

Characterization of novel W alloys produced by HIP

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Abstract: W and W alloys containing 0.5 wt% Y₂O₃, x wt% Ti and (x wt% Ti + 0.5 wt% Y₂O₃) have been prepared, x = 2 or 4. Elemental powders were blended or ball milled, canned, degassed and finally consolidated by a two stage HIP process under a pressure of 195 MPa. It is found that Ti addition favours the densification attaining a fully dense material. XRD, SEM and EDX analyses of the material with Ti addition reveal the formation of a microstructure consisting of tungsten particles embedded in a W-Ti matrix. The microhardness of these materials increased noticeably with the titanium content.

1. Introduction

W is considered as a candidate material for plasma facing components (PFCs) in a future fusion power reactor because of its refractory characteristics, low tritium retention and low sputtering yielding [1,2]. However, its use in PFCs requires the development of W alloys that, in addition to these properties, maintain good mechanical properties after a prolonged exposure at high temperature. Sintering would be the most suitable method to produce W materials for these applications if their recrystallization temperature is high enough and the grain growth is restrained. The usual sintering conditions for W require very high temperatures that induce a coarse grained structure in the sintered material, and a low recrystallization temperature in the hot worked material. This causes the failure of its mechanical properties. The combined addition of a sintering activator, which lowers the sintering temperature and favours the densification, and an insoluble oxide that produces a dispersion strengthening and grain growth inhibition, may result in a W material with improved mechanical characteristics.

Moreover, oxide dispersed W alloys have been envisaged as potential structural materials for a modular He cooled divertor of the fusion demonstration reactor DEMO [1]. In particular, conceptual designs based on sintered W tiles joined to a W-La₂O₃ thimble structure are being considered [3,4]. In the case of developing a W alloy suitable for PFCs, a critical issue would be the joining between different W material components. Joining experiments performed on sintered W/W-La₂O₃ have demonstrated the feasibility of the joint if a Ti interlayer is used [3]. These results suggest that W/W-Ti would be more effectively joined than W/W-La₂O₃. Then,

oxide dispersed W-Ti alloys appear to be promising materials for the fabrication of divertor components.

The present work assesses the possibility of using jointly Ti as sintering activator and Y₂O₃ particles as strengthening dispersoids in W.

2. Experimental

The starting materials were 99.9% pure W and 99.8% pure α -Ti powders with an average particle size of 14 and 20 μ m, respectively, and 99.5% pure nanometric Y₂O₃ powder with particle sizes between 10 and 50 nm. Powder blends with compositions W-4 wt% Ti were prepared mixing together the powder inside a container sealed under a high purity Ar atmosphere. Blends with compositions W-0.5 wt% Y₂O₃, W-4 wt% Ti-0.5 wt% Y₂O₃ and W-2 wt% Ti-0.47 wt% Y₂O₃ were mixed by high energy milling for 2 h in a planetary mill under a high purity Ar atmosphere. In this case, a WC container and WC balls of \varnothing 10 mm, as grinding media, were used. The blends were canned and degassed at 673 K for 24 h in vacuum and then the can sealed. First, the cans were HIP treated at 1550 K for 2 h at a pressure of 195 MPa. After removing the can, the consolidated billet underwent a second HIP at 1973 K for 30 min in a pure Ar atmosphere at 195 MPa.

The microstructural characteristics were investigated using light microscopy, scanning electron microscopy (SEM) and energy dispersive X-ray (EDX) analyses. Moreover, X-ray diffraction (XRD) analyses were made. Vickers microhardness measurements were performed applying a load of 2.94 N during 20 s. Transmission electron microscopy (TEM) observations and EDX analyses were performed on ion milled samples in a Philips Tecnai microscope at 200 kV. The ion milling conditions were $\theta = 6^\circ$, $V = 7$ kV and $I = 2.6$ mA.

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3. Results and discussion

3.1. W and W 0.5Y₂O₃

After the first HIP at 1550 K the consolidated W material resulted in a relative density of 92.7%. The second HIP at 1973 K produced no meaningful increase in the density. Fig. 1a and b show the microstructure of the W material after the second HIP stage. A strong pore shrinkage and formation of high angle grain boundaries induced by coalescence of the tungsten particles were observed. The mean grain size D_{50} for W was $\sim 4.4 \mu\text{m}$ after the second HIP at 1973 K.

The microstructure of the W 0.5Y₂O₃ material HIP treated at 1550 K was similar to that corresponding to W. Pores appear mainly formed at triple or quadruple junctions between the W grains, see Fig. 2a. The relative density and the mean grain size measured after the first HIP treatment result in $\sim 90\%$ and $\sim 4.0 \mu\text{m}$, respectively. EDX analyses showed no evidence of Y and O on the sample surfaces. Despite of the open porosity, estimated in $\sim 8\%$ from metallographic analyses, the porosity of this material decreased after the second HIP treatment. Fig. 2b and c show micrographs of W 0.5Y₂O₃ after the second HIP treatment at 1973 K. SEM EDX analyses revealed the presence of a new phase containing Y and O in the residual marks of previous pores like areas 1 and 2 in Fig. 2b and c. Fig. 2c shows the appearance of grains of these oxides surrounding a W particle; the average composition of these grains correspond to 51 at.% O, 21 at.% W and 18 at.% Y. The mean size of the oxide grains was estimated in $0.62 \mu\text{m}$.

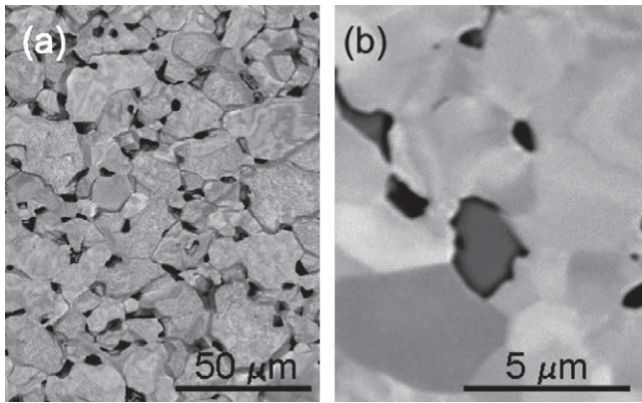


Fig. 1. SE-SEM micrographs of W consolidated by HIP after 30 min at 1973 K and 195 MPa (a) and (b) etched with Murakami reagent.

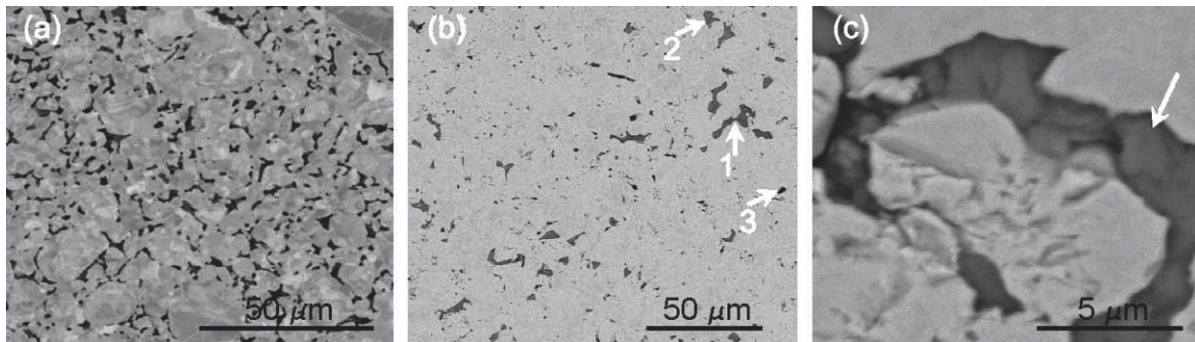


Fig. 2. BSE-SEM micrographs of W-0.5Y₂O₃ after HIP consolidation: (a) 2 h at 1550 K and 195 MPa, (b) and (c) after second HIP for 30 min at 1973 K and 195 MPa. The samples have not been etched to avoid the preferential etching of the new (W - Y) oxide phase (dark grey contrast) in areas 1 and 2 of (b), and white arrow (c).

The new (W - Y) oxide phase filled most of the pores present after the first HIP treatment stage. This evidences the pore shrinkage by segregation of the new phase, and it may be the main cause of the apparent densification of the alloy. The relative density can not be accurately determined because of the appearance of this new phase. However, from metallographic analyses the volume fraction of this phase was estimated in $\sim 3.8\%$ and the porosity in $\sim 0.2\%$.

The TEM observations performed on W 0.5Y₂O₃ after the second HIP treatment showed tungsten grains with typical sizes between 2 and 5 μm . These sizes might not be representative of the characteristic grain size of the sample because only small areas of sample were transparent to the electron beam. These tungsten grains and their boundaries were free of second phase precipitates and the EDX analyses did not detected presence of impurities. Moreover, they were apparently dislocation free. In some visible areas of these W 0.5Y₂O₃ samples, small grains of phases containing O, Y, W, Cr and Ti appeared alongside of the tungsten grains, as Fig. 3 shows. These grains, which had sizes $< 1 \mu\text{m}$, exhibited different composition and crystallographic structure from each other. The analyses revealed that the grains were complex oxides containing Y and W, and unwanted Cr and Ti impurities. These impurities might be either concomitant impurities of the starting powders or contamination from the pot and grinding media. The

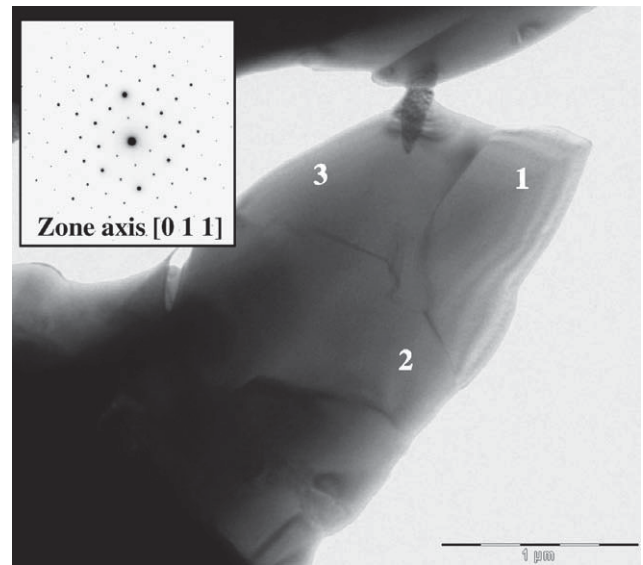


Fig. 3. TEM image showing the complex oxides formed alongside of the W grains in W-0.5Y₂O₃ samples after the second HIP treatment at 1973 K. The SAD pattern with zone axis [011] corresponds to grain 1.

identification of these oxides from the selected area diffraction patterns (SAD) is difficult because agreement with the lattice parameters reported for Y–W oxides have not found. For instance, the grain denoted by 1 in Fig. 3 is identified from the SAD patterns as an fcc crystal with a lattice parameter of 0.989 ± 0.005 nm. TEM analyses are still in progress to identify these complex oxide phases.

3.2. W–4Ti alloy

The W–4Ti material obtained after the first HIP treatment resulted in a density of 17.09 ± 0.05 g/cm³ in agreement to the theoretical value. After the first HIP treatment practically the whole content of Ti appears as β Ti, only a meaningless residual signal from α Ti was detected. The analyses of the patterns using the program Fullproof suggest that the observed peaks could be the convolution of two peaks corresponding to two different bcc structures with very similar lattice parameters. After the second HIP, evidence for α Ti disappeared and the diffraction peaks slightly shifted to positions corresponding to solid solutions with less Ti content.

The microstructure of the W–4Ti material after the first and the second HIP is shown in Fig. 4. Pools of Ti(W), which are the black spots surrounded by W grains, were observed uniformly distributed, but no pores were observed. Fig. 4b shows the formation of areas into the Ti pools with a very high W content after HIP at 1550 K. After the second HIP, the W particles are surrounded by an inter particle W–Ti phase with a Ti content of ~ 7.2 wt.% as Fig. 4d reveals. The second HIP treatment reduced the number and size of these pools and produced a bimodal distribution of grain sizes due to grain growth in the larger Ti pools, as Figs. 4 and 5 illustrate. The effect of the HIP treatments on the cumulative distribution of grain sizes in the Ti pools is shown in Fig. 5. Despite of the grain growth in the large pools, it should be noted that the average grain size D_{50} did not increase, see Table 1. This fact suggests that small Ti(W) grains formed after the second HIP treatment.

3.3. W–Ti–Y₂O₃ alloys

Both milled powders, W–2Ti–0.47Y₂O₃ and W–4Ti–0.50Y₂O₃, rendered fully dense materials after the first HIP treatment. The

measured final density was 16.8 g/cm³ and 16.9 g/cm³, respectively. Fig. 6 shows the microstructure of these materials. No pores, and a bimodal distribution of Ti pools similar to that observed in W–4Ti, were also found in these materials after the first HIP treatment. The distribution of Ti pools in W–4Ti–0.50Y₂O₃ was apparently equal to the one for W–4Ti. The second HIP treatment tended to make round the small Ti pools and reduced the size of the large ones. In the case of W–2Ti–0.47Y₂O₃, SEM and EDX analyses showed a distribution of Ti pools less dense and with a number of large Ti pools smaller than the corresponding to W–4Ti–0.50Y₂O₃. The large pools tended to be more elongated than in W–4Ti–0.50Y₂O₃, and the small ones appeared to be outlining the W particles. The XRD patterns of these alloys exhibited very similar behaviour to that already described for W–4Ti, but no α Ti phase in W–2Ti–0.47Y₂O₃ was detected after the first HIP at 1550 K. SEM images of W–2Ti–0.47Y₂O₃ HIP treated at 1550 K revealed inter particle areas with dark grey contrast, and irregular and diffuse contour, see Fig. 6a. EDX analyses of these areas resulted in compositions of W (4–5) wt.% Ti without any signal of Y or impurities. As in W–4Ti, W particles were surrounded by a W(Ti) phase with a Ti content of ~ 7 wt.%, as the areas indicated by black arrows in Fig. 6f. EDX analyses revealed that the second HIP at 1973 K induced the formation of complex oxide aggregates in the previous Ti pools. Ti, Y and O in a variable concentration were present in these

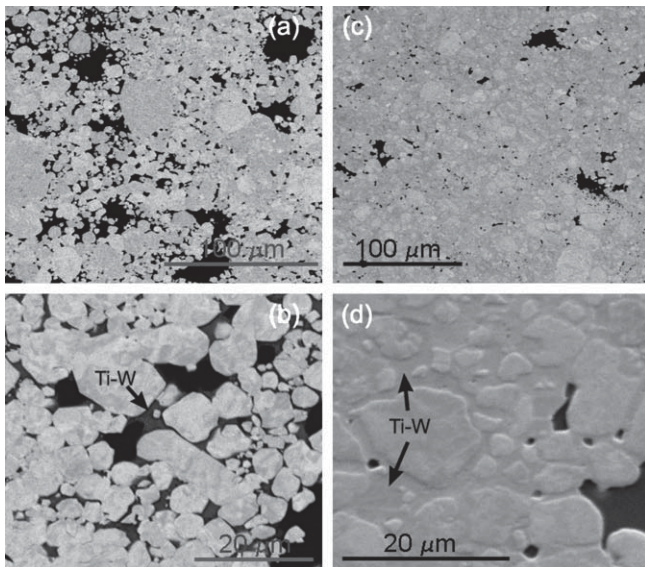
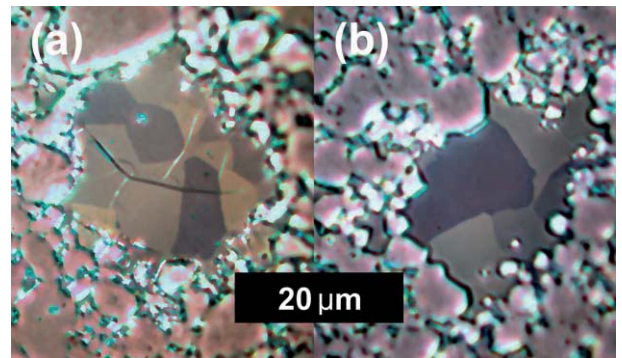


Fig. 4. BSE-SEM images of the W–4Ti material. (a) and (b) after the HIP at 1550 K, (c) and (d) after the second HIP at 1973 K.

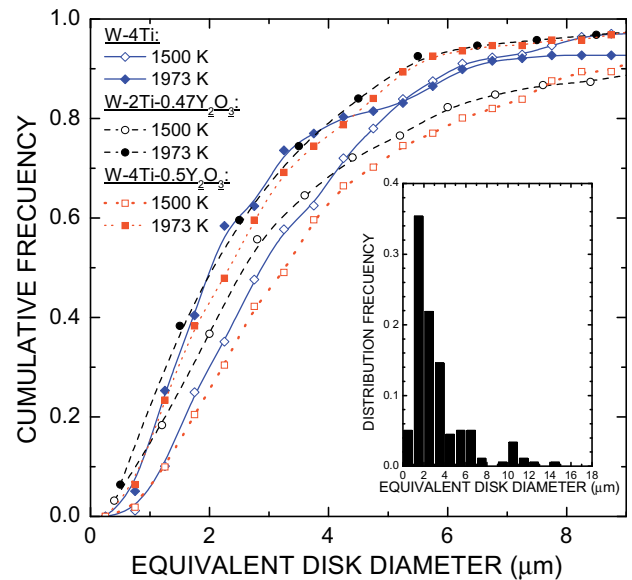


Fig. 5. Micrographs of W–4Ti using crossed polarized light illumination showing the grain growth in the larger Ti pools; after HIP at (a) 1550 K and (b) 1973 K. The graph depicts the cumulative size distributions of the grains in the Ti pools after the HIP treatments at 1550 K and 1973 K, for: (◇, ◆) W–4Ti, (○, ●) W–2Ti–0.47Y₂O₃ and W–4Ti–0.50Y₂O₃ (□, ■). The inset shows the grain size distribution for W–4Ti after HIP at 1973 K.

Table 1

Grain size in the Ti(W) pools corresponding to 50% of the cumulative size distribution and Vickers microhardness values for the materials successively HIP treated.

Material	Grain size D_{50} for inter-particle Ti(W) phase		Vickers microhardness	
	HIP at 1550 K (μm)	HIP at 1973 K (μm)	HIP $T = 1550$ K (GPa)	HIP $T = 1973$ K (GPa)
W	–	–	2.91 ± 0.10	2.85 ± 0.10
W-0.5Y ₂ O ₃	–	–	2.65 ± 0.15	2.44 ± 0.05
W-4Ti	2.6 ± 0.3	2.0 ± 0.3	4.46 ± 0.04	4.84 ± 0.12
W-2Ti-0.47Y ₂ O ₃	2.5 ± 0.3	1.9 ± 0.3	4.61 ± 0.12	3.6 ± 0.5
W-4Ti-0.50Y ₂ O ₃	3.2 ± 0.3	1.9 ± 0.3	6.36 ± 0.15	4.4 ± 0.4

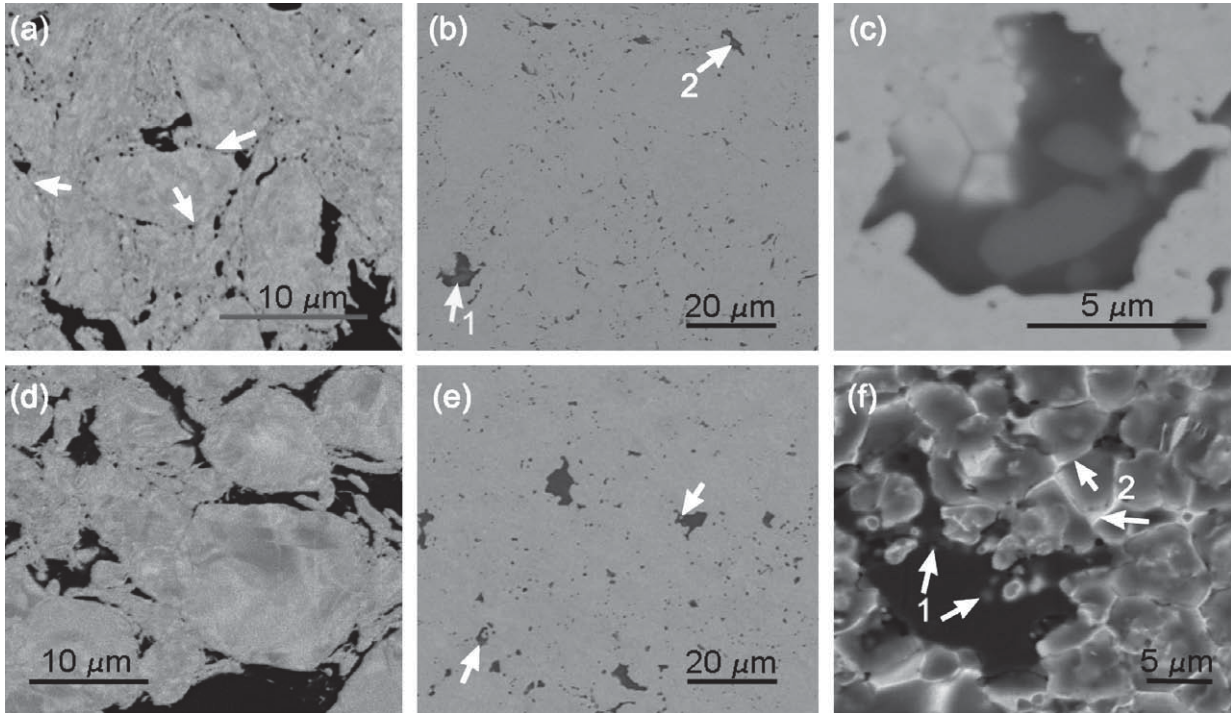


Fig. 6. BSE-SEM images of (a – c) W-2Ti-0.47Y₂O₃, and (d – f) W-4Ti-0.50Y₂O₃; (a) and (d) after HIP at 1550 K, (b), (c), (e) and (f) after HIP at 1973 K. (a) Arrows indicate the Ti-rich particle boundaries (black contrast). In figures (b), (e) and (f), the arrows indicate new W-Ti-O phases formed after the second HIP treatment (medium grey level contrast).

oxides. The composition and distribution of these oxides appeared to be similar in both materials. Fig. 6c shows a previous Ti pool (black contrast) with several inclusions of the new oxide phase (dark grey contrast) formed after HIP at 1973 K. Similar oxide particles were also found inside the previous Ti pools in W 4Ti 0.50Y₂O₃, as those indicated by white arrows in Fig. 6f.

The β Ti(W) grains in these alloys exhibited the same behaviour than the one observed for W 4Ti, i.e. the second HIP at 1973 K produced a reduction of the average grain size D_{50} , see Fig. 5 and Table 1. The grain size distribution of the W particles was also obtained for both alloys. The average grain sizes D_{50} resulted in 4.6 and 3.3 μm for W 2Ti 0.47Y₂O₃ and W 4Ti 0.50Y₂O₃, respectively.

The TEM analyses performed on W 4Ti 0.50Y₂O₃ samples HIP treated at 1973 K also revealed the presence of oxide particles between the W grains, which sizes ranged between 0.5 and 1 μm . EDX and electron diffraction analyses identified them as complex Ti Y oxides with different compositions and crystalline structure.

3.4. Microhardness measurements

Table 1 shows the effect of Ti and Y₂O₃ addition on the microhardness. 4 wt.% Ti addition to W containing Y₂O₃ produced microh

ardness increments of $\sim 120\%$ and 80% in the materials HIP treated at 1550 K and 1973 K, respectively. Y₂O₃ addition to W reduced the microhardness. The second HIP lowered the microhardness except for W 4Ti. This suggests that the formation of the Ti Y oxide phases softened the alloys.

4. Conclusions

The HIP treatment at 1973 K applied to W previously consolidated by HIP at 1550 K did not produce further densification because of the presence of open porosity in the material. However, addition of 0.5 wt.% Y₂O₃ yielded a material with very low porosity after the second HIP stage at 1973 K. In this case, complex (W Y) oxides were found filling the residual pores.

Addition of 2 or 4 wt.% Ti to W resulted in a fully dense material irrespective of the Y₂O₃ presence in the materials HIP treated at 1550 K. The present results demonstrate that sound W Ti/Y₂O₃ materials for full mechanical testing can be fabricated by HIP.

The validation of these alloys as PFC materials requires investigating their mechanical characteristics and behaviour under irradiation and plasma exposure. The mechanical characterization of these alloys is reported in Ref. [5].

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