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Published online 31 July 2019 | https://doi.org/10.1007/s40843-019-9469-4 Sci China Mater 2019, 62(12): 1898–1909



# Dense and pure high-entropy metal diboride ceramics sintered from self-synthesized powders *via* boro/ carbothermal reduction approach

Junfeng Gu<sup>1</sup>, Ji Zou<sup>1,2\*</sup>, Shi-Kuan Sun<sup>3</sup>, Hao Wang<sup>1</sup>, Su-Yang Yu<sup>2</sup>, Jinyong Zhang<sup>1</sup>, Weimin Wang<sup>1</sup> and Zhengyi Fu<sup>1\*</sup>

Equimolar quinary diboride powders, with ABSTRACT nominal composition of (Ti<sub>0.2</sub>Hf<sub>0.2</sub>Zr<sub>0.2</sub>Nb<sub>0.2</sub>Ta<sub>0.2</sub>)B<sub>2</sub>, were synthesized by boro/carbothermal reduction (BCTR) of oxide mixtures (MO<sub>2</sub>, M = Ti, Hf, Zr, Nb and Ta) using  $B_4C$  as source of B and C in vacuum. By adjusting the B<sub>4</sub>C/MO<sub>x</sub> ratios, diboride mixtures without detectable MO<sub>x</sub> were obtained at 1600°C, while high-entropy diboride (HEB) powders with particle size of < 1 µm was obtained at 1800°C. The phase, morphology and solid solution evolution process of the HEB powders during the BCTR process were comprehensively investigated. Although X-ray diffraction pattern indicated the powders synthesized at 1800°C were in a single-phase AlB<sub>2</sub> structure, elemental mappings showed that (Ta, Ti)-rich and (Zr, Nb)-rich solid solution coexisted in the HEB powders. The distribution of niobium and zirconium atoms in HEB was unable to reach uniform until the HEB powders were spark plasma sintered at 2000°C. (Ti<sub>0.2</sub>Hf<sub>0.2</sub>Zr<sub>0.2</sub>Nb<sub>0.2</sub>Ta<sub>0.2</sub>)B<sub>2</sub> ceramics with a relative density of 97.9% were obtained after spark plasma sintering the HEB powders at 2050°C under 50 MPa. Rapid grain growth was found in this composition when the sintering temperature was increased from 2000 to 2050°C, and the averaged grain size increased from 6.67 to 41.2 µm. HEB ceramics sintered at 2000°C had a Vickers hardness of 22.44  $\pm$ 0.56 GPa (under a load of 1 kg), a Young's modulus of ~500 GPa and a fracture toughness of 2.83  $\pm$  0.15 MPa m<sup>1/2</sup>. This is the first report for obtaining high density HEB ceramics without residual oxide phase, benefiting from the high quality HEB powders obtained.

Keywords: ultra-high temperature ceramics, high-entropy borides, solid solution, rapid grain growth, powder synthesis

#### INTRODUCTION

High-entropy alloys (HEAs) have attracted much research interest in recent years. Although the effects of "entropy stabilized" are still controversial, some HEAs exhibit superior mechanical and physical properties [1-3]. HEAs refer to the solid solutions that are formed by mixing equal or relatively large proportions of five or more metallic elements [1-4]. The concept of highentropy ceramics (HECs) was extended from HEAs. Up to now, studies on the preparation and characterization of HECs including oxides [5-14], carbides [15-21], borides [22-26], nitrides [27-29], and silicides [30] have been extensively carried out. Analogous to HEAs, HECs also show some unique properties including high hardness [15,16,20,22] and excellent thermo-mechanical properties [16], implying their promising applications in various areas including cutting tool [16], thermal protecting coating [24,28], etc.

High-entropy metal diborides (HEBs) were the firstly proposed and synthesized high-entropy non-oxides, which have greatly broadened the scope of high-entropy materials and the family of ultra-high temperature ceramics (UHTCs) [22]. These transition metal diborides exhibit a unique layered hexagonal crystal structure with alternating rigid two-dimensional (2D) boron nets and high-entropy 2D layers of metal cations [22]. Up to now, a series of HEBs including  $(Ti_{0.2}Hf_{0.2}Zr_{0.2}Nb_{0.2}Ta_{0.2})B_2$ ,  $(Hf_{0.2}Zr_{0.2}Ta_{0.2}Mo_{0.2}Ti_{0.2})B_2$ ,  $(Hf_{0.2}Zr_{0.2}Ta_{0.2}Nb_{0.2}Ti_{0.2})B_2$ ,  $(Hf_{0.2}Zr_{0.$ 

<sup>&</sup>lt;sup>1</sup> State Key Laboratory of Advanced Technology for Materials Synthesis and Processing, Wuhan University of Technology, Wuhan 430070, China

<sup>&</sup>lt;sup>2</sup> School of Metallurgy and Materials, University of Birmingham, Birmingham B15 2TT, United Kingdom

<sup>&</sup>lt;sup>3</sup> Department of Materials Science and Engineering, University of Sheffield, Sheffield S1 3JD, United Kingdom

<sup>\*</sup> Corresponding authors (emails: j.zou@bham.ac.uk or zouji1983@aliyun.com (Zou J); zyfu@whut.edu.cn (Fu Z))

performance of five individual counterparts [22]. HEB also offers a large space to tune the properties of diborides through controlling the composition of the solid solutions. In addition, reports confirmed that some unstable diborides like MoB<sub>2</sub>, could be stabilized at an AlB<sub>2</sub> structure in HEB [22,25]. This kind of cations, if segregated at grain boundary, might improve the mechanical property and oxidation resistance of UHTCs [31].

However, similar to other transitional metal diborides, the fabrication of dense HEB bulks still faces a challenge due to the impurities in starting powders and the strong covalent bonding therein. For example, all the aforementioned HEB ceramics densified from single phase diborides reached about ~92% of final relative density, although the starting powders had been ground to a fine level through high energy ball-milling [22]. Tallarita *et al.* [25] proposed a new processing route to densify HEBs that combined self-propagating high-temperature synthesis (SHS) with spark plasma sintering (SPS). Although the processing time had been shortened since ball-milling process was no more needed, the final relative density of HEBs did not exceed 92.5%.

The presence of porosity and impurities might make us greatly underestimate/misunderstand some performances of HEBs. High quality starting powder is the prerequisite to obtain HEBs with high purity and density. Boro/carbothermal reductions (BCTR) have been widely used for the synthesis of transition metal diboride powders [32–36]. However, this approach has not been well explored to synthesize HEB powders. Recently, Zhang *et al.* [26] showed that HEBs with relative density above 95% including (Hf<sub>0.2</sub>Zr<sub>0.2</sub>Ta<sub>0.2</sub>Cr<sub>0.2</sub>Ti<sub>0.2</sub>)B<sub>2</sub>, (Hf<sub>0.2</sub>Zr<sub>0.2</sub>Mo<sub>0.2</sub>Nb<sub>0.2</sub>Ti<sub>0.2</sub>)B<sub>2</sub> and (Mo<sub>0.2</sub>Zr<sub>0.2</sub>Ta<sub>0.2</sub>Nb<sub>0.2</sub>Ti<sub>0.2</sub>)B<sub>2</sub> could be sintered from HEB powders that were synthesized *via* borothermal reduction. Nevertheless, all the HEB ceramics from reference [26] show the presence of oxide impurities up to 7.2 wt%.

Therefore, the present work has two purposes: (1) to develop a feasible BCTR route for fine HEB powders to minimize the oxide levels and (2) to investigate the solid solution formation and densification behavior of the assynthesized powders to achieve dense HEBs with a finer microstructure.

#### **EXPERIMENTAL SECTION**

Commercially available TiO<sub>2</sub> (99.9% purity,  $D_{50} = 50$  nm, Aladdin, China), ZrO<sub>2</sub> (99.9% purity,  $D_{50} = 50$  nm, CSG Holding, China), HfO<sub>2</sub> (99.9% purity, average gain size 300 nm, Founde Star, China), Nb<sub>2</sub>O<sub>5</sub> (99.9% purity, Aladdin, China), Ta<sub>2</sub>O<sub>5</sub> (99.9% purity, Aladdin, China), and B<sub>4</sub>C (97% purity, average gain size 2.36 µm, Mudanjiang Diamond Boron Carbide, China) powders were used as raw materials. It should be noted that the reactions between B<sub>4</sub>C and oxides at higher temperature in vacuum can generate both B<sub>2</sub>O<sub>3</sub> and BO. If only B<sub>2</sub>O<sub>3</sub> is formed during the BCTR process, the process should proceed according to Reaction (1); vice versa, if BO is the only gaseous product, Reaction (2) should take place.

$$2\text{TiO}_{2}+2\text{ZrO}_{2}+2\text{HfO}_{2}+\text{Nb}_{2}\text{O}_{5}+\text{Ta}_{2}\text{O}_{5}+52/7\text{B}_{4}\text{C}$$

$$\rightarrow 10\text{Ti}_{0.2}\text{Zr}_{0.2}\text{Hf}_{0.2}\text{Nb}_{0.2}\text{Ta}_{0.2}\text{B}_{2}+34/7\text{B}_{2}\text{O}_{3}(\text{g})$$

$$+52/7\text{CO}(\text{g}), \qquad (1)$$

$$2 \text{Ti}O_2 + 22 \text{r}O_2 + 2\text{H}O_2 + \text{Nb}_2O_5 + \text{Ta}_2O_5 + 42/5B_4C$$
  

$$\rightarrow 10 \text{Ti}_{0.2} \text{Zr}_{0.2} \text{Hf}_{0.2} \text{Nb}_{0.2} \text{Ta}_{0.2} \text{B}_2 + 68/5\text{BO}(\text{g})$$
  

$$+42/5\text{CO}(\text{g}). \qquad (2)$$

In order to minimize both the residual  $B_4C$  and  $MO_x$  in the final powders, the  $B_4C/MO_x$  ratio should be set in the range regulated by Reactions (1) and (2). Accordingly, four compositions with different  $B_4C/MO_x$  ratios denoted as HEB1, HEB2, HEB3 and HEB4 were designed. Corresponding molar ratios of TiO<sub>2</sub>/ZrO<sub>2</sub>/HfO<sub>2</sub>/Nb<sub>2</sub>O<sub>5</sub>/ Ta<sub>2</sub>O<sub>5</sub>/B<sub>4</sub>C are 2/2/2/1/1/7.43 (all B<sub>2</sub>O<sub>3</sub>, Reaction (1)), 2/2/2/1/1/7.8, 2/2/2/1/1/8.17, and 2/2/2/1/1/8.4 (all BO, Reaction (2)). The designations for the different specimens are listed in Table 1.

The mixtures (each for 100 g) were blended in a ballmiller for 12 h, using ethanol and  $ZrO_2$  balls with a diameter of 5 mm as the mixing media. The weight loss of  $ZrO_2$  balls after mixing for 12 h was less than 0.1 g. After mixing, rotary evaporation was used to remove the excess solvent from the mixture. Then, the mixtures were

Table 1Sample designations

Composition	$TiO_2/ZrO_2/HfO_2/Nb_2O_5/Ta_2O_5/B_4C$	Excess B <sub>4</sub> C	1300°C	1400°C	1500°C	1600°C	1700°C	1800°C
HEB1 <sup>a</sup>	2/2/2/1/1/7.43	0 wt%	HEB1-13	HEB1-14	HEB1-15	HEB1-16	HEB1-17	HEB1-18
HEB2	2/2/2/1/1/7.8	5 wt%	HEB2-13	HEB2-14	HEB2-15	HEB2-16	HEB2-17	HEB2-18
HEB3	2/2/2/1/1/8.17	10 wt%	HEB3-13	HEB3-14	HEB3-15	HEB3-16	HEB3-17	HEB3-18
HEB4 <sup>b</sup>	2/2/2/1/1/8.4	13 wt%	HEB4-13	HEB4-14	HEB4-15	HEB4-16	HEB4-17	HEB4-18

a) According to Reaction (1); b) according to Reaction (2).

ground in an agate mortar by pestle and sieved through a 200-mesh screen. Field-assisted synthesis has been proven to be efficient for the fast synthesis of high purity UHTC powders [37,38]. In this work, a quantity of 20 g mixtures for each composition was then placed in a graphite crucible for reduction processing in a SPS system (ED-PAS-111, ELENX, Japan). The details for the setup could be found in our previous work [39,40]. This experiment was conducted in vacuum with a heating rate of 100°C/min to the designed temperature and held for 10 min. The initial vacuum reading was ~10 Pa. This number dropped when the reduction process started during heating, and the maximum vacuum level reached ~300 Pa during the powder synthesis. After holding at 1700°C or above, the vacuum level returned to ~10 Pa. The temperature was monitored by an optical pyrometer focusing on the external wall of the graphite crucible. Finally, 2.5 g of powders for each pellet was poured into a graphite die with inner diameter of 15 mm and densified using the same furnace: the sintering temperatures were 1900, 2000 and 2050°C, with a heating rate of 100°C/min and holding time of 5 min.

The phase assembly of the as-prepared samples was examined by X-ray diffraction (XRD, Empyrean, Panalytical, Netherlands). The microstructure was investigated by a scanning electron microscope (SEM, JEOL7000F, Tokyo, Japan) equipped with energy/wavelengthdispersive X-ray (EDS/WDS, Oxford, UK) analyzer system and a scanning transmission electron microscope (STEM, TalosF200S, FEI, USA). Final density of the bulks was measured using the Archimedes' method. Oxygen and carbon contents in the selected powder samples were measured by a nitrogen/oxygen determinator (TC600, Leco Corporation, USA) and a carbon/sulphur analyzer (CS-2000, Eltra Elemental Analyzers, Germany), respectively. Theoretical densities of HEBs were calculated according to the lattice parameters of HEB3, by fitting the peaks from the XRD results. XRD patterns were analyzed using Rietveld refinement method [41]. The program EXPGUI 2 was used to refine a structural model based on  $ZrB_2$  (in hexagonal space group *P6/mmm*) [42]. On basis of the calculated lattice parameter, the theoretical density was determined by the quotient between unit cell mass and unit cell volume. The thermodynamic calculations were performed via HSC Chemistry v. 7.1 (Outokumpu research Oy, Pori, Finland).

The grain size was measured from the SEM images. Each reported value was averaged from ~400 grains. The modulus and Vickers hardness of the as-sintered ceramics were determined by a mechanical resonance frequencies machine (GrindoSonic, Model MK7, Sweden) and a Vickers Hardness Tester (Durascan50, Struers, Austria), respectively. For the hardness test, hardness values were compared under different loads (0.2, 1 and 5 kg) at the same holding time of 15 s. The fracture toughness of HEB was calculated by direct crack measurements according to Anstis equation [43], based on the crack generated by the indents with a load of 5 kg. A typical indentation image for the crack measurement was shown in Fig. S1.

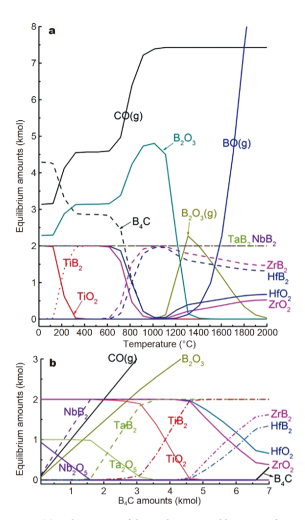
#### **RESULTS AND DISCUSSION**

#### Thermodynamic consideration

Though all the  $MO_x$  could be reduced by  $B_4C$  with increasing temperature, the favorable temperatures for the different BCTR reactions are different. Equilibrium amounts of phases in  $MO_x$ - $B_4C$  system were calculated as a function of temperature (Fig. 1a) under a system partial pressure of 10 Pa. The prediction shows that the reaction between  $Ta_2O_5$  and  $B_4C$  at P = 10 Pa could start even at room temperature, so does the reaction between  $Nb_2O_5$  and  $B_4C$ . The required temperature for the reduction of  $MO_x$  by  $B_4C$  increases in an order of  $TiO_2$ ,  $ZrO_2$  and  $HfO_2$ . The proceeding of the  $HfO_2$ - $B_4C$  reduction is the most difficult; a temperature over 1000°C is needed.

If the input amount of the raw materials is based on Reaction (1), the calculation also implies that certain amounts of  $ZrO_2$  and  $HfO_2$  gradually reappear in the system when temperature exceeds 1100°C. Interestingly, the increasing amounts of  $ZrO_2$  and  $HfO_2$  are associated with the occurrence of BO(g). At a sufficiently high temperature,  $B_2O_3(l, g)$  could react with  $B_4C$  to generate BO(g). When it happens, more  $B_4C$  will be consumed. The lack of  $B_4C$  is the reason for the occurrence of  $ZrO_2$  and  $HfO_2$  in Fig. 1a when temperature is over 1100°C. It is the theoretical basis for adding excess  $B_4C$  in the starting powder mixtures, in order to obtain HEB powders free of residual oxides, as will be described later.

The reaction order in  $MO_x$ -B<sub>4</sub>C system at 1600°C can be seen from Fig. 1b. As shown in the plots, among  $MO_{x^3}$  $Nb_2O_5$  is the easiest compound which could be reduced by B<sub>4</sub>C, followed by Ta<sub>2</sub>O<sub>5</sub>, TiO<sub>2</sub>, ZrO<sub>2</sub> and HfO<sub>2</sub>. It should be noted that no solid solutions of various oxides or diborides were considered in these calculations (Fig. 1), due to the lack of thermodynamic database. Solid solutions normally have lower Gibbs' free energies compared with the end members; therefore, more complicated oxide or boride solid solutions may co-exist in  $MO_x$ -B<sub>4</sub>C system, which fail to be taken into account from the current calculations.



**Figure 1** (a) Molar content of the products in equilibrium as a function of the temperature from room temperature (RT) to 2000°C at a partial pressure of 10 Pa. The input amounts of the reactants are based on Reaction (1). (b) The effects of  $B_4C$  amount on the progress of reactions in  $MO_x$ - $B_4C$  system at 1600°C.

#### **Reaction process**

Fig. 2a shows the XRD patterns for the powders synthesized at 1800°C. Peaks for residual oxides were found in HEB1-18 and HEB2-18, but disappeared when more  $B_4C$ were added in the starting mixtures (HEB3-18 and HEB4-18), indicating the reactions proceeded according to both Reactions (1) and (2). The optimized excess amount of  $B_4C$  to completely remove  $MO_x$  based on Reaction (1) should be in a range from 5 to 10 wt%. No apparent peak shift was detected among HEB1–HEB4, as seen from an enlarged range in Fig. 2b, indicating the compositions of the high-entropy diboride phase (AlB<sub>2</sub> structure with a space group of *P6/mmm*, solid solution 1, SS1) were similar when different  $B_4C/MO_x$  ratios were adopted. However, the peak assigned to the (101) plane in Fig. 2b reveals the presence of trace unsolved boride (SS2) in HEB1-18 and HEB2-18. Based on the XRD patterns of HEB3-18, lattice parameters of  $(Ti_{0.2}Zr_{0.2}Hf_{0.2}Nb_{0.2}Ta_{0.2})B_2$  were refined as  $a=3.1062 \pm 0.0013$  Å and  $c=3.3755 \pm 0.0021$  Å, slightly higher than those in Gild's work (a=3.101 Å, c=3.361 Å), but lower than the first-principle predicted values (a=3.116 Å, c=3.391 Å) [22,23]. According to the lattice parameters of HEB3-18, theoretical density of  $(Hf_{0.2}Zr_{0.2}Ta_{0.2}Nb_{0.2}Ti_{0.2})B_2$  was calculated to be 8.237 g cm<sup>-3</sup>, assuming the lattice is fully occupied.

The phase evolution of HEB3 at different temperatures is displayed in Fig. 2c, d. Clearly, with the increase of temperature, oxides were gradually converted into diborides, and disappeared completely at 1500°C. Interestingly, peaks assigned to TiO<sub>2</sub>, Nb<sub>2</sub>O<sub>5</sub> and Ta<sub>2</sub>O<sub>5</sub> were not detectable in the mixtures at 1300°C, suggesting that the boro/carbothermal reductions of TiO2, Nb2O5 and Ta<sub>2</sub>O<sub>5</sub> took place earlier compared with those of ZrO<sub>2</sub> and HfO<sub>2</sub>. Such observation well fits the thermodynamic prediction shown in Fig. 1. In fact, Ta<sub>2</sub>O<sub>5</sub> can react with B<sub>4</sub>C and completely convert into TaB<sub>2</sub> at 1100°C, while corresponding temperature for the conversion of ZrO<sub>2</sub> needs at least 1400°C [34,36]. The difference of favorable temperatures between thermodynamic prediction and experimental observation suggests the kinetics play a major role in the advancement of BCTR in MO<sub>x</sub>-B<sub>4</sub>C systems.

The contents of residual oxygen and carbon in the assynthesized HEB3 powders are 7.52 and 0.18 wt% in HEB3-16, respectively, as listed in Table 2. The low carbon content suggested most of the B4C in the powder mixture had been reacted with MO<sub>x</sub> at 1600°C. Considering no oxide peaks were found in HEB3-16 (Fig. 2b), the high level of oxygen content in this sample indicated that significant amount of B<sub>2</sub>O<sub>3</sub> was still left in the powders after the vacuum reduction at 1600°C. A sharp drop of the oxygen content from 7.52 to 0.64 wt% was found with the increase of synthesis temperature to 1700°C, implying most of the B<sub>2</sub>O<sub>3</sub> had been removed out of the boride powders at this stage. It should be noted that the HEB3-17 powders synthesized in this work shows substantially lower oxygen/carbon content, compared with the other HEB powders prepared via various routes [35,36].

#### Morphology evolution of HEB powders during BCTR

The morphology of HEB3-13 powders is shown in Fig. 3a. Clearly, there are still unreacted  $B_4C$  and oxide particles as arrowed in HEB3-13. As the temperature increased, the boride particles gradually transformed into regular shapes

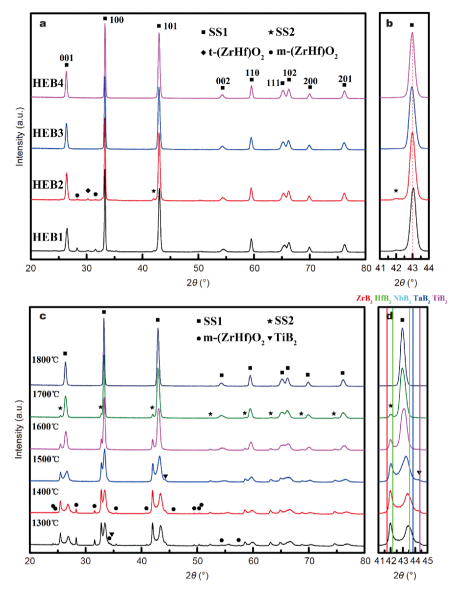


Figure 2 XRD patterns of (a, b) HEB powders synthesized at 1800°C and (c, d) HEB3 powders synthesized at 1300-1800°C.

Table 2	Oxygen and	carbon	contents in	ı the	selected	HEB	nowders
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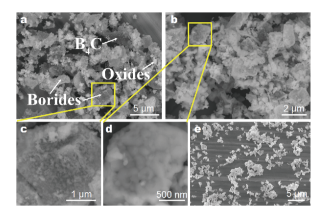
Composition	Oxygen content (wt%)	Carbon content (wt%)
HEB3-16	7.52	0.18
HEB3-17	0.64	0.041
HEB3-18	0.52	0.032

and their sizes became coarser (Fig. S2a–d). When the synthesis temperature reached 1600°C (Fig. 3b), the contrast of backscattered electron signal from the powders became uniform, revealing most of the oxides and  $B_4C$  had been consumed. Interestingly, the newly formed

ultra-fine boride powders in Fig. 3a were agglomerated in a shape that resembled unreacted  $B_4C$  particles. A higher magnification image of HEB3-13 in Fig. 3c indicated the boride particles were embedded in the unreacted  $B_4C$ matrix with a darker contrast. Such phenomenon implied that the surfaces of  $B_4C$  particles were the sites where the reduction reactions took place, and boride powders were generated through the reactions between the  $B_4C$  particles and oxide particles attached on them. Inversely, if the diffusion of boron atoms into oxides is the controlling step of BCTR, grains should be observed in the shape resembling original oxides rather than  $B_4C$ . Similar features in Fig. 3c can be seen in the powders synthesized at

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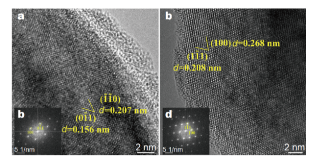
**Figure 3** Back scattered electron (BSE) image of the (a) HEB3-13 and (b) HEB3-16 powders, showing the presence of unreacted oxides, unreacted  $B_4C$  and newly formed ultra-fine diboride particles as-arrowed. (c, d) Higher magnification images of the ultra-fine diboride particles in (a) and (b). (e) SEM image of the HEB3-18 powders.

1600°C (Fig. 3d), i.e., some diborides agglomerated into a shape of raw  $B_4C$  particles. However, no region with darker contrast could be recognized in HEB3-16, indicating that residual  $B_4C$  was completely consumed at 1600°C.

The morphology of the HEB powders synthesized with various amounts of B4C additions at 1800°C is presented in Fig. 3e and Fig. S2e-g. Slight difference in morphology can be seen from these images, i.e., necking is prone to take place in the powders synthesized at higher temperature and/or with more B<sub>4</sub>C additions. The average particle size of HEB1-18, HEB2-18, HEB3-18, and HEB4-18 powders is  $0.35 \pm 0.07$ ,  $0.38 \pm 0.10$ ,  $0.44 \pm 0.12$ , and  $0.63 \pm 0.15 \,\mu\text{m}$ , respectively. Fig. 4 presents the high resolution TEM (HRTEM) images of the HEB3-13 and HEB3-18 powders, both of which are well crystalline. The interplanar distances of HEB3-13 match well with those of (110) and (011) planes of SS1 (0.208 and 0.154 nm, from XRD patterns). The interplanar distances of HEB3-18 match well with those of (100) and  $(1\overline{1}1)$  planes (0.269)and 0.210 nm, from XRD patterns) in HEB with AlB<sub>2</sub> structure.

#### The formation of solid solution during BCTR

Although five different oxides were used, only three different diboride phases (SS1, SS2, TiB<sub>2</sub>, as seen in Fig. 2c) were detected from the XRD pattern of the powders synthesized at 1300°C. The relative peak intensity of SS2 and TiB<sub>2</sub> gradually decreased with temperature, suggesting that SS1, SS2 and TiB<sub>2</sub> diffused into each other to form a single-phase solid solution and such an entropy increase process was activated by temperature. According



**Figure 4** HRTEM images of HEB3 powders synthesized at (a) 1300 and (c) 1800°C. The fast Fourier transform images of (a) and (c) were inserted as (b) and (d).

to the XRD pattern of HEB3-13 (Fig. 2c), the lattice parameters of SS2 are a=3.1504 Å, c=3.4958 Å, slightly smaller than those of the as-reported  $Zr_{0.5}Hf_{0.5}B_2$  (a= 3.1542 Å, c=3.4954 Å [44] and a=3.155 Å, c=3.497 Å [45]). An enlarged region of the peaks assigned to (101) plane in Fig. 2d reveals that the peak shift of SS2 is very limited for increasing processing temperature. On the contrary, SS1 shifted significantly towards lower angles, indicating that the composition of SS2 kept almost unchanged during its diffusion into SS1. It is worth noting that intensity of (Zr,Hf)O2 peaks also decreases in this temperature range. If there are other major metallic elements apart from Hf and Zr in SS2, the incorporation of more HfB<sub>2</sub> and ZrB<sub>2</sub> should cause a further peak shift of SS2 between 1300 and 1500°C. Accordingly, SS2 must be rich in HfB<sub>2</sub> and ZrB<sub>2</sub>, in accordance with the evolution of the XRD patterns.

To further understand the solid solution process, STEM-mappings of HEB3-13 and HEB3-18 were chosen to evaluate the compositional uniformity in the powders after BCTR. However, the scale of HAADF-STEM mapping is not large enough to fully recognize the phase separation verified by XRD. The mappings on the different areas from HEB3-13 and HEB3-18 show the signals of five transition metal elements (Fig. 5). This indicates the formation of quinary solid solution (SS1) started at as low as 1300°C; however, the distribution of these elements in the solid solution seems to be inhomogeneous. For instance, several regions rich in Ti were found in HEB3-13 (Fig. 5a), implying the as-indexed TiB<sub>2</sub> in Fig. 2c actually was a solid solution but concentrated on Ti. As the temperature increased to 1800°C, the distribution of Nb was slightly different from other elements which had diffused uniformly (Fig. 5b). However, based on the XRD pattern (Fig. 2a), a single phase HEB has been indexed at this temperature.

The above assumption on the formation of solid solu-

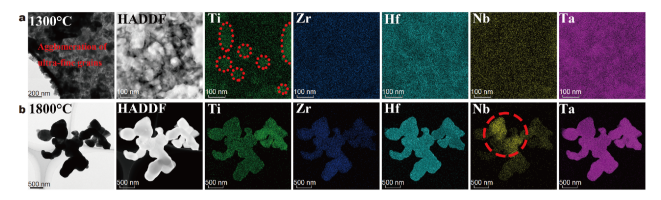


Figure 5 (a) TEM image of the HEB3-13 powders shows an agglomeration of newly formed ultra-fine diborides grains. STEM-mappings of one of the agglomerate regions indicate the quinary solid solution reaction had occurred at 1300°C. (b) TEM image and STEM-mappings of the HEB3-18 powders reveal the unevenly atomic distribution within grains. Note that some circled regions have different concentrations of Nb.

tion was further examined by SEM and element mappings. In Fig. 6a, regions with different features/contrasts are identified on HEB3-13. Among them, some spherical particles (marked 3 in Fig. 6a, ~0.5  $\mu$ m) are randomly distributed in the pellet, which are mainly composed of Zr, Hf and O, as indicated by the EDS patterns (Fig. 6d). Hence, these particles are likely assumed to be (Zr,Hf)O<sub>2</sub>, as previously indexed from the XRD patterns. The distribution of elements in the other regions of HEB3-13 is also inhomogeneous, and two typical regions are labelled as "1" and "2" in Fig. 6b and c, respectively. EDS spectra confirm the presence of all five elements in these areas; however, region "1" is richer in Ti and Ta while Hf, Zr and Nb are the dominated metallic elements in region "2". Combined with the XRD pattern (Fig. 2c), region "1" and "2" are likely assumed to SS1 and SS2, due to the differences in lattice parameters, i.e., the lattice parameters of  $ZrB_2$  and  $HfB_2$  are larger than those of  $TiB_2$ , NbB<sub>2</sub> and TaB<sub>2</sub>.

If it were correct, the extents of areas rich in Hf, Zr and Nb would be obviously smaller, compared with that in TiTa-rich region in HEB3-16, due to the lower intensity of SS2 in HEB3-16 (Fig. 2c). However, it was not the case. In HEB3-16 (Fig. 7a), two regions still could be easily recognized, and the area rich in Zr and Nb is of the similar extent as that rich in Ta and Nb, although the distribution of Hf seems to be homogenous. The deconvolution of the XRD peaks for HEB3-16 also indicates

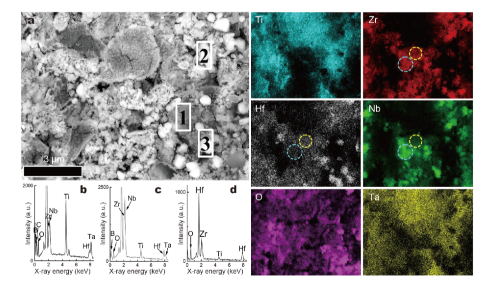


Figure 6 (a) BSE image of the HEB3-13 powders and corresponding EDS-mappings, (b–d) point EDS results showing the presence of (Ti, Ta)-rich regions, (Zr,Hf)-rich regions and unreacted (Zr,Hf) $O_2$ . The circled regions in the Nb, Zr and Hf mappings revealing the ZrNb-rich smaller particles are superimposed on the ZrHfNb-rich area.

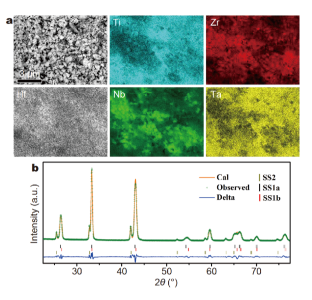


Figure 7 (a) BSE image and corresponding EDS-mappings of HEB3-16, and (b) Rietveld refinement of the XRD patterns of HEB3-16 using MAUD software (Rwp = 9.29%).

that SS1 is constituted of two parts (Fig. 7b). By further zooming the region "2" in HEB3-13, it could be found that smaller particles with stronger signals of Zr and Nb were superimposed on the ZrHfNb-rich area, as circled in the Zr, Hf and Nb mappings in Fig. 6. Therefore, apart from TiB<sub>2</sub>-rich area, there are three other boride solution phases in HEB3-13 powders: (1) (Ti, Ta)-rich regions, (2) (Zr, Hf, Nb)-rich regions and (3) (Zr, Nb)-rich regions. Note that the locations of (2) and (3) are normally overlapping at 1300°C and (2) is decreasing in size with temperature increase.

According to the Vegard's law, the lattice parameters of solid solution should locate in the area among their single component counterparts. Considering the lattice parameters of TiB<sub>2</sub> and TaB<sub>2</sub> are smaller than others, the right part (SS1b) of SS1 should be assigned to the (Ti, Ta)-rich regions. The (Zr, Nb, Hf)-rich regions which contain more Hf should have the largest lattice parameters, which are supposed to be SS2. This is also in favor of the fact that SS2 is rich in Zr and Hf according to XRD patterns. On the contrary, the lattice parameters of (Zr, Nb)-rich regions position between the above two, which correspond to the left part of SS1 (SS1a). With the temperature increasing, the distribution of Hf became homogenous and not concentrated on specific particles (as seen in Figs 6, 7). This indicated the Hf in (Zr, Nb, Hf)-rich solid solution gradually diffused into the matrix, which was in well accordance with the weakened signal of SS2 as observed in Fig. 2d.

# The sinterability of self-synthesized (Ti<sub>0.2</sub>Hf<sub>0.2</sub>Zr<sub>0.2</sub>Nb<sub>0.2</sub>Ta<sub>0.2</sub>)B<sub>2</sub> powders

In order to evaluate the sinterability of the as-synthesized powders, HEB3-17 powders were selected to densify HEBs, due to the relatively finer particle size and better composition homogeneity in this batch. Relative density of the HEBs sintered at 1900, 2000, and 2050°C is 85.8%, 94.4%, and 97.9%, respectively. The HEB powders obtained from BCTR route in this work exhibited better sinterability than those prepared from the existing routes (e.g., high energy ball milling [22] and self-propagating high-temperature synthesis [25]). Fig. 8 shows the polished surface of these HEBs sintered at different temperatures. Pores with a black contrast and size of ~1 µm are visible on the sample sintered at 1900°C, and are mainly located at grain boundaries. Both XRD patterns (Fig. 9) and oxygen maps (Fig. S3) indicate that all these ceramics have formed into a single phase, without notable residual oxides. However, the oxides have been recently observed in the HEBs sintered from the powders synthesized *via* borothermal reduction [26].

Although a single HEB phase has formed, apart from the pores, areas with two contrasts could be clearly distinguished from the BSE image: the darker one is rich in Nb and Zr (Fig. 10a) while the elemental distribution in the lighter area seems to be homogenous. Similar phase separation was previously noticed in the HEB powders (Fig. 7). Again, it supported the conclusion that SS1 peak was contributed by both (Ti, Ta) and (Nb, Zr)-rich solid solution.

When the sintering temperature further rose to 2000 and 2050°C, the porosity was evidently reduced and the distribution of Nb and Zr became homogenous, i.e., the final pore removal occurred simultaneously with the further diffusion of Nb and Zr atom from its rich to depleted regions in (Ti<sub>0.2</sub>Zr<sub>0.2</sub>Hf<sub>0.2</sub>Nb<sub>0.2</sub>Ta<sub>0.2</sub>)B<sub>2</sub> (Fig. 10b, c). Surprisingly, a sudden grain growth occurred when the sintering temperature was elevated from 2000 to 2050°C. The average grain sizes of HEB3 ceramics are  $0.71 \pm 0.14 \,\mu m$  (1900°C),  $6.67 \pm 1.20 \,\mu m$  (2000°C) and 41.2  $\pm$  8.1 µm (2050°C), respectively. Such rapid grain growth from 2000 to 2050°C is contradictory with the existing knowledge that high configurational entropy lowers the grain boundary energy, thus lowering the driving force for grain growth in high entropy materials [46,47]. It has not been observed in diboride ceramics. The mechanism for such a rapid grain growth of HEBs is still unclear. One possible reason is that entropy alone might not be sufficient to stabilize the (Ti<sub>0.2</sub>Zr<sub>0.2</sub>Hf<sub>0.2</sub>-Nb<sub>0.2</sub>Ta<sub>0.2</sub>)B<sub>2</sub> boride solid solution phase, and the en-

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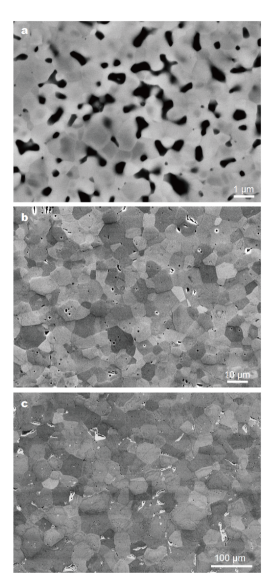


Figure 8 Polished surfaces of HEB3 ceramics sintered at (a) 1900°C, (b) 2000°C, and (c) 2050°C.

thalpy of mixing in this system has to be evaluated to fully understand whether HEB is more stable compared with other single-phase counterparts.

The typical mechanical properties of the as-sintered HEB ceramics are summarized in Table 3. Young's modulus (*E*) of the HEB3 sintered at 2050°C (527 GPa) is slightly higher than that sintered at 2000°C (500 GPa), probably due to the less porosity in the former sample. The Vickers hardness of HEB3 ceramics sintered at 2000°C exhibits a typical indentation load dependence (ISE) behavior, i.e., the as-measured hardness values decrease with the increasing indentation load. ISE has been commonly observed in many structural ceramics [48].



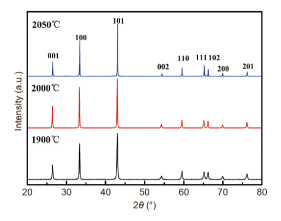


Figure 9 XRD patterns of the three ceramics.

For example, under loads of 0.2, 1, and 5 kg, the hardness values for HEB3 are  $25.61 \pm 0.83$ ,  $22.44 \pm 0.56$  and  $19.44 \pm 0.50$  GPa, respectively. These values are obviously higher than the results of previous reports on HEB ceramics with the same composition, e.g., 21.7 GPa under 0.2 kg [49] and 17.5 GPa under ~1 kg [22]. Such differences may be attributed to the high-purity and dense bulks achieved in this work. Apparently, any residual oxide and pores observed in references [22,49] will lower the hardness for HEB ceramics.

The hardness of each single phase diborides under the same indentation load is also compared in Table 3 [50–54]. The hardness of HEB ceramics (22.44 GPa under 1 kg load) is comparable to the averaged value (23.4 GPa, under 1 kg load) of the five. Considering the residual porosity left, higher intrinsic hardness value could be expected in HEB. Unfortunately, the fracture toughness of HEB shows no superiority or even worse than the other single-phase counterparts, possibly due to the lack of toughening mechanisms in this sample, as indicted by the cracks which propagated straightly in Fig. S1.

#### CONCLUSION

This work demonstrates that conventional boro/carbothermal reduction could be applied for the synthesis of HEB powders.  $(Ti_{0.2}Hf_{0.2}Zr_{0.2}Nb_{0.2}Ta_{0.2})B_2$  powders were successfully synthesized by  $B_4C$  reduction of corresponding oxide mixtures. As-synthesized powders showed fine particle size, high purity and good sinterability. The following conclusions can be drawn regarding this study on the powder preparation, densification and mechanical properties of HEB ceramics:

(1) To minimize the residual oxide in the HEB powders, 10 wt% excess  $B_4C$  according to Reaction (1) has to be added in the starting powder mixtures, as the forma-

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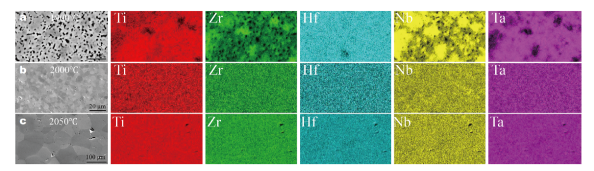


Figure 10 WDS-mappings of the specimens sintered at (a) 1900°C, (b) 2000°C, and (c) 2050°C, indicating the uneven elemental distribution at 1900°C while the elemental distributions at 2000 and 2050°C are homogenous.

Table 3 Properties of the high-entropy diboride ceramics

Composition	Sintering temperature (°C)	Relative density (%)	Average grain size (µm)	E (GPa)	$K_{\rm IC}$ (MPa m <sup>1/2</sup> )	Hardness (GPa)	Ref.
$(Ti_{0.2}Zr_{0.2}Nb_{0.2}Hf_{0.2}Ta_{0.2})B_2$	2000	94.4	6.67 ± 1.20	500	2.83 ± 0.15	25.61 ± 0.83 (HV0.2) 22.44 ± 0.56 (HV1)	This work
$(Ti_{0.2}Zr_{0.2}Nb_{0.2}Hf_{0.2}Ta_{0.2})B_2$	2050	97.9	$41.2\pm8.1$	527	/	26.82 ± 1.77 (HV0.2)	This work
$(Ti_{0.2}Zr_{0.2}Nb_{0.2}Hf_{0.2}Ta_{0.2})B_2$	2000	92.4	/	/	/	$17.5 \pm 1.2 (HV \sim 1)$	[22]
$(Ti_{0.2}Zr_{0.2}Nb_{0.2}Hf_{0.2}Ta_{0.2})B_2$	2000	96.3	1.59	/	$4.06 \pm 0.35$	$21.7 \pm 1.1$ (HV0.2)	[49]
$TiB_2$	1800	96.1	/	$497 \pm 15$	$4.64 \pm 0.45$	28.6-32.4 (HV1)	[50]
$ZrB_2$	2000	99.8	~6	489	$3.5\pm0.3$	$23 \pm 0.9$ (HV1)	[51]
NbB <sub>2</sub>	1900	97.7	/	539	$4.0~\pm~0.6$	$20.3 \pm 0.6 (HV0.5)^{a}$	[52]
$HfB_2$	1850	98.8	8	/	$3.5 \pm 0.5$	$18.1 \pm 0.9$ (HV1)	[53]
TaB <sub>2</sub>	1850	96	5.3	/	$5.61 \pm 0.17$	$25.1 \pm 0.5$ (HV1)	[54]

a) HV1 data is not available.

tion of BO(g) (Reaction 2) will consume more  $B_4C$ .

(2) All the oxides could be reduced by  $B_4C$  in mild vacuum at as low as 1500°C in this system. Study of the solid solution process showed that different solid solution, i.e., (i) TiB<sub>2</sub> dominated; (ii) (Ti, Ta)-rich; (iii) (Zr, Hf, Nb)-rich and (iv) (Zr, Nb)-rich, appeared during the BCTR process. Homogenization of them cannot be realized until the as-synthesized HEB powders were sintered at 2000°C or above.

(3) Powders synthesized at 1700°C (HEB3-17) exhibit relatively good sinterability. Single phase HEBs with high relative density of 94.4% were achieved at 2000°C/50 MPa /5 min using this powder. Corresponding (Ti<sub>0.2</sub>Hf<sub>0.2</sub>Zr<sub>0.2</sub>Nb<sub>0.2</sub>Ta<sub>0.2</sub>)B<sub>2</sub> ceramics exhibits a combination of good mechanical properties, except fracture toughness, including a hardness of 22.44 ± 0.56 GPa, a fracture toughness of 2.83 ± 0.15 MPa m<sup>1/2</sup> and a Young's modulus of 500 GPa.

(4) Rapid grain growth for HEBs was found on the samples sintered at 2050°C, suggesting the high configurational entropy might not be enough to stabilize the  $(Ti_{0.2}Zr_{0.2}Hf_{0.2}Nb_{0.2}Ta_{0.2})B_2$  ceramics at ultra-high temperature.

# Received 9 April 2019; accepted 1 July 2019; published online 31 July 2019

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Acknowledgements This work was financially supported by the National Natural Science Foundation of China (51521001 and 51832003), and the Fundamental Research Funds for the Central Universities.

**Author contributions** Zou J and Fu ZY designed the research. Gu J and Zou J performed the experiments with the help of Wang H, Wang W, Zhang J and Fu ZY. Sun SK performed the XRD refinement. Yu SY and Zou J performed the hardness test. Fu Z supervised and acquired funding for the research. Gu J and Zou J wrote the paper. All authors discussed the results and commented on the manuscript.

Conflict of interest These authors declare no conflict of interest.

**Supplementary information** Supporting data are available in the online version of this paper.



**Junfeng Gu** is currently a PhD student at the State Key Laboratory of Advanced Technology for Materials Synthesis and Processing, Wuhan University of Technology. His research focuses on the sintering mechanisms and properties of ultra-high temperature ceramics.



**Ji Zou** is a Research Fellow at the University of Birmingham, UK and an adjunct professor at Wuhan Institute of Technology, China. He received his PhD degree from Shanghai Institute of Ceramics, Chinese Academy of Sciences. He has been active in the processing-structure-property correlation of ceramics, especially for boride ceramics.



Zhengyi Fu is a chief professor at the State Key Laboratory of Advanced Technology for Materials Synthesis and Processing, Wuhan University of Technology. He received his PhD degree from Wuhan University of Technology in 1994. His current research interests include advanced sintering and bioprocess-inspired fabrication.

#### 硼热/碳热还原技术制备高致密度、高纯度高熵 硼化物陶瓷

顾俊峰<sup>1</sup>, 邹冀<sup>1,2\*</sup>, 孙世宽<sup>3</sup>, 王皓<sup>1</sup>, 于苏洋<sup>2</sup>, 张金咏<sup>1</sup>, 王为民<sup>1</sup>, 傅正义<sup>1\*</sup>

摘要 以过渡金属氧化物(MO<sub>x</sub>)和碳化硼(B<sub>4</sub>C)为原料,采用硼热/ 碳热还原技术在1800°C下制备得到了等摩尔比的亚微米级五元高 熵硼化物(Ti<sub>0.2</sub>Hf<sub>0.2</sub>Zr<sub>0.2</sub>Nb<sub>0.2</sub>Ta<sub>0.2</sub>)B<sub>2</sub>粉体,并深入探讨了硼热/碳热 还原过程中产物的物相、形貌以及固溶体变化过程.X射线衍射谱 表明所制备的粉体在1800°C即形成了单相结构,但直到2000°C样 品中各元素才分布均匀.将得到的粉体在2050°C/50 MPa的条件下 进行放电等离子烧结,获得了相对密度为97.9%的高熵硼化物陶瓷. 研究发现,当烧结温度从2000°C增加到2050°C时,高熵陶瓷晶粒迅 速长大,平均晶粒尺寸从6.67 μm增大到41.2 μm. 2000°C下制备得 到的高熵硼化物陶瓷具有良好的力学性能,其硬度、杨氏模量和 断裂韧性分别为22.44 ± 0.56 GPa,~500 GPa, 2.83 ± 0.15 MPa m<sup>1/2</sup>.