests. The mechanical characteristics of the W-2Ti alloy appear to be related to solution hardening. In W-1TiC, the residual porosity should be responsible for the poor behavior observed in comparison with W-2Ti.

1. Introduction

In the last years a great number of research works related to the development and characterization of W-based materials have been carried out [1-3]. It is due to the fact that W and W alloys are candidate materials to be used in the future fusion devices, in particular for the divertor and other first wall components [4-6]. W accomplishes some of the requirements for the plasma facing components of the fusion reactors, such as high melting point, good thermal conductivity, resistance to high thermal shocks, minimal tritium retention and low sputtering erosion rates. However, W has some critical issues that will have to be overcome or reduced for being used in fusion devices. In particular, it is brittle with a ductile-brittle transition temperature around 400 °C and a recrystallization temperature between 1200-1300 °C. Two possible strategies are considered for increasing W ductility: alloying and nanostructuring [7-9]. Concerning alloying, W-Re alloys are more ductile than pure W, however they have been rejected for fusion applications because they suffer irradiation embrittlement due to Re impurity transmutation into deleterious impurities, besides being very expensive [10,11]. Moreover, other property required to the first wall materials is to be classified as a low or reduced activation material, thus only Fe, Cr, V, Ti, Si and C should be considered as potential alloying elements. On the other hand, materials with an ultrafine-grained (UFG) microstructure usually exhibit enhanced ductility. In addition, it would be expected that a UFG microstructure improves the irradiation resistance of W, as grain boundaries can be strong sinks for the irradiation induced defects. In relation to the recrystallization temperature, a fine dispersion of hard particles such as La_2O_3 , Y_2O_3 or TiC may have capability for inhibiting grain growth and increases the recrystallization temperature. W materials are usually fabricated by powder metallurgy routes due to its high melting point. In the case of processing W alloys, the mechanical milling is a critical step for alloying and attaining both grain refinement and homogeneous solution or dispersion of

nanosized particles, i.e. for developing a nanostructured material [12]. For achieving a consolidated W material with a suitable density and mechanical properties, very high sintering temperatures under reducing conditions are required. Hot isostatic pressing (HIP) is a consolidation technique that is very successful for producing W alloys, in particular untextured UFG oxide dispersion strengthened (ODS) W-Ti and W-V alloys containing a nanosized dispersion of oxide particles [13,14]. Furthermore, these research works have shown that both Ti and V are sintering activators for the W alloys, as their addition increases the densification degree. According to the W-V and W-Ti phase diagrams, the solubility in W is total for any V content and for a weight percent of Ti below 10 %. For small contents of these impurities the melting temperature of the alloy would maintain quite close to that for pure W. Recently, it has been showed that Ti addition might alter the core structure of the screw dislocations and their slip mechanism increasing W ductility such as it occurs for Re impurities [15]. The present communication reports the development of W-2%Ti and W-1%TiC (wt.%) materials processed by mechanical alloying and HIP. For W-2Ti, the goal of the mechanical milling process will be to achieve the proper Ti alloying in W, and to analyze its effect on the mechanical behaviour of the alloy. For W-1TiC the purpose is to achieve a good dispersion of the TiC nanoparticles in the W matrix. The characterization of the materials during the different steps of the processing has been carried out to establish the optimum parameters in the fabrication process. The microstructure of the consolidated materials has also been studied, and the particular mechanical properties explored.

2. Experimental procedure

The alloys were prepared from W and Ti elemental powders with purity of 99.9 %, and 99 % and pure nano-sized TiC powder (Skyspring Nanomaterials). The particle sizes of the elemental powders were in the range 1-5 μm for W, and -150 mesh ($<100 \mu\text{m}$) for Ti. The

average particle size for TiC was ~ 40 nm. The processing of the materials consisted of: 1) blending of the starting powders in a Turbular T2F mixer for 4 h; 2) mechanical alloying of the blends at 150 rpm in a high-energy planetary ball mill; 3) alloyed powder encapsulation and degassing; and 4) consolidation by HIP for 2 h at 1573 K and 195 MPa. The mechanical alloying was carried out under a high purity Ar atmosphere in a WC lined pot, using WC balls of 10 mm in diameter as grinding media at a ball-to-powder ratio of 4:3. The milling time was 75 hours for W-2Ti and 50 hours for W-1TiC. Before the consolidation, the alloyed powder encapsulated in a steel container was degassed for 24 h at 400 °C in a vacuum of $\sim 2 \times 10^{-3}$ mbar. This degassing process removes part of the volatile components that could have been incorporated to the powder during the milling process and handling.

The O content in the powders and consolidated materials was determined using a LECO TC500 analyzer. The initial O content of the elemental W, Ti and TiC powders was 0.037 %, 0.069 % and 4.5%, respectively, after handling in the glove box in Ar atmosphere. The density measurements of the consolidated alloys carried out in a He ultracycrometer resulted in ~ 98 % for W-2Ti and ~ 95 % for W-1TiC of the expected densities calculated applying the mixture rule for the corresponding composition (Table I). The evolution of the particle morphology and structure of the powder during the milling process, and the microstructure of the consolidated alloys were investigated by scanning electron microscopy (SEM) and X-ray diffraction (XRD). For compositional and microstructural analyses of the milled powder by SEM and energy dispersive spectroscopy (EDS), powder samples were embedded into a conductive epoxy resin and the surface of the specimens suitably polished. The microstructure was also studied with an atomic force microscope (AFM) (XE-100, Park Systems). XRD patterns were analyzed by the Rietveld method using the Fullprof software [16]. To obtain correct measurements of the average crystallite size and crystal lattice strain produced by the milling process, the contribution of the diffractometer to the peak shape was

determined from a precise XRD pattern from a reference sample. The instrumental and the microstructure effects on the peak broadening were extracted applying the integral breadth analysis assuming a pseudo-Voigt function for the peak shapes and isotropic broadening for strain and crystallite size [17].

The Vickers microhardness was measured applying a load of 2.94 N for 20 s. The flexural strength of $2 \times 2 \times 25 \text{ mm}^3$ samples in vacuum was determined from three-point bending tests carried out in a universal testing machine (Instron. Model 8800), at a crosshead speed of 0.1 mm/min, with the samples on a self-aligning stage having a span of 16 mm. The stage with the sample was placed at the center of a high vacuum furnace mounted onto the testing machine, heated up to the test temperature using a heating rate of 5 °C/min and a dwell time of 15 min. At least, two samples were tested for each temperature. Before testing, the samples were annealed for one hour at 1200 °C in H₂-rich atmosphere to relieve the possible residual stresses induced by the electro-discharge machining and preparation of the samples.

3. Results and discussion

3.1 Milled powders

Although the powder handling and milling were carried out under a high purity Ar atmosphere, oxygen contamination was obvious. Fig. 1 shows a continuous increase of the O content with a similar increasing rate for both alloys, which indicates that the O intake is the same irrespective of the composition of the powder, at least for 50 h milling under the present conditions. Let us point out that the initial O content in the W-1TiC blends is remarkably higher than in W-2Ti.

The milling effect on the XRD patterns is shown in Fig. 2. The broadening of the diffraction peaks is an obvious result of the grain refinement and plastic deformation in the powder particles. After milling for 5 h the XRD patterns exhibit diffraction peaks that correspond to

a single bcc phase; peaks from hcp Ti or fcc TiC are not detected. The evolution of the average crystallite size and lattice strain with milling time is shown in Fig. 3. The cumulative grain refinement and lattice strain for W-1TiC is apparently faster than for W-2Ti. From the linear increase of the lattice strain with milling time a strain rate of 1.00×10^{-4} and $1.33 \times 10^{-4} \text{ h}^{-1}$ are respectively determined for W-2Ti and W-1TiC. At least milling for 25 h at the present conditions are required for getting grain refinement with average crystallite size of ~ 20 nm in W-1TiC, but in the case of W-2Ti, 50 h are necessary for attaining similar crystallite sizes. The addition of 1 wt.% of hard TiC particles in W-1TiC, in contrast with 2 wt.% of ductile Ti in W-2Ti, accounts for the observed evolution of crystallite size and lattice strain. During milling a significant part of the impact energy would be consumed to produce solid solution alloying, in the case of W-2Ti in addition to plastic deformation, particle fragmentation, grain refinement and cold welding of particles. However, in W-1TiC the impact energy consumption for interdiffusion or solid solution would not be significant if it is accepted that the TiC dispersion is directly accomplished by the mechanical effect of the milling, and not via TiC decomposition, i.e. Ti and C dissolution in the W lattice, and finally re-precipitation into nanosized TiC particles. Fig. 4 shows backscattered electron (BSE) images of the W-2Ti and W-1TiC powders after being mechanically alloyed for 75 and 50 h, respectively. For W-2Ti, the BSE image (Fig. 4a) reveals the presence of sub-micron Ti-rich areas, indicated by arrows in the image. EDS analyses performed on the tungsten matrix indicate a Ti fraction slightly below the nominal one, indicating that a nearly complete solid solution is attained after milling for 75 h under the present conditions. In W-1TiC milled for 50 h (Fig. 4b), the observed sub-micron Ti-rich regions correspond to TiC particles. These particles are homogeneously distributed indicating that a good dispersion of submicron TiC particles may be developed in W under the present conditions of milling.

3.2 Consolidated alloys

The Vickers microhardness of consolidated alloys, along with the densities and O contents are given in Table I. As expected, the relative density of the alloy W-2Ti alloy is significantly higher than the one measured for the W-1TiC alloy because Ti favors the densification of the material [20]. The O contents measured in the consolidated alloys were considerably lower than corresponding contents in the as-milled powders. The Vickers microhardness for W-2Ti is twice that for W-1TiC, i.e. 14.0 against 7.0 GPa. The value obtained for W-1TiC is somewhat lower than the one of 8.1 reported for W-5TiC [20], and that found for W-0.5TiC (8.8 - 10.8 GPa, depending on the grain size) [21]. Figs. 5 and 6 show SEM and AFM images of the consolidated alloys. Both alloys exhibit an intermingled grain structure that consists of a population of submicron-sized grains and another one of coarser grains with sizes $\sim 1 \mu\text{m}$ (see Figs. 5b and 6b). In the case of W-2Ti, the EDS-SEM analyses reveal the presence of small Ti pools localized in the interstices between some W prior particles. The formation of similar pools has also been reported in ODS W-Ti and ODS W-V alloys processed under similar conditions but mechanically alloyed for shorter period of time [18, 19]. Since the milling time for the present alloys was up to three times longer than those for the ODS W alloys and the formation of Ti pools is also produced, the present results confirm that this phenomenon is due to Ti segregation to the prior particle boundaries induced by the HIP process.

3.3 Three-point bending tests

Fig. 7 shows the temperature effect on the bending behavior of the alloys. The stress-strain curves for W-2Ti reveal plastic deformation at $T \geq 1000 \text{ }^\circ\text{C}$ and completely brittle behavior at lower temperatures. The bending strength and the yield strength at 0.2 % of plastic

deformation are calculated using the equation:

$$\sigma = \frac{3PL}{2tw^2} \quad (1)$$

and the strain is determined from the expression:

$$\varepsilon = \frac{6w\delta}{L^2}100 \quad (2)$$

where P is the breaking load, L is the span (distance between the support points), w is the specimen width, t is the thickness of the sample and δ is the deflection. This alloy appears to exhibit better bending strengths at low temperature than corresponding strengths reported for the W-4Ti alloy, which did not show plastic deformation up to 1000 °C [14]. The W-1TiC alloy exhibits plastic deformation at 1200 °C and a completely brittle behavior at $T \leq 1000$ °C. The yield strength values at 0.2% of plastic strain above 1100 °C are shown in Table II. It should be noticed that the bending strength at 1100 °C for other W alloys processed by the same route is noticeably lower, i.e. around 300 MPa for pure W and around 350 MPa for ODS W-2Ti-0.5Y₂O₃ and W-4Ti-0.5Y₂O₃ [18, 22] .

According to the three point bending tests, these W-2Ti and W-1TiC alloys have the ductile-brittle transition above 1000 °C, which is higher than that of pure W. The powder metallurgy route is able to dissolve a part of the Ti particles in the W matrix for W-2Ti and, in case of W-1TiC, a certain amount of TiC particles are well dispersed inside the W grains. However, the compositional and microstructural analyses indicated that Ti and TiC particles are found segregated at the grain boundaries. Furthermore, a non-negligible O concentration is also present in the alloys, which may also be segregated at the grain boundaries forming complex oxides. These phases segregated would develop a high concentration of internal stress in the grain boundaries promoting the grain boundary decohesion and brittleness of the alloys, as it occurs in Mo alloys [23]. In the case of W-1TiC, the segregation of TiC particles at the boundaries might be avoided by substituting the mechanical alloying by

some wet-chemical process like that applied for processing Mo alloys [23].

4. Conclusions

W-2Ti and W-1TiC processed by mechanical alloying and HIP exhibit a grain structure containing a population of submicron-sized grains and another of coarser grains with sizes $\sim 1 \mu\text{m}$. The XRD measurements and the EDS analyses of the alloyed W-2Ti powders reveal the virtual dissolution of Ti into W under the present milling conditions. However, after HIP consolidation a fraction of Ti is found segregated in the interstices among prior powder particles. The SEM and EDS analyses show an apparent dispersion of TiC into W in the alloyed W-1TiC powder, but very small TiC particles appears decorating grain boundaries of the consolidated material.

The high microhardness value and the mechanical behavior at high temperature of the W-2Ti alloy may be attributed to solid solution hardening and grain boundary decohesion due to Ti segregation. In contrast, the much lower microhardness value in W-1TiC, as well as the bending strain at 1200 °C, could be mainly caused by residual porosity excess since the densification grade attained was low, i.e. 95.3% against 98 % for W-2Ti.

Acknowledgements

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Figure captions

Fig. 1. Oxygen content versus milling time for W-2Ti and W-1TiC powders.

Fig. 2. XRD patterns for W-2Ti and W-1TiC powders: blended (solid) and after mechanical alloying (dashed) for 75 and 50 hours, respectively.

Fig. 3. Effect of milling time on the average crystallite size and internal strain of the W-2Ti and W-1TiC powders.

Fig. 4. EBS-SEM images of powder particles. (a) W-2Ti milled for 75 h; the arrows point to sub-micrometer Ti-rich regions. (b) W-1TiC milled for 50 h; the arrows indicate sub-micrometer TiC particles.

Fig. 5. Microstructure of consolidated W-2Ti. (a) EBS-SEM image; the Ti pools are indicated by arrows. (b) AFM image.

Fig. 6. Microstructure of consolidated W-1TiC . (a) EBS-SEM image. (b) AFM image.

Fig. 7. Stress-strain curves of three point bending tests for W-1TiC and W-2Ti at different temperatures.

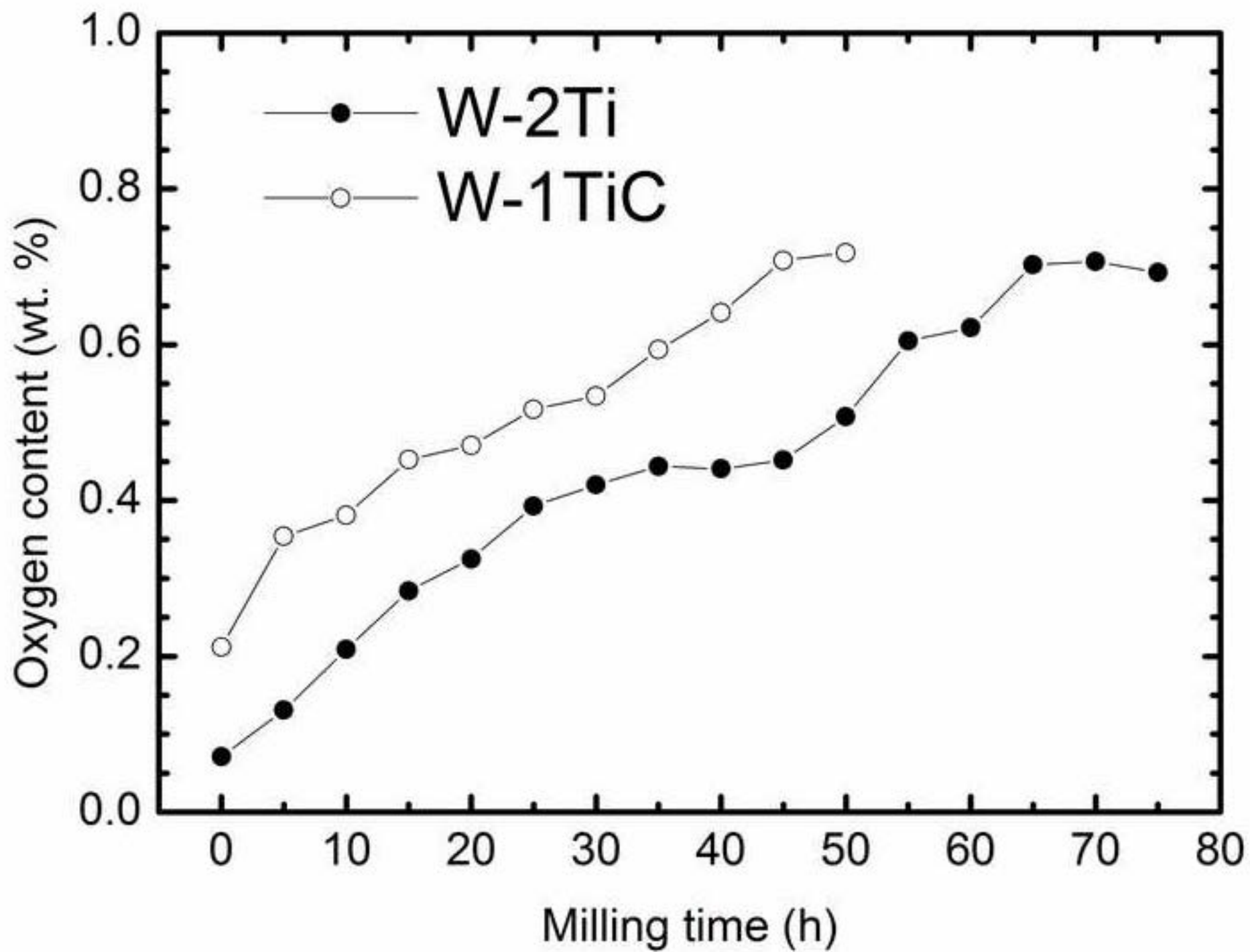
Table I. Density, oxygen content and Vickers microhardness for consolidated W-2Ti and W-1TiC .

	ρ_{theo} (g/cm ³)	ρ_{expe} (g/cm ³)	$\rho_{\text{exp}}/\rho_{\text{theo}}$	O (wt.%)	Hv (GPa)
W-2Ti	18.069	17.708 ± 5	98.0 %	0.418	14.0 ± 0.7
W-1TiC	18.706	17.827 ± 9	95.3 %	0.454	7.0 ± 0.4

Table II. Three point bending yield strength for W-2Ti and W-1TiC.

	1100 °C (MPa)	1200 °C (MPa)
W-2Ti	526 ± 4	381 ± 4
W-1TiC	---	340 ± 10

Figure 1
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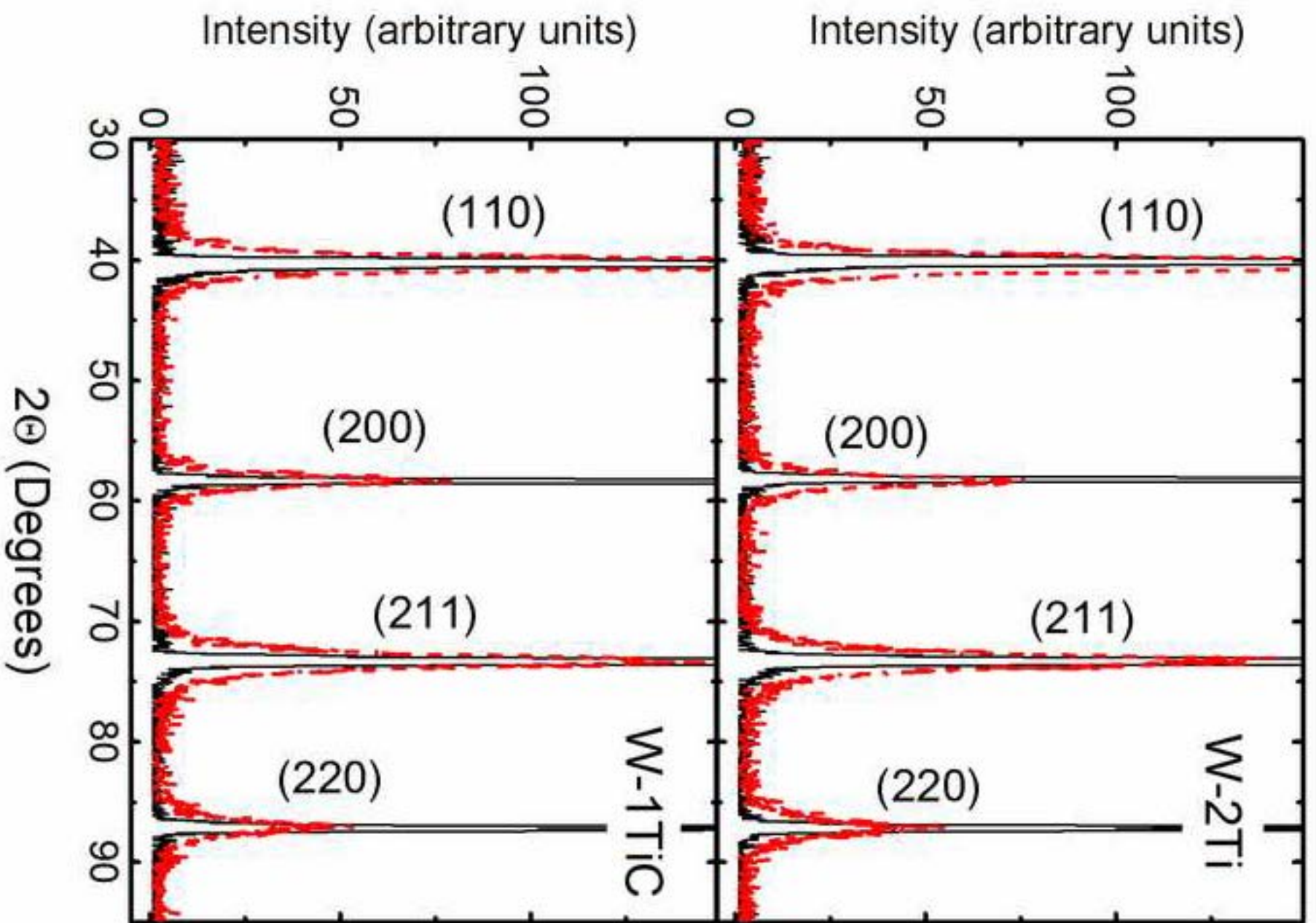


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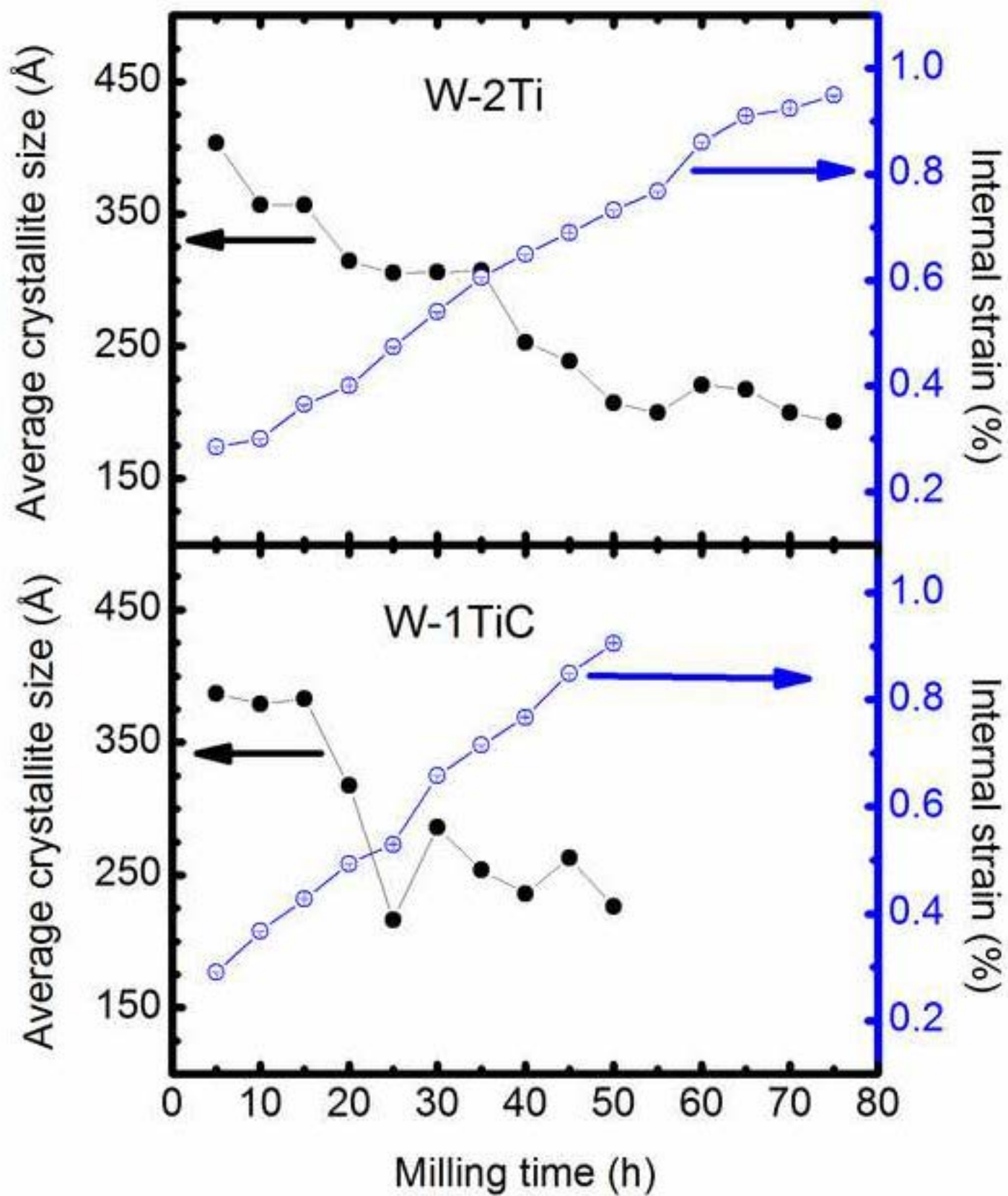


Figure 4
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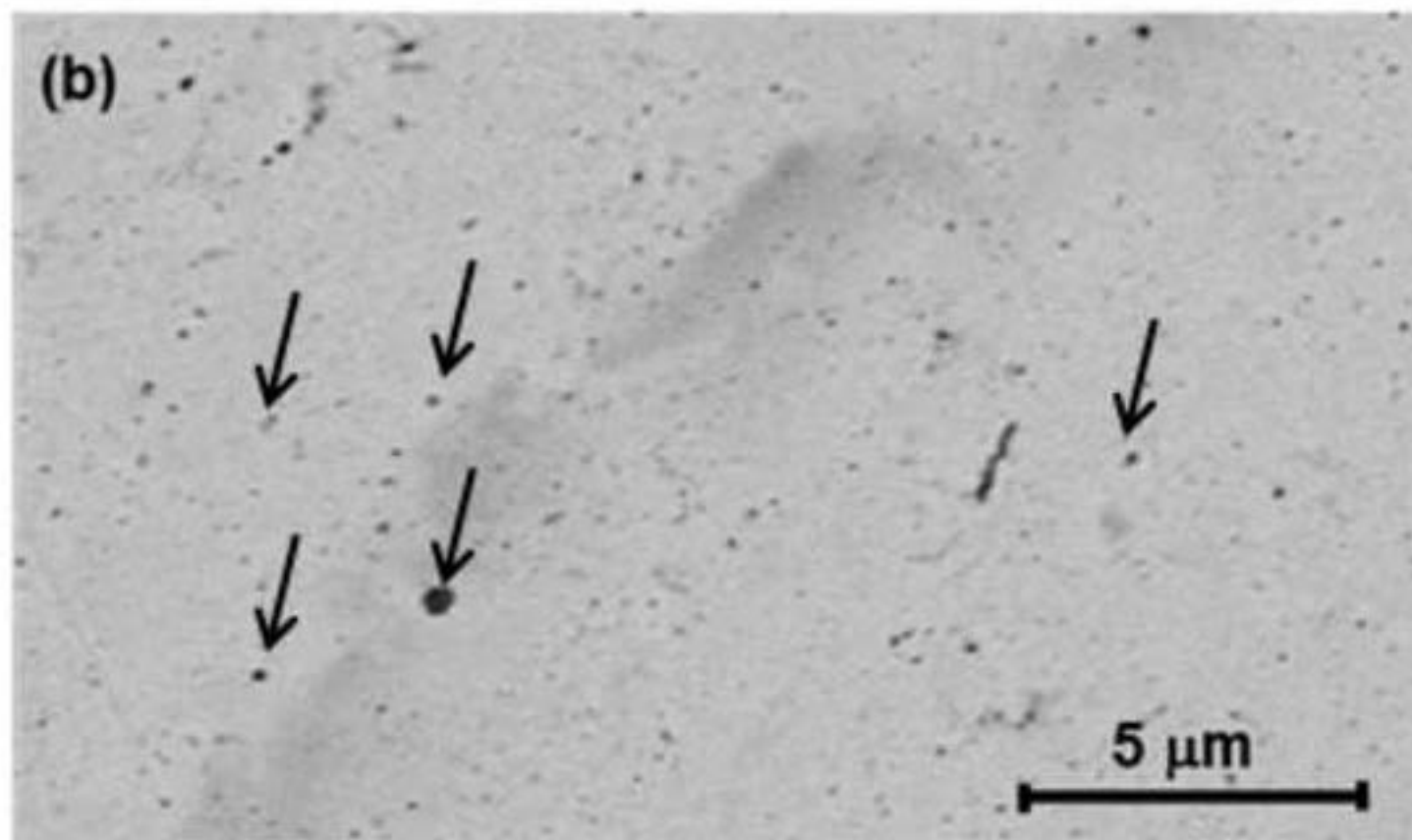
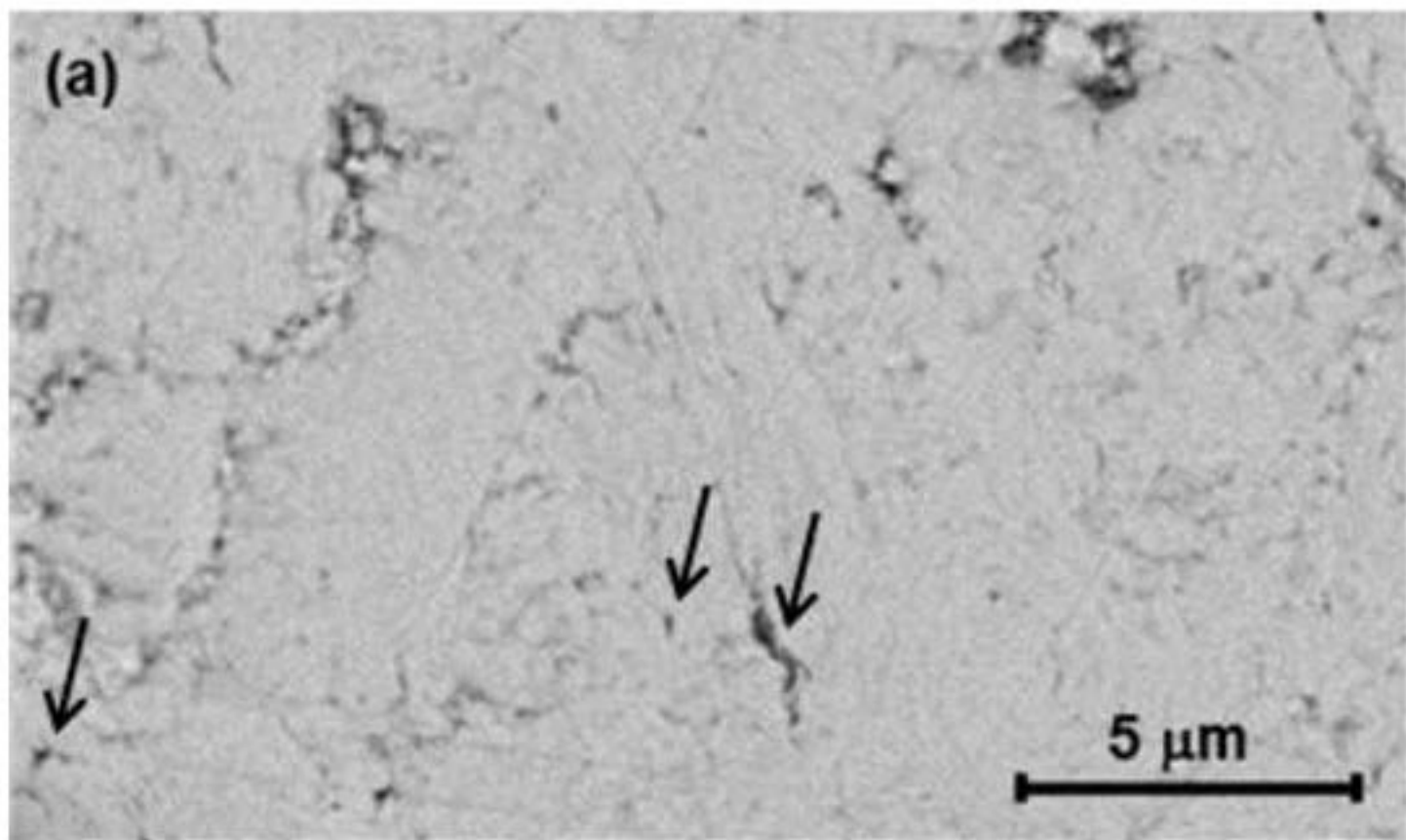


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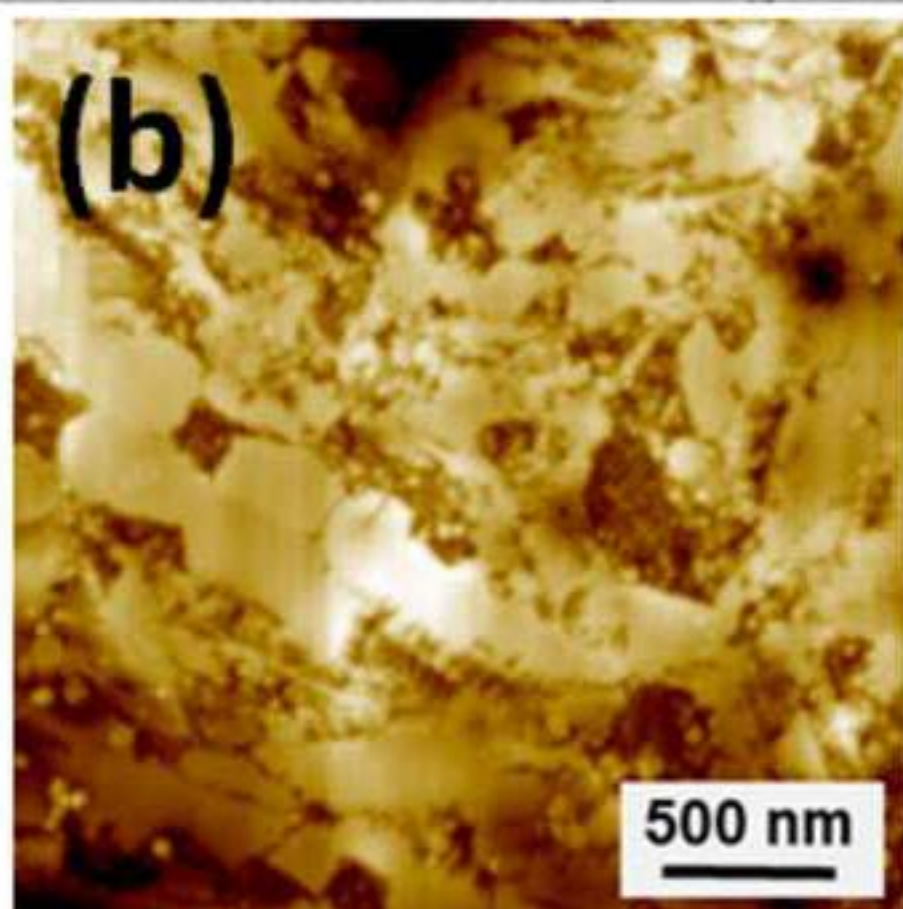
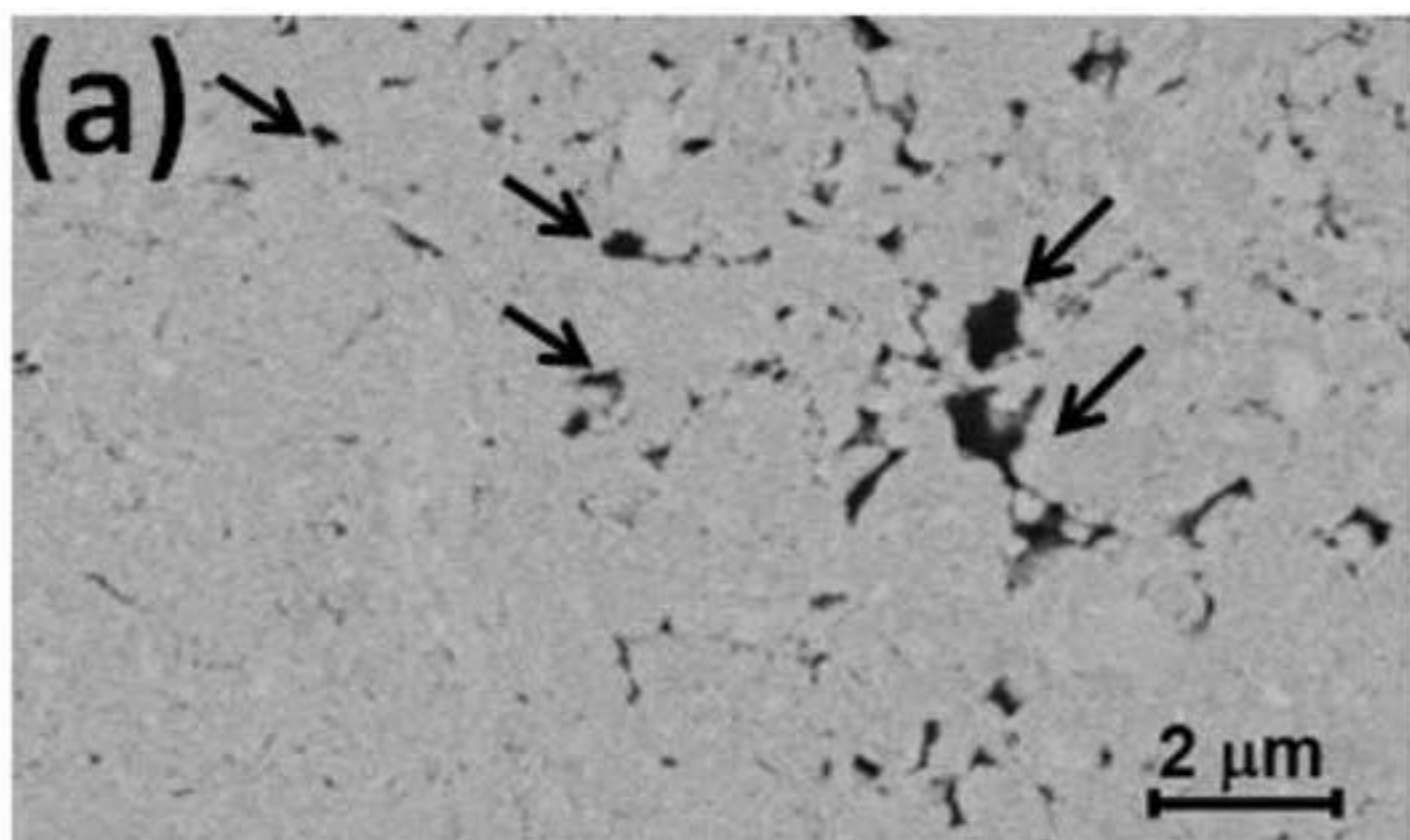


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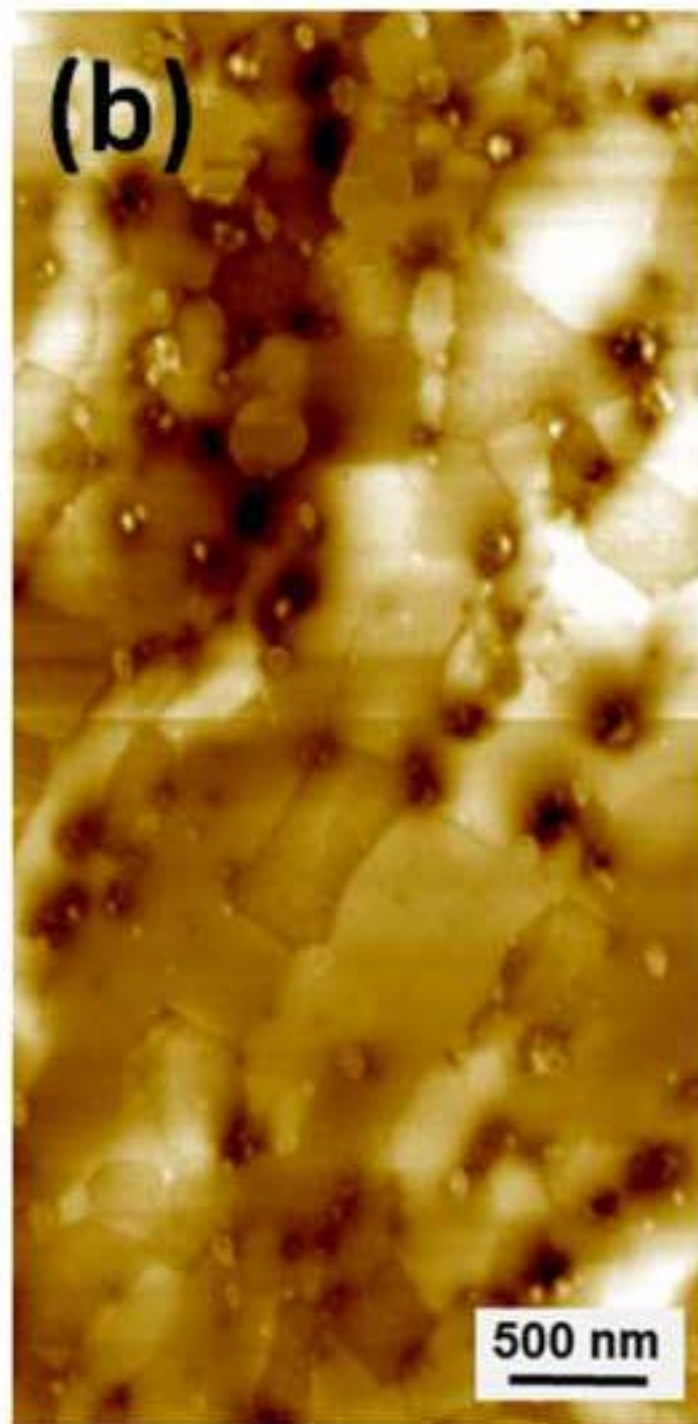
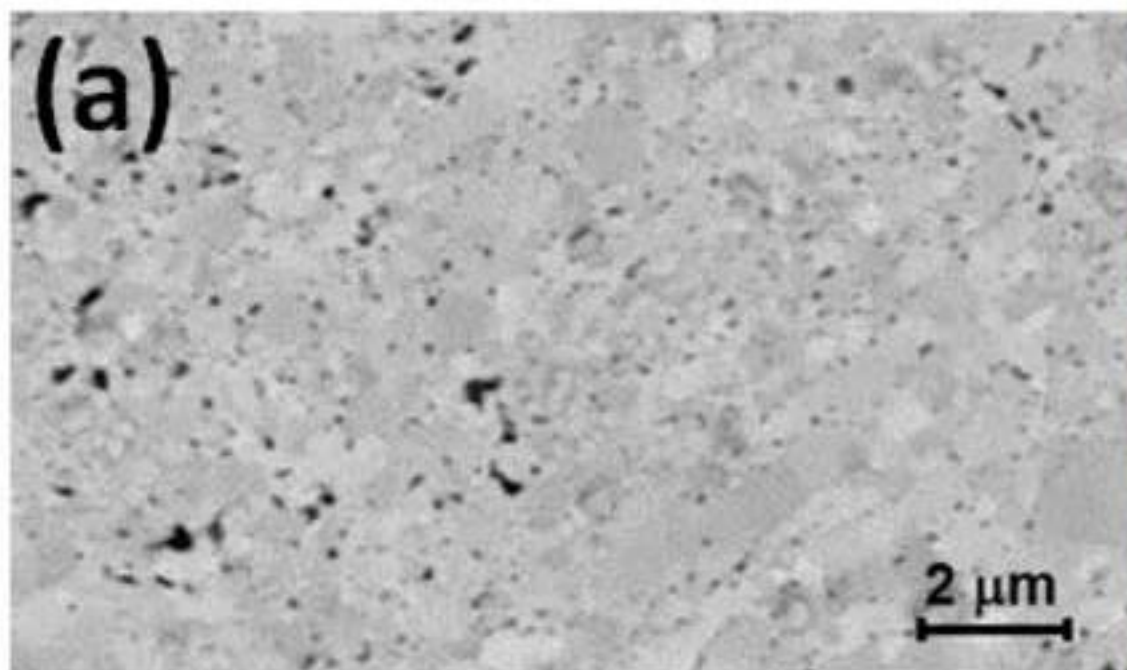


Figure 7
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