New Zirconium Diboride Polymorphs—First-Principles Calculations

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Abstract: Two new hypothetical zirconium diboride (ZrB₂) polymorphs: (hP6₃-P6/mmc-space group, no. 194) and (oP6-Pmmm-space group, no. 59), were thoroughly studied under the first-principles density functional theory calculations from the structural, mechanical and thermodynamic properties point of view. The proposed phases are thermodynamically stable (negative formation enthalpy). Studies of mechanical properties indicate that new polymorphs are less hard than the known phase (hP3-P6/mmm-space group, no. 191) and are not brittle. Analysis of phonon band structure and density of states (DOS) also show that the phonon modes have positive frequencies everywhere and the new ZrB₂ phases are not only mechanically but also dynamically stable. The estimated acoustic Debye temperature, Θ_D, for the two new proposed ZrB₂ phases is about 760 K. The thermodynamic properties such as internal energy, free energy, entropy and constant-volume specific heat are also presented.

Keywords: zirconium diboride; ab initio calculations; mechanical properties; elastic properties; phonons

1. Introduction

During latest years, the transition metal borides have attracted attention among materials researchers due to the combination of their outstanding physical properties such as electric and thermal conductivity comparable with metals, low compressibility, high shear strength and exceptionally high hardness [1,2]. Even in the form of thin films, they possess extraordinary properties such as very high hardness with increased flexibility, great thermal properties and very good corrosion and wear resistance [3–5]. Among borides, zirconium diboride (ZrB₂) deserves special attention. ZrB₂, with melting temperature 3245 °C [6], is a member of a family of materials known as ultra-high temperature ceramics (UHTCs). In addition to high melting temperatures, ZrB₂ has a unique combination of chemical stability, high electrical and thermal conductivities and resistance to erosion and corrosion that makes its suitable for the extreme chemical and thermal environments associated with, for example, hypersonic flight, atmospheric re-entry and rocket propulsion [7]. According to the Zr–B phase diagram [8,9] there are three phases, namely, ZrB, ZrB₂, and ZrB₁₂, which have been reported and widely studied for this system. Theoretical investigations show that ZrB can create different crystallographic structures. The basic phase is NaCl-type face-centered cubic ZrB (Fm-3m-space group, no. 225) with lattice constant a = 4.900 Å [10]. Furthermore, ZrB also can crystallize in a FeB-type structure with a primitive orthorhombic (Pnma) crystal structure [9,11], CrB-type orthorhombic structure with Cmcm-space group [11,12] and hexagonal Pmmm [13]. A literature review shows that ZrB₁₂ is only stable in one structure type. The cubic LuB₁₂ structure (Fm3m-space group, no. 225) with lattice constant a = 7.4085 Å was studied theoretically and experimentally for example in Reference [14]. They compared electric-field gradient measurements...
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at the B sites and first-principles calculations in order to analyse the chemical bonding properties. Both experimental and theoretical results were in good agreement. In Reference [15], however, the mechanical properties of ZrB$_2$ were calculated. The Vickers hardness of zirconium boride was 32.9 GPa, which is in good agreement with other theoretical results [14,16,17].

In contrast to ZrB$_2$, depending on the calculation method, different hardness values have been determined for zirconium diboride ZrB$_2$. The calculated values of hardness range from 12.82–55.53 GPa [18], whereas experimentally measured hardness reached 23 ± 0.9 GPa [6,19]. All mentioned structures were assigned as ZrB$_2$ with the crystal hexagonal structure of AlB$_2$-type with the P6/mmm-space group, no. 191. Such large differences in the values of the analysed properties may, however, come not only from differences in calculation methods but also from the possibility of the existence of other stable forms of ZrB$_2$, which may form nanocomposites of different polymorphs of ZrB$_2$. A similar conclusion about the possibility of the existence of other ZrB$_2$ crystal types can be drawn on the basis of other studies on possible forms of transition metal diborides, for example, WB$_2$ [20,21] or ReB$_2$ [22,23]. For comparison, in Reference [21] the authors proposed six different phases of WB$_2$. In the case of ZrB$_2$, it is hard to find such a study.

It should also be noted that in addition to the phases appearing in the Zr-B equilibrium diagram [8,24], other zirconium and boron compounds have been theoretically determined. There are hypothetical Zr-B phases such as: Zr$_3$B$_4$, Zr$_2$B$_3$, Zr$_3$B$_2$ [16], ZrB$_3$ [25], ZrB$_4$ [26] and ZrB$_6$ [13]. All polymorphs are both mechanically and dynamically stable but have not been confirmed experimentally yet. In this work, structural, mechanical and thermodynamic properties of stable ZrB$_2$ polymorphs from density functional calculations will be studied.

2. Computational Methods

First-principle calculations based on density functional theory (DFT) [27,28] within the pseudopotential plane-wave approximation (PP-PW) implemented in ABINIT code were performed in this work. Projector augmented-wave formulation (PAW) pseudopotentials [31] were used to describe the interactions of ionic core and non-valence electrons.

To enhance the confidence of the calculations as an exchange-correlation (XC) functional, three approximations were used: local density approximation (LDA) [32,33], classical Perdew-Burke-Ernzerhof (PBE) generalised gradient approximation (GGA) [34] and modified Perdew-Burke-Ernzerhof GGA for solids (PBEsol) [35]. There is a strong view that the PBEsol is the overall best performing XC functional for identifying the structure and elastic properties [36–38].

Projector augmented wave method (PAW) pseudopotentials used for LDA and PBE XC functionals were taken from PseudoDojo project [39]. PAW pseudopotentials for PBEsol exchange-correlation functional [35] were generated using ATOMPAW software [40] and a library of exchange-correlation functionals for density functional theory, LibXC [41].

All calculations were made by tuning the precision of the calculations, which was done by automatically setting the variables at accuracy level 4 (accuracy = 4 corresponds to the default tuning of ABINIT). The cut-off energy consistent with PAW pseudopotentials of the plane-wave basis set was 15 Ha with 4$d^55s^2$ valence electrons for Zr and $2s^22p^1$ valence electrons for B. K-PoinTs grids were generated with kptrlen = 30.0 (grids that give a length of smallest vector LARGER than kptrlen). Metallic occupation of levels with the Fermi-Dirac smearing occupation scheme and tsmear (Ha) = 0.02 was used in all ABINIT calculations.

2.1. Optimization of Structures

As mentioned earlier, tungsten diboride [20] and rhenium diboride [23] crystallise in various space groups. Searching for new structures of ZrB$_2$, we started with basic cells of hP6-P6$_3$/mmc-WB$_2$ and oP6-Pmmm-WB$_2$ and replaced tungsten atoms with zirconium atoms, wherein the designations mean: Pearson symbol, Space group and Chemical formula [20]. Then, all structures were relaxed by
using the Broyden-Fletcher-Goldfarb-Shanno minimisation scheme (BFGS) with full optimisation of cell geometry and atomic coordinates. Maximal stress tolerance (GPa) was set to $1 \times 10^{-4}$.

2.2. Formation Enthalpy and Cohesive Energy

The formation enthalpy and cohesive energy were determined as follows \[23,42\]:

$$\Delta_f H(ZrB_2) = E_{coh}(ZrB_2) - E_{coh}(Zr) - 2E_{coh}(B),$$

$$E_{coh}(ZrB_2) = E_{total}(ZrB_2) - E_{iso}(Zr) - 2E_{iso}(B),$$

where $\Delta_f H(ZrB_2)$ is the formation enthalpy of the $ZrB_2$; $E_{coh}(ZrB_2)$ is the cohesive energy of the $ZrB_2$; $E_{coh}(Zr)$ is the cohesive energy of Zr; $E_{coh}(B)$ is the cohesive energy of B; $E_{tot}(ZrB_2)$ is the total energy of the ZrB$_2$; $E_{iso}(Zr)$ is the total energy of a Zr atom and $E_{iso}(B)$ is the total energy of a B atom.

The cohesive energy of the $E_{coh}(Zr)$ is calculated, taking into account stoichiometry, as the total energy difference of zirconium crystal ($hp2$-$P6/mmc$-space group, no.194) and single Zr atom in the box, whereas $E_{coh}(B)$ is the total energy difference of a-boron crystal ($hR12$-$R-3m$-space group, no. 166) and single B atom in a sufficiently large box \[23\].

2.3. Mechanical Properties Calculations

The theoretical ground state elastic constants $C_{ij}$ of all structures were established with the metric tensor formulation of strain in density functional perturbation theory (DFPT) \[43\]. Isotropised bulk modulus $B$, shear modulus $G$, Young’s modulus $E$ and Poisson’s ratio $\nu$ were estimated by means of a Voigt–Reuss–Hill average \[44,45\].

In order to verify the elastic stability of all the structures, positive definiteness of the stiffness tensor was checked \[46\] by calculating Kelvin moduli, that is, eigenvalues of stiffness tensor represented in second-rank tensor notation \[47\].

Hardness of ZrB$_2$ polymorphs in the present paper was calculated with the use of semi-empirical relation proposed in Reference \[48\]. The equation is defined as follow:

$$H_v = 0.92 \left( \frac{G}{B} \right)^{1.137} G^{0.708}.$$  

$G/B$ ratio appearing in the above formula named Pugh’s modulus ratio \[49\] is commonly used as a universal ductile-to-brittle criterion.

2.4. Phonon and Thermodynamic Properties Calculations

To calculate phonons, DFPT was utilised \[29,30\]. The phonon dispersion curves \[50\] of the analysed structures were then used to determine their dynamical stability \[46,51\], complementary to elastic stability. Acoustic Debye temperature was calculated from the phonon densities of states (DOS).

Using the calculated phonons under the harmonic approximation, that is, in the range up to Debye temperature, thermal quantities: phonon internal energy, free energy, entropy and constant-volume heat capacity as a function of the temperature were determined \[24,52\].

3. Results

Using the approach described in Section 2.1, the first step in our calculations was the geometry optimisation of the two new hypothetical (ZrB$_2$) polymorphs—($hp6$-$P6_3/mmc$-space group, no.194) and ($oP6$-$Pmmn$-space group, no. 59), and the formerly known polymorph ($hp3$-$P6/mnm$-space group, no. 191).
3.1. Structural Properties

The basic cells for all three analysed polymorphs are depicted in Figure 1, whereas the crystallographic data, calculated with three different XC functionals (LDA, PBE and PBEsol), are stored in crystallographic information files (CIFs) in Supplementary Materials.

Determined lattice parameters, formation enthalpy and cohesive energy for known \textit{hP3-P6/mmm} polymorph (Figure 1a) are comparable to those of other authors [18] (see Table 1). We treat this as a verification of the correctness of the methodology used.

The first new hypothetical phase, \textit{hP6-P63/mmc} (Figure 1b), also crystallises in the hexagonal system but has 6 atoms in the cell, whereas the second new hypothetical phase, \textit{oP6-Pmmn} (Figure 1c), crystallises in the orthorhombic system and also has 6 atoms in the cell. There is little sense to compare lattice parameters for phases in different systems, but it is worth comparing the formation enthalpy and the cohesive energy. It can be seen that the formation enthalpy, $\Delta_f H$, for new phases is significantly lower than for the known \textit{hP3-P6/mmm} phase and comparable between the new phases (see Table 1). The calculated cohesive energy, $E_c$, is a little higher than for the known \textit{hP3-P6/mmm} phase and again comparable between the two new phases. These two facts suggest that the new phases are comparably thermodynamically stable but less stable than the known \textit{hP3-P6/mmm} phase.

![Figure 1. ZB$_2$-Basic cells: (a) hP3-P6/mmm, (b) hP6-P63/mmc and (c) oP6-Pmmn.](image)

3.2. Mechanical Properties

Computed elastic constants, Kelvin moduli, isotropised bulk, shear and Young’s modulus, Poisson’s ratio, $C/B$ Pugh’s modulus ratio, Debye temperature and estimated hardness of all analysed zirconium diboride structures are listed in Table 1. For known \textit{hP3-P6/mmm} phase (Figure 1a), these quantities are comparable to those of other authors [18], which is a further verification of the validity of the methodology used.
Analysing the data received, it can be concluded that the new phases have lower mechanical parameters than the known \( hP3-P6/mmm \) phase, except for the Poisson’s ratio. Both new phases have a similar isotropised bulk modulus of about 170 GPa, but the shear modulus for the \( hP6-P6_3/mmc \) phase is much lower and is only about 40 GPa. The consequence of this is that the hardness of the \( hP6-P6_3/mmc \) phase is only 2 GPa, while for the \( oP6-Pmmn \) phase it is about 14 GPa. A high \( G/B \) Pugh’s modulus ratio would correspond to a more brittle than ductile character of material. The critical value, separating ductile from brittle materials, is approximately 0.571 \([49,53]\). It can be seen that the known \( hP3-P6/mmm \) phase is brittle, the \( hP6-P6_3/mmc \) phase is ductile and the \( oP6-Pmmn \) phase is somehow between brittle and ductile.

All analysed structures have a positive definite stiffness tensor and positive Kelvin moduli, that is, eigenvalues of stiffness tensor represented in \textit{second-rank tensor} notation, so there are mechanically stable (see Table 1).
Table 1. Lattice parameters (Å); formation enthalpy $\Delta_f H$ (eV/Atom); cohesive energy $E_c$ (eV/Atom); elastic constants $C_{ij}$ (GPa); Kelvin moduli $K_i$ (GPa); bulk modulus $B$ (GPa); shear modulus $G$ (GPa); Young’s modulus $E$ (GPa); Poisson’s ratio $\nu$; $G/B$ Pugh’s modulus ratio; Debye temperature $\Theta_D$ (K); hardness $H_v$ (GPa); of ZrB$_2$ phases: ZrB$_2$ (hP$_3$-P6/mmm-space group, no. 191; hP6-P6$_3$/mmc-space group, no. 194; and oP6-P6/mmm-space group, no. 59). Experimental and calculated data for hP3-P6/mmm phase are taken from Reference [18].

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<th>Exp.</th>
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<th>PBEsol</th>
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<td>554</td>
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3.3. Phonon and Thermodynamic Properties

Phonon dispersion curves along the high symmetry q-points [50] and phonon densities of states (DOS) calculated with the use of PBEsol exchange-correlation (XC) functional for known hP3-P6/mmm phase, Figure 1a), are depicted in Figure 2, for the phase hP6-P6$_3$/mmc, Figure 1b), in Figure 3 and for the phase oP6-Pmnn, Figure 1c), in Figure 4, respectively. Phonon results for all exchange-correlation (XC) functionals are stored in Supplementary Materials. Analysis of the calculated curves allows us
to state that phonon modes everywhere have positive frequencies and the new ZrB₂ phases are not only mechanically but also dynamically stable. The estimated acoustic Debye temperature Θ_D for the two new proposed ZrB₂ phases is about 760 K and is about 200 K lower than that for the known hP3-P6/mmm phase and it is consistent with the mechanical properties, see Table 1. Results for thermodynamic properties up to 760 K for the three zirconium diboride polymorphs calculated with the use of PBEsol exchange-correlation (XC) functional, that is, phonon internal energy, free energy, entropy, constant-volume heat capacity are depicted in Figures 5–7 and it can be seen that are very similar for the two new proposed ZrB₂ polymorphs. This additional fact suggests again that the new phases are comparably thermodynamically stable up to Debye temperature Θ_D, but less stable than the known hP3-P6/mmm phase.

ZrB₂-191 Phonon bands + DOS in cm⁻¹ (PBEsol)

Figure 2. ZrB₂(hP3-P6/mmm-space group, no. 191)-phonon band structure and densities of states (DOS).
**Figure 3.** $\text{ZrB}_2$ ($\text{hP6}-\text{P6}_3$/mmc-space group, no. 194)-phonon band structure and DOS.

**Figure 4.** $\text{ZrB}_2$ ($\text{oP6}-\text{P6}$/mmm-space group, no.59)-phonon band structure and DOS.
Figure 5. ZrB$_2$(hP3-P6/mmm-space group, no. 191)-thermodynamic properties: internal energy, free energy, entropy and constant-volume specific heat.

Figure 6. ZrB$_2$(hP6-P6/mmc-space group, no.194)-thermodynamic properties: internal energy, free energy, entropy and constant-volume specific heat.
Temperature (K)
0.5
1.0
U(T) (eV/cell)
internal_energy

Temperature (K)
0.5
0.0
F(T) + ZPE (eV/cell)
free_energy

Temperature (K)
0.0000
0.0005
0.0010
0.0015
c(V,T) (eV/cell)
cv

Figure 7. ZrB$_2$\((oP6-P6/mmc\)-space group, no.59) - thermodynamic properties: internal energy, free energy, entropy and constant-volume specific heat.

4. Conclusions

In the present paper, extensive analysis of two new hypothetical and one previously known zirconium diboride (ZrB$_2$) polymorphs within the framework of density functional theory from the structural, mechanical and thermodynamic properties point of view was performed. We can conclude that:

- two new hypothetical zirconium diboride (ZrB$_2$) polymorphs: \((hP6-P6\text{}/mcm\)-space group, no. 194) and \((oP6-Pmnn\)-space group, no. 59) are mechanically and dynamically stable;
- these phases are comparably thermodynamically stable but less stable than the known \(hP3-P6/mcm\) phase;
- \(hP6-P6\text{}/mcm\) phase is ductile and \(oP6-Pmnn\) phase is intermediate between brittle and ductile;
- both new phases have a lower hardness than the known \(hP3-P6/mmm\) phase.

We hope that results relating to new hypothetical zirconium diboride (ZrB$_2$) polymorphs will be confirmed by other calculations, as well as through experiments. Knowledge about these new phases can be very useful when doping metal borides with zirconium.

**Supplementary Materials:** The following are available online at http://www.mdpi.com/1996-1944/13/13/3022/s1.

**Author Contributions:** M.M. contributed to the conceptualization, investigation, methodology, analysis, interpretation of results, visualization, and the writing—original draft preparation. T.M. contributed to the conceptualization, investigation, writing—review and editing, funding acquisition, and supervision. All authors have read and agreed to the published version of the manuscript.

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**Conflicts of Interest:** The authors declare no conflict of interest.
Abbreviations
The following abbreviations are used in this manuscript:

- UHTCs: ultra high-temperature ceramics
- DFT: density functional theory
- DFPT: density functional perturbation theory
- PAW: projector augmented-wave
- PP-PW: pseudopotential, plane-wave
- XC: exchange-correlation
- LDA: local density approximation
- GGA: generalized gradient approximation
- PBE: Perdew-Burke-Ernzerhof

References


