

WB_{5-x} : Synthesis, Properties, and Crystal Structure—New Insights into the Long-Debated Compound

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The recent theoretical prediction of a new compound, WB₅, has spurred the interest in tungsten borides and their possible implementation in industry. In this research, the experimental synthesis and structural description of a boron-rich tungsten boride and measurements of its mechanical properties are performed. The ab initio calculations of the structural energies corresponding to different local structures make it possible to formulate the rules determining the likely local motifs in the disordered versions of the WB₅ structure, all of which involve boron deficit. The generated disordered WB4 18 and WB4 86 models both perfectly match the experimental data, but the former is the most energetically preferable. The precise crystal structure, elastic constants, hardness, and fracture toughness of this phase are calculated, and these results agree with the experimental findings. Because of the compositional and structural similarity with predicted WB₅, this phase is denoted as WB_{5-v}. Previously incorrectly referred to as "WB₄," it is distinct from earlier theoretically suggested WB₄, a phase with a different crystal structure that has not yet been synthesized and is predicted to be thermodynamically stable at pressures above 1 GPa. Mild synthesis conditions (enabling a scalable synthesis) and excellent mechanical properties make WB_{5-x} a very promising material for drilling technology.

1. Introduction

Tungsten borides have attracted great attention from the scientific community, with phases like WB₂, WB₃, and hotly debated WB4 found to display fascinating mechanical properties. Wide regions of homogeneity of the W-B phases^[1-5] may be caused, at least partly, by extensive polysomatism,^[6] which leads to significant difficulties in the synthesis of single crystals with a well-defined structure and stoichiometry. This factor and, more importantly, difficulties in determining the exact positions of the boron atoms using the X-ray diffraction (XRD) also result in inaccurate crystallographic descriptions of the synthesized phases.

A long-standing debate has been the crystal structure of the highest tungsten boride phases. Observed for the first time in 1961 by Chretien and Helcorsky^[7] and determined as WB₄ with a tetragonal ThB₄ arrangement, several years later this phase was reported to have compositions WB₄,^[8] W_{2-x}B₉,^[9] WB₁₂,^[10] and

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 $W_{1-x}B_3$,^[11,12] and a hexagonal structure. Various experimental techniques were used to characterize the obtained samples.^[8,11,13]

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In 1966, Romans and Krug^[8] proposed a structural model of WB4 based on the X-ray diffraction and density measurements, suggesting a possible excess of boron. The model has the hexagonal symmetry P6, /mmc, with boron dimers and graphenelike boron sheets. A model by Nowotny et al.^[9] introduced the $W_{2-x}B_{0}$ composition with a fractional occupancy of the tungsten atoms, suggesting that the boron atoms form B₆ octahedra instead of the dimers and that there is only a half of the hexagonal honeycomb boron layers between the metal layers, compared to the model by Romans and Krug.^[8] A model proposed by Lundström and Rosenberg^[11] and further discussed by Zeiringer et al.^[12] suggested the $W_{1-r}B_3$ composition with a fractional occupation of the tungsten atoms, which are sandwiched between planar graphene-like boron layers (a MoB₃-type structure). In the experimental work by Gu et al.,^[14] the crystal structure of the synthesized higher tungsten boride has been analyzed using the model of WB4 by Romans and Krug.^[8] Cheng et al.^[3] have carried out a comprehensive theoretical investigation of the W–B phase diagram, predicting several metastable WB₄, and showed that the structural model of WB4 proposed and used by experimentalists^[8,14,15] is thermodynamically and dynamically unstable, and therefore this structure cannot exist in principle. In the theoretical study of the W-B system, Zhao et al.^[5] have predicted a very different structure for WB₄ (which we denote as WB'₄) and showed it to be metastable at normal conditions, but becoming stable at pressures >1 GPa.

In 2015, to resolve the positions of the boron atoms in the crystal structure of this boron-rich phase, Lech et al.^[13] have performed a neutron diffraction study, combined it with the analysis of previous structural models,^[8–12] and proposed a configuration that suggests a structural disorder and boron triangles instead of dimers^[8] or octahedra.^[9] These boron triangles are partially occupied and form a stacking where all the triangles in a layer have the same orientation different from that of the triangles in an adjacent layer. The resulting composition of "WB₄" was determined as WB_{4,2}.^[13]

Recent theoretical studies using USPEX,^[16–18] an ab initio global optimization technique, have predicted a new stable compound, superhard tungsten pentaboride WB_5 .^[4,5] The reported composition is pseudohexagonal, with an orthorhombic structure (space group *Pnnm*) and a similar structural motif containing boron triangles of different mutual in-plane orientation located within a metal layer. As we show below, the long-debated "WB₄" and the newly predicted WB₅ are actually the same material.

In this work, we present the results of the theoretical prediction, laboratory synthesis, and mechanical testing this potentially superhard material. We synthesized this material, measured its properties, and revealed the unexpected connection between theoretically predicted WB₅ and experimentally known WB₄: The new material has a crystal structure derived from the WB₅ structure type, with some amount of disorder and nonstoichiometry resulting in a wide range of chemical compositions described by formula WB_{5-x}. The calculations allowed us to identify the averaged crystal structures matching the experiment, reveal the preferable local structure of the material, and elaborate the specific patterns of atomic arrangements. Such structural models are suitable for detailed calculations of properties, which we did here to compare the results with experiment.

2. Results and Discussion

2.1. Crystal Structure Models and Structure Refinement

The crystal structure of the synthesized highest tungsten boride sample was determined using a combination of the theoretical structure models (**Figure 1**) and subsequent Rietveld refinement of the experimental XRD patterns. The sample, synthesized at 5 GPa and 1400 °C, was a mixture of two tungsten borides crushed into a fine powder to collect the XRD data.

To perform the Rietveld refinement, we constructed several new theoretical models on the basis of the previously predicted structure of $WB_5^{[4,5]}$ and our experimental results. On the one hand, the experimental data show a hexagonal symmetry compatible with space group $P6_3/mmc$. On the other hand, the orthorhombic WB_5 structure, which is pseudohexagonal and differs from the experimental hexagonal XRD pattern by only a few weak reflections resulting in the *R*-factor of 22%, can be used as an initial approximation for building hexagonal models that would match the experiment. Both the theoretically discovered $WB_5^{[4]}$ and experimentally proposed $WB_{4.2}$ phases^[13] contain triangular boron units, which we used for designing structural models for Rietveld refinement.

The first approximation, with the composition WB₅, belongs to $P\bar{6}m2$ space group (Figure 1a) and has a structural motif similar to orthorhombic WB₅ with boron triangles located in the vacancies of metal layers and the tungsten and boron positions fully occupied. The comparison of the simulated and experimental XRD patterns gives the *R*-factor of 14.6%, an improved but still unsatisfactory value. The failure of this model indicated that the hexagonal symmetry is achieved by disorder, therefore we decided to investigate disordered structures.

The second approximation (Figure 1b) is based on the previous one, $P\bar{6}m2$ -WB₅. To satisfy the likely symmetry $P6_3/mmc$ suggested by the experimental XRD patterns, we introduced two boron triangles with fractional occupations (i.e., the orientational disorder of the boron triangles) located in the vacancies of the tungsten honeycomb sublattice, in the same positions as the boron dimers in the Romans and Krug model.^[8] Thus, either the tungsten atoms in the Wyckoff position 2b or the boron atoms in position 12j (Table 1) have fractional occupancies. This model was used as a basis for the third approximation (Figure 1c), in which the center of gravity of boron triangles has the same position as the tungsten atom in site 2b (the Wyckoff position of each boron atom in the triangle is 6j), and both have partial occupancies (Table 1). This is structurally equivalent to the model proposed by Lech et al.^[13] The details of the crystal structure are summarized in Table 2.

The second and third models show a particularly close match with the experimental XRD patterns (Figure 1d,e). Performing spatial averaging, we obtained structure descriptions with partial occupancies and used these averaged structures in the Rietveld refinement. The final *R*-factors were 3.9% and 3.6% (Table 2). Refining the atomic positions and site occupancies in the second (Figure 1b) and third (Figure 1c) models, we obtained compositions WB_{4.86} and WB_{4.18}, respectively. The third model predicts a lower boron concentration (WB_{4.18}), compared with the composition defined by Lech et al.^[13] as WB_{4.31} (Table 1).

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Figure 1. Crystal structures of the proposed WB₅-based models: a) $P\bar{6}m2$ -WB₅, b) WB_{4.86}, and c) WB_{4.18}. Rietveld refinements of the synthesized sample according to models d) WB_{4.86} and e) WB_{4.18}. The experimental and calculated XRD patterns and the difference between them are shown in red, green, and purple, respectively. The positions of the Bragg reflections of WB₂ and WB_{4.86} (WB_{4.18}) are indicated by vertical ticks in the upper and lower rows, respectively.

2.2. Thermodynamic Stability of Proposed Models

To compare the thermodynamic stability of the proposed structural models, we performed first-principles total energy computations using density functional theory (DFT). Fractional occupancies of tungsten and boron in the second and third models mean a degree of disorder in these structures. To calculate their energies, supercells with different concentrations and orientations of boron triangles were constructed. For the second model (Figure 1b), we considered a supercell with composition WB4 89 and 159 atoms. For the third model (Figure 1c), six supercells were considered: two with composition $WB_{4,2}$ (156) and 52 atoms in the supercell), two WB_{445} (158 atoms each, with different relative positions of the boron triangles), one WB_{5.55} (478 atoms), and one WB_{5.71} (141 atoms). The local symmetry of each of these structures is low, whereas their global symmetry, corresponding to the averaged structure, is P6₃/mmc, consistent with the experimental data.

The resulting energies of formation are shown in **Figure 2b**. The energy of $P\bar{6}m2$ -WB₅ is higher by 11 meV per atom than that

of orthorhombic Pmmn-WB₅. The WB_{4.89} structure has a positive energy of formation of 0.11 eV per atom, which indicates its instability. The structures corresponding to the third structure model have much lower energies. Two supercells with composition WB_{4,2} are the best energetically, being only 3 meV per atom above the convex hull. The structures with a higher boron concentration, WB_{445} , are above the convex hull by only 7 and 13 meV per atom. Further increase in the boron concentration leads to a higher energy, which is above the convex hull by 66 and 82 meV per atom for $WB_{5.55}$ and $WB_{5.71}$, respectively. All of these structures are much more realistic than the previously proposed models for WB_4 . The energy of formation of WB_4 in the structure proposed by Romans and Krug^[8] is high and positive, 0.40 eV per atom, implying that this structure is impossible (Figure 2b). The structure proposed by Nowotny et al.^[9] has an even higher energy of formation of 0.96 eV per atom. After full relaxation, this structure changes significantly: its energy of formation becomes slightly negative (-0.091 eV per atom), but the structure no longer matches the experimental XRD pattern. Therefore, this model is also unrealistic.

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Table 1. Coordinates and site occupancy of tungsten and boron, *R*-factor, and calculated energy relative to the convex hull for the proposed models. Theoretical lattice parameters of *Pmmn*-WB₅ are: a = 5.199 Å, b = 6.369 Å, c = 8.993 Å. Experimental lattice parameters of hexagonal WB_{5-x} are: a = 5.20122 Å, c = 6.33601 Å.

Phase	Atom	Wyckoff position	Occupancy	x	Ŷ	Z	R [%]	ΔH [meV per atom]
Pmmn-WB ₅	W1	2b	1.0	0.500	0.000	0.918	22	7
	W2	2a	1.0	0.500	0.500	0.752		
	W3	2a	1.0	0.000	0.000	0.588		
	B1	8g	1.0	0.757	0.669	0.915		
	B2	8g	1.0	0.752	0.167	0.752		
	B3	8g	1.0	0.757	0.335	0.582		
	B4	4e	1.0	0.000	0.319	0.807		
	B5	2b	1.0	0.000	0.500	0.628		
Pēm2-₩B₅	W1	1b	1.0	0.000	0.000	0.500	10	18
	W2	3ј	1.0	0.333	0.667	0.000		
	W3	3ј	1.0	0.667	0.333	0.000		
	B1	120	1.0	0.333	0.333	0.250		
	B2	3k	1.0	0.547	0.453	0.500		
P6 ₃ /mmc-WB _{4.86}	W1	2c	1.0	0.333	0.667	0.250	3.9	318 ^{a)}
	W2	2b	0.663(2)	0.000	0.000	0.250		
	B1	12i	1.0	0.333	0.000	0.000		
	B2	12j	0.347(5)	0.333	0.500	0.750		
P6 ₃ /mmc-WB _{4.18}	W1	2c	1.0	0.333	0.667	0.250	3.5	3 ^{b)}
	W2	2b	0.690(2)	0.000	0.000	0.250		
	B1	12i	1.0	0.333	0.000	0.000		
	B2	6h	0.3569(5)	0.119	0.238	0.250		
P6 ₃ /mmc-WB _{4.31} ^[13]	W1	2c	1.0	0.333	0.667	0.250	3.5	-
	W2	2b	0.641(2)	0.000	0.000	0.250		
	B1	12i	1.0	0.33167	0.000	0.000		
	B2	6h	0.3569(1)	0.11887	0.23775	0.250		

^{a)} Energy obtained for the WB_{4.89} composition ^{b)} Energy obtained for the WB_{4.2} composition.



Figure 2. a) Calculated convex hull of the W–B system. b) Enlarged boron-rich part of the convex hull shown in yellow in panel (a). Metastable and thermodynamically stable phases and model highest tungsten borides are shown by white circles, blue squares, and red triangles, respectively.

The relation between the arrangement of boron atoms and energy of the structures allowed us to formulate the following rules determining the most energetically favorable local structures:

- atoms in the Wyckoff position 2b replaced with three boron atoms arranged in triangles whose plane coincides with that of the tungsten atoms (Figure S5a, Supporting Information).
- 1) As the boron content increases, the structure of WB_x (x > 3) may be described as $P6_3/mmc$ -WB₃ with some of the tungsten

2) Along the *c*-axis, the preferred neighbor of a boron triangle is a tungsten atom, not another boron triangle (Figure S5b, Supporting Information).

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Table 2. Rietveld refinement of the proposed models.

	Crystal data	
Chemical formula	WB _{4.86}	WB _{4.18}
Chemical weight	236.28	229.04
Space group	P6 ₃ /mmc	P6 ₃ /mmc
a [Å]	5.201	22(2)
c [Å]	6.336	01(4)
V [Å ³]	148.3	81(1)
Ζ	4	Ļ
ho [g cm ⁻³]	8.798	8.665
Weight ratio	86	(6)
Radiation type	Cul	Κα1
Wavelength [Å]	1.540	5981
Temperature [K]	29	93
Diffractometer	Guinier Imaging Plate Camera	
Refinement	GS	AS
R _F	0.0390	0.0358
R _P	0.0440	0.0418
R _{WP}	0.0677	0.0651

3) The boron triangles are oriented so as to maximize the distance between the boron atoms and the nearest neighboring in-plane tungsten atoms. Therefore, the boron triangles of even and odd planes are rotated by 180° with respect to each other (Figure S5c, Supporting Information).

The theoretical WB_5 structure and $WB_{4,2}$ model proposed by Lech et al.^[13] both satisfy these rules of low-energy structures. Many other low-energy structures with various compositions can be constructed on the basis of these rules.

A systematic construction of intermediate compositions for the WB_{5-x} structures can be accomplished using the lattice model developed for the study of boron-rich molybdenum borides.^[19] This model accounts for the rules of low-energy local structures formulated above. After the parameterization of the lattice model for the boron-rich W-B system, we considered a $2 \times 2 \times 3$ supercell and calculated the lowest-energy arrangements of the triangular boron units for compositions from WB₃ to WB₉. The results are summarized in Figure 3, where the energies of formation of the DFT-calculated boron-rich phases are shown together with those from the lattice model. A broad range of low-energy compositions of WB_{5-x} may be obtained within this structure type. The most stable composition, closest to the convex hull line, corresponds to WB_{4.2}. The same situation has been observed in the Mo-B system^[19] where the most stable composition was found to be MoB_{47} , which is even closer to MoB_5 .

The rules formulated above that determine local structures make it possible to easily generate structures for any intermediate composition. The crystal structure of the studied material has a structural motif of predicted WB_5 and the composition WB_{5-x} caused by a disorder and fractional occupancy of the boron atoms (Table 1 and ref. [13]).

Because tungsten borides are synthesized at a high temperature, we also consider the vibrational contributions to the Gibbs



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Figure 3. Boron-rich part of the convex hull diagram. Stable and metastable phases obtained during the evolutionary global optimization are shown by filled squares and hollow circles, respectively. Boron-rich phases constructed on the basis of the proposed structural models and used for the XRD refinement are shown by triangles. Small blue dots depict the WB_{5-x} structures obtained using the parameterized lattice model.

free energy of boron-rich phases within the quasiharmonic approximation. The resulting convex hull diagrams for different temperatures are shown in **Figure 4**. An increase in temperature leads to the destabilization of the previously theoretically proposed WB'_4 structure, whereas $WB_{4,2}$ and WB_5 become stable at high temperatures. These results confirm that boron-rich phases in the W–B system have a broad homogeneity region extending up to WB_5 , and are consistent with the conclusion that the obtained material has a disordered structure with the composition close to $WB_{4,2}$.

Because of the disordered nature of the WB_{5-x} system where many different arrangements of the triangular boron units (B₃) with similar energy exist, it is important to discuss the influence of the configurational entropy on the thermodynamic stability of these compounds. The estimates made within the lattice model for a 2 × 2 × 4 supercell show the configurational entropy values of 2.0 and 0.4 J mol⁻¹ K⁻¹ at 2000 K for WB_{4.2} and WB₅, respectively. The lower value for the WB₅ composition is explained by the strong repulsive interaction between the nearest-neighbor B₃ units, which limits the number of low-energy configurations.

2.3. Mechanical Properties of the Synthesized Samples

Compacts based on the highest tungsten boride were obtained at temperatures of the sintering reaction (1200–1300 °C) and pressures of about 1.5 GPa and have a fine-grained structure with particles smaller than 1 μ m. The results of the transmission electron microscopy (TEM) analysis are shown in **Figure 5**a. The WB_{5-x} crystals formed under high-pressure conditions consist of slightly disoriented blocks.

An excess of boron is required to obtain the maximum content of the highest tungsten boride WB_{5-x} , and it was not possible to fully realize the physical and mechanical characteristics of the material because of the residual boron in the sample.



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Figure 4. Boron-rich part of the convex hull calculated at a) 500 K, b) 1000 K, c) 1500 K, and d) 2000 K. The stabilization of the WB₅ and WB_{4.2} phases and destabilization of WB₄ occur with the temperature increase. Thermodynamically stable and metastable phases are shown by full and open circles, respectively.

The Rockwell hardness of the highest tungsten boride and hard alloy 94WC-6Co was measured with a load of 600 N. The average area of imprints of a diamond indenter is 1.5 times smaller compared with the hard alloy (Figure 5b), and the hardness of the sample of WB_{5-x} is about 30 GPa.

The Vickers microhardness was measured on EMCO-TEST DuraScan-20 with a 2 N load on the indenter. The average value of 15 measurements of the microhardness of WB_{5-x} is 29.3 GPa (**Table 3**), lower than the calculated value because an excess of amorphous noncrystalline boron softens the compact. However, this value is comparable to 32.8 GPa obtained by Mohammadi et al.^[20] for "WB₄" with the same load of 2 N. The Vickers hardness of the hard alloy 94WC-6Co, measured using the same experimental setup, is 22 GPa.

For the sample made using the arc melting technique, the average value of the Vickers hardness is 33.9 GPa, whereas the maximum value is close to 37 GPa, approaching the level of superhard materials (Table 3). These values are close to those that have been previously measured for "WB₄" also prepared by arc melting, which showed the Vickers hardness of 32.8 GPa with the load of 2 N.^[20,21] This sample has a much denser structure compared with the one made using the thermobaric treatment (both composites contained more than 80% of WB_{5-x} and less than 20% of WB₂ and pure boron). The microhardnesses of the pressuresintered and arc-melted composites are higher than that of tungsten carbide by \approx 30% and 50%, respectively (Table 3). However, the arc-melted sample (Figure S2, Supporting Information) contains grains as large as 100 µm, which makes the composite more brittle. The structural elements in the sintered sample have an average size of 1 µm.

The elastic moduli were measured on samples with a diameter of 10–13 mm and a height of 5–7 mm. The bulk modulus *B* is 180–210 GPa and the shear modulus *G* is 190–230 GPa. The samples with the best quality have B = 205 GPa and G = 220 GPa.





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Figure 5. a) TEM images of crystals of WB_{5-x}. b) Imprints of a diamond Rockwell indenter on WB_{5-x} (left) and on a hard alloy (right).

 Table 3. Vickers microhardness of the tungsten boride and tungsten carbide samples.

No. of measurement	H _V [GPa]					
	94WC-6Co	WB _{5-x} -WB ₂ -B, sintering	WB _{5-x} -WB ₂ -B, arc melting			
1	23.5	28.9	36.5			
2	20.6	30.3	34.7			
3	24.2	27.2	35.5			
4	19.7	30.8	34.9			
5	21.0	28.9	34.6			
6	24.9	30.4	32.2			
7	22.6	27.7	29.5			
8	24.1	29.4	35.1			
9	22.2	29.5	34.4			
10	23.9	28.3	33,8			
11	19.7	30.7	33.5			
12	21.3	28.8	32.7			
13	20.4	28.3	34.9			
14	20.9	30.4	36.8			
15	21.4	29.9	28.9			
Average	22.0	29.3	33.9			

The residual boron, which could amount to 20% of the sample volume, is poorly connected with tungsten boride. The porosity of the samples was low but potentially nonzero, approximately several percent. Considering that these factors could sharply decrease the measured moduli, the actual elastic moduli of the highest tungsten boride may be 25–30% higher.

To calculate the Vickers hardness, the empirical model proposed by Chen et al.^[22] and the Mazhnik–Oganov model^[23] were used, whereas the fracture toughness was calculated using the recently developed Mazhnik–Oganov^[23] and Niu–Oganov^[24] models. The calculated mechanical properties of the proposed disordered models (**Table 4**) are lower than those of *Pmmn*-WB₅.^[4] The values for the experimental sample were calculated from the measured bulk and shear moduli using the corresponding empirical models.^[22,24] Our best theoretical estimate of hardness, by the Mazhnik–Oganov model for WB_{4,2}, is 39.2 GPa, which is extremely high and very close to the highest experimental value for the sample obtained by arc melting.

3. Conclusion

The synthesis of the boron-richest tungsten boride was guided by the crystal structure prediction using the evolutionary algorithm USPEX.^[16–18] The careful theoretical examination and

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Table 4. Mechanical properties of the proposed models compared with the experimental values of the bulk and shear moduli *B* and *G*, and the energy above the convex hull ΔH . The Vickers hardness H_V and fracture toughness K_{IC} were calculated using the Mazhnik–Oganov model.^[23] The values in parentheses were calculated using the empirical models.^[22,24]

Phase	B [CPa]	C [CPa]	C/B	H [CPa]	K [MPa m ^{0.5}]	AH [meV per atom]
	D [OI a]	0 [01 a]	0/6		KIC [IVII a III]	
Pmmn-WB ₅	295	270	0.915	45.2 (45.1)	4.59 (4.01)	7
<i>P</i> 6 <i>m</i> 2-WB ₅	288	224	0.775	30.2 (32.2)	4.27 (3.62)	18
WB _{4.2}	280	245	0.875	39.2 (36.6)	3.97 (3.64)	3
WB _{5.71}	286	258	0.902	42.7 (42.5)	4.21 (3.86)	82
WB ₂	313	248	0.792	33.5 (35.8)	5.0 (4.06)	0
WB _{5-x} -WB ₂ -B	205	220	1.073	40.7 ^{a)} (47.9 ^{a)})	3.31^{a} (3.01 ^a)	-
"WB ₄ "	339 ^[20]			$43.3 \pm 2.9^{[20]^{\rm b)}}$		-

^{a)} Calculated using the empirical models^[22,24] on the basis of the experimentally measured *B* and *G*. The values of *B* and *G* were measured for the WB_{5-x} - WB_2 -B sample, therefore they cannot fully characterize the mechanical properties of pure WB_{5-x} ; ^{b)} The applied load was 0.49 N.

comparison of the experimental and theoretical data show that the obtained material has a disordered structure with the composition close to WB_{4,2}, but we expect a broad homogeneity region between WB₄ and WB₅. The comprehensive theoretical investigation allowed us to determine the crystal structure of the synthesized tungsten boride and resolve the puzzle of "WB4" widely discussed over the past 50 years. The experimentally obtained material is structurally derived from previously predicted WB₅ and can be referred to as WB_{5-x} . It is different from WB'_4 —a phase with a predicted stability at pressures above 1 GPa but not synthesized as yet. The measured mechanical properties of WB_{5-x} are in close agreement with predictions that indicate exceptional characteristics (theoretical hardness of 39 GPa and fracture toughness of 4 MPa m^{0.5}). The hardness of composites based on this phase is 30-50% higher than that of the hard alloy 94WC-6Co. Furthermore, the WB_{5-x} -based composite has excellent thermal stability: the sample shows no degradation to at least 1000 °C, whereas 94WC-6Co oxidized and broke at 800 °C (see the Supporting Information). The combination of excellent mechanical properties, thermal stability, and inexpensive synthesis at mild conditions means that such composites may replace traditional hard alloys in many applications.

4. Experimental Section

To determine the optimal thermobaric conditions for the synthesis of WB₅-based compacts, a wide range of pressures suitable for large-scale production was tested. The aim was not only to synthesize this particular phase, but to achieve the desired physical and mechanical properties of the obtained samples. Initially, the powders of tungsten and boron had sub-micrometer-size particles. Their mixture with the atomic ratio W:B = 1:7 was used for synthesis. It was found that both the synthesis of borides and sintering of crystals into sufficiently strong compacts occur at high pressures of 4–7 GPa and 1000 °C. The X-ray phase analysis of the samples showed the presence of two borides, WB_{5-x} and WB₂, and the content of tungsten diboride increased with pressure and temperature (Table S1, Supporting Information). Some amount of unreacted boron remained in the samples after the thermobaric synthesis and sintering.



Figure 6. Microstructure of the WB_{5-x} sample with a minimum content of WB_2 showing the dense fine crystalline material.

Compacts with a high content of WB_{5-x} were also obtained at a lower pressure of 1.5 GPa and temperatures of 1100-1300 °C from a powder mixture of tungsten and boron. The content of WB_{5-x} was in the range from 75% to 95%. Samples with the maximum content of the highest tungsten boride were obtained at temperatures of 1000–1100 °C, but in this case, WB_{5-x} had many defects (very broad diffraction peaks). Therefore, a wellcrystallized sample containing 86(6)% of the highest tungsten boride and 14(4)% of WB2 was taken for the structure refinement (see below). During the cleavage of the samples with a maximum content of WB5-x, micrometer and sub-micrometer crystals with a clearly visible faceting were observed (Figure 6). For comparative analysis, compacts of the highest tungsten boride were sintered by hot isostatic pressing at a pressure of 0.03 GPa and a temperature of 1400 °C. In contrast to the high-pressure synthesis, hot isostatic pressing produces porous compacts containing about 70% of WB_{5-x} and 30% of WB_2 . Arc melting was performed in argon at a temperature of about 2100 °C, leading to the synthesis of dense and hard samples.

The synthesis of WB_{5-x} was carried out in high-pressure cells of the toroid and piston-cylinder type. Pressed calcite strips were

used for sealing and electrically insulating the piston. The experiments were conducted at a pressure of 1.5 GPa. Heating was done by a cylindrical graphite resistive heater. The pressure in the toroid-type chamber was generated in a limestone cell placed between hard alloy anvils with a special profile. The pressure calibration was done at room temperature by measuring the electrical conductivity and observing the phase transition of Bi (at 2.55 and 7.7 GPa). The temperature was measured by thermocouples. The high-pressure treatment was carried out at 4–7 GPa in the temperature range of 1100–1500 °C. After the stabilization of the applied pressure, the samples were heated up to the required temperature, and then the cells were quenched to room temperature before the pressure release.

The exposure time at the maximum temperature during the synthesis was 0.5–1.5 min in the toroid high-pressure cell and about 10 min in the piston-cylinder cell.

The particles of tungsten and boron powders used in this work were smaller than 1 µm and had a distribution close to isotropic (Figure S3, Supporting Information). The experiments were performed with tungsten and boron mixed in 3:10 proportion by weight. The mixture was pressed into tablets with a diameter of 5 mm and a height of 3 mm for sintering in the toroid highpressure cell and those with a diameter of 15 mm and a height of 7 mm for sintering in the piston-cylinder high-pressure cell.

Another sample was obtained by melting in an electric arc vacuum furnace LK200I (Leybold-Heraeus) with a nonexpendable tungsten electrode. A sample weighing 20 g formed from a powder mixture of tungsten with 30 wt% of boron was placed in a copper water-cooled mold. During melting, the voltage on an electric arc was varied from 20 to 30 V, and the current was 1000 A.

Guinier Imaging Plate Camera G670 (Huber) with Cu K α_1 radiation was used for the X-ray phase analysis of samples; the structure refinement was done using the GSAS software package. The microstructure study and elemental analysis were performed using the scanning electron microscope (JEM) JEOL JSM-6390 equipped with the EDS analyzer INKA.

The measurements of the elastic properties of compacts were carried out using the pulsed ultrasonic method with a device based on the PXI platform (National Instruments) with the system registering the passed and reflected ultrasonic signals. The velocities of propagation of longitudinal and shear waves were determined with an error below 0.5%. The bulk and shear moduli were calculated in the approximation of a homogeneous isotropic medium.

The microhardness was measured with the load of 2 N for two composites made of a mixture of tungsten and boron powders, one of the composites sintered at a pressure of 1.5 GPa, and the other obtained by arc melting.

5. Computational Section

Our calculations are based on DFT^[25,26] within the generalized gradient approximation (the Perdew–Burke–Ernzerhof functional)^[27] and the projector augmented-wave method^[28,29] as implemented in VASP^[30–32] code. The plane wave energy cutoff of 400 eV, the Methfessel–Paxton smearing^[33] of electronic occupations, and Γ -centered *k*-point meshes with a resolution of $2\pi \times 0.025$ Å⁻¹ for the Brillouin zone sampling were used, ensuring the convergence of the energy differences and stress tensors.

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The Vickers hardness and fracture toughness were calculated using the recently developed Mazhnik–Oganov model $^{\left[23\right] }$

$$H_{\rm V} = \gamma_0 \,\chi \left(\nu\right) E \tag{1}$$

where $\gamma_0 = 0.096$, *E* is Young's modulus, and $\chi(v)$ is a dimensionless function of Poisson's ratio^[23]

$$\chi(\nu) = \frac{1 - 8.5\nu + 19.5\nu^2}{1 - 7.5\nu + 12.2\nu^2 + 19.6\nu^3}$$
(2)

The fracture toughness was calculated as^[23]

$$K_{\rm IC} = \alpha_0^{-\frac{1}{2}} \cdot V_0^{\frac{1}{6}} \cdot [\zeta(\nu) E]^{\frac{3}{2}}$$
(3)

where α_0 depends on the chemical bonding in the material and has the units of pressure, V_0 is the volume per atom, and $\zeta(v)$ is a dimensionless function of Poisson's ratio

$$\zeta(\nu) = \frac{1 - 13.7\nu + 48.6\nu^2}{1 - 15.2\nu + 70.2\nu^2 - 81.5\nu^3}$$
(4)

To get an idea of uncertainties of these estimates, we also applied Chen's model to evaluate the Vickers hardness and the Niu–Oganov model to calculate the fracture toughness. According to Chen's model,^[22] the Vickers hardness $H_{\rm V}$ is

$$H_{\rm V} = 2 \cdot (k^2 \cdot G)^{0.585} - 3 \tag{5}$$

where *k* is the Pugh ratio (k = G/B), *G* is the shear modulus, and *B* is the bulk modulus (both moduli were obtained from the Voigt–Reuss–Hill averaging and expressed in GPa).

In the Niu–Oganov empirical model, $^{[24]}$ the fracture toughness $K_{\rm IC}$ is

$$K_{\rm IC} = \alpha \cdot V^{\frac{1}{6}} \cdot G \cdot \left(\frac{B}{G}\right)^{\frac{1}{2}}$$
(6)

where α is the enhancement factor accounting for the degree of metallicity, *V* is the volume per atom, and *G* and *B* are the shear and bulk moduli. For insulators, semiconductors, carbides, nitrides, and borides, $\alpha = 1$.

Supporting Information

Supporting Information is available from the Wiley Online Library or from the author.

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Conflict of Interest

The authors declare no conflict of interest.

Keywords

mechanical properties, superhard materials, tungsten borides, USPEX, X-ray diffraction

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