Comparison of Functionals for Metal Hexaboride Band Structure Calculations

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Abstract

Density functional calculations of the electronic band structure for superconducting and semi-conducting metal hexaborides are compared using a consistent suite of assumptions and with emphasis on the physical implications of computed models. Spin polarization enhances mathematical accuracy of the functional approximations and adds significant physical meaning to model interpretation. For YB6 and LaB6, differences in alpha and beta projections occur near the Fermi energy. These differences are pronounced for superconducting hexaborides but do not occur for other metal hexaborides.

Keywords

Hexaborides; Density Functional Theory; Band Structure; Spin Polarization; Superconductivity

1. Introduction

The flexibility of the boron framework in metal hexaborides accommodates a wide range of metal atoms or cation charges that results in a diversity of electronic and magnetic properties [1] [2]. We have previously evaluated properties of the hexaboride framework using precise experimental data and density functional theory [3] and recognized that, in general, property trends are defined by bond length adjustments within and between framework atoms.

The metal hexaborides offer an ideal structural system for comparison of computational models with physical properties as well as for validation of theories on bonding. The system is of relatively simple symmetry—cubic with space group Pm\(\overline{3}\)m—and contains only two elements. This simplicity of composition and structure affords
facile computation and interpretation of model outcomes. Depending on the metal ion within the structure, boron accommodates both covalent and ionic bonding to varying degree within and between the $\text{B}_6$ octahedra that comprise the structural framework. Boron-boron and metal-boron bonds clearly influence the electronic properties extant in the hexaboride suite [3] and are explored in more detail in this work.

Electronic properties are generally controlled by the behavior of electrons within a small energy range (~$k_B T$, where $k_B$ is Boltzmann’s constant and $T$ is the absolute temperature) around the Fermi energy [4]-[6]. Electronic band structure calculations for the electrons in the normal state of LaB$_6$ and YB$_6$, two of the metal hexaborides that display superconductivity, attribute predominant d metal character to the electrons at the Fermi level, with large contributions from boron 2p character [7]-[9]. In this paper, we use density functional theory (DFT) to model electron behaviors at the Fermi energy, including all interactions in which the electrons participate, and the characteristics of bonding electrons in metal hexaborides.

Boron bonding in metal hexaborides is complex and integral to a boron framework that is highly responsive to structural variations [3]. This work evaluates the factors that control electron dynamics at the Fermi energy in metal hexaborides, particularly in regions of interest identified in an earlier paper [3]. Model calculations are subjected to a wide range of initial setup conditions, which include more accurate functional approximations to account for exchange interactions and correlations, consideration of the material system as metal or non-metal and evaluation of spin polarization.

2. Calculation Methods

We have performed an extensive comparison of DFT calculations using the CASTEP module [10] of Materials Studio 6.1, with various functionals, applied to a selection of metal hexaborides. Starting with the simplest local density approximation (LDA), a range of functionals with particular intrinsic errors are tested including the generalized gradient approximation (GGA), Hartree-Fock approximation (HF-LDA), the exchange-correlation function by Ceperley and Alder (CP) [11] parameterized by Perdew and Zunger (PZ) [12], screened Hartree-Fock (sX-LDA), the hybrid functional by Becker, Lee, Yang and Parr (B3LYP) and the screened hybrid functional by Heyd, Scuseria and Ernzerhof (HSE06) [13] [14]. All calculations use an ultrafine grid choice and plane-wave (PW) basis, linear response functions, norm-conserving pseudo potentials, Pulay density mixing schemes (when the method allowed, otherwise the program choice is All Bands/EDFT) and fine interpolation methods. The PW basis set cut-off is typically $>290$ eV and the energy cut off is 10 eV. For display of results, the instrument broadening default value of 0.050 eV is retained, as this value is technique and/or instrument specific.

Although the oxidation state of the atom can be specified during parameter input for the crystal structure, results from test calculations that assumed different oxidation states for the atoms did not differ from those for zero oxidation state. Given this result, subsequent calculations reported here assume zero oxidation state for constituent atoms.

For calculations that utilized full geometry optimization, an estimate of the physical accuracy of these first-principles calculations is obtained. These optimizations use as input values the refined unit cell parameters from X-ray or neutron diffraction of well determined hexaboride single crystal studies [3]. For calculations, which do not include optimization of the unit cell in the geometry optimization, lattice parameters determined from refinements of X-ray and/or neutron diffraction data are used [3], but fractional atomic coordinates are optimized in order to compare approximations for the various electronic band structure calculations. For cases where geometry optimization is not carried out, the band structure is calculated in the energy task of the setup, using the full set of structural parameters refined from X-ray and/or neutron diffraction data [3].

Possible effects of spin polarization on the band structure are calculated using the CASTEP program. For this software, spin polarization models require treating the compound as a non-metal. The CASTEP program confers explicit constraints on electronic models depending on the choice of “metal” or “non-metal” configuration. The “metal” option in CASTEP [10] corresponds to a tighter program setting (that is usually associated with the modeling of metals, hence the name “metal”). However, this term as used in the program does not of itself mean that the subject material is a metal [15].

In essence, the number of occupied bands for insulators is calculated as half the total number of valence electrons. However, this approach is not suitable for metals [13]. For metallic systems, partial occupancies are introduced to eliminate discontinuous changes in total energy that are created when an energy band crosses a Fer-
during self consistent field (SCF) minimization [13]. This change in orbital occupancy from fixed to un-fixed automatically introduces smearing of the energy levels as described in detail below. Spin polarization calculations focus on YB₆ and CaB₆, since these compounds represent an electron-doped and a complete orbital configuration, respectively.

Most calculations are completed using Materials Studio 6.1 via the Microsoft Windows mode of a 12 core Mac Pro Xeon 64bit workstation. This workstation operates with an Intel chip and mimics up to 24 cores. When memory requirements exceed the capacity of this computer, calculations are undertaken via the High Performance Computing (HPC) facility at QUT using multiples of 16 cores. The optimum choice for calculations on the HPC facility is 64 cores.

3. Bonding in Metal Hexaborides

The literature shows a range of approaches to analyze and describe boron bonds in metal hexaborides. Some approaches allow easier visualization of electron distribution, for example, while others facilitate model calculations. These broad approaches are briefly reviewed below.

3.1. Molecular Orbital Models

Molecular orbital (MO) models that build the structure as a linear combination of boron atomic orbitals of valence electrons (not including core electrons) using a tight-binding approximation occurs in the early work of Longuet-Higgins and de V. Roberts [16]. A similar tight-binding approach with detailed group theoretical analysis of MO in metal hexaborides is given by Yamazaki [17]. A slightly modified version (normally referred to as the Valence Bond approach) where the inter-octahedral bonds are considered simple localized bonds using sp hybrids from each boron atom, can be found in the book by Urch [18]. Although these models do not embrace the conventional paradigm with ligands coordinated to a metal cation, the lack of a cation implies that the interactions between ligands hold the B₆ cluster together [18]. Both MO models lead to a suite of 18 orbitals, six with primary sp hybrid character (a₁g, t₁u, and e_g) and 12 with p hybrid character (t₁g, t₂g, t₁u, t₂u) involving the 6 borons in an octahedral cluster. Of these 18 orbitals, seven (a₁g, t₁u, and t₂g, where t₁u and t₂g have p hybrid character) have bonding character. This configuration requires 14 electrons to hold the B₆ cluster together (two of which would be donated by the metal atom), and 11 orbitals (e_g, t₁u, t₂u and t₁g) have antibonding character [18].

3.2. Cluster Approach to B₆ Octahedra

While largely pertaining to a MO approach, cluster properties of the boron octahedra [19]-[22] can provide further insight. These structures can be viewed as a metal boron cluster where the stability of cluster assembly [23] is determined by the nature of boron bonding. The quantum states in metal clusters adopt a super shell structure with associated orbitals that mimic the characteristics of atoms and give rise to the concept of a super atom [24] [25]. The quantum confinement in nearly spherically symmetric clusters results in the electronic shell sequence:

\[ 1s^2, 1p^6, 1d^{10}, 2s^2, 1f^{14}, 2p^6, \ldots \]

originally introduced through the concept of a jellium spherical potential [19] [25], that is attractive, constant within a given radius and zero outside that radius [26]. Electrons contributed by the valence orbitals of the atoms in the cluster fill nl orbitals in keeping with the Pauli exclusion principle. However, due to the nature of charge distribution in the clusters, the ordering of these nl levels is different to conventional configurations. Although these electronic shells are introduced via the jellium model, the existence of electronic shells in fermionic systems is known for a far wider range of potentials [25].

Furthermore, when interacting electrons are considered and the octahedral geometry of the cluster is included, a partial breaking of the 1d degeneracy is expected [27]. This loss of degeneracy results in cluster orbitals with the symmetry sequence: a₁g, t₁u, t₂g and e_g, which matches the MO configuration for all the bonding orbitals up to the first antibonding orbital [16] [18]. The t₂g and e_g orbitals of the boron cluster indicate that electrons at the highest occupied molecular orbital (HOMO) and at the lowest unoccupied molecular orbital (LUMO) resemble “d-like” symmetry behavior. Thus, clusters may open detectable gaps via splitting of the shells due to the arrangement of ionic cores similar to crystal field splitting of atomic d states through the electric fields of surrounding ions in solids [24] [27]. As such, Hund’s rules favor high spin states in open shell systems stabilized by
exchange coupling, and similarly, orbital occupation is expected to be determined by a trade-off between Hund’s rules and possible Jahn-Teller effects.

3.3. United Atom Configuration

A complete understanding of MO requires recognition of the atomic orbitals into which they revert at large inter-nuclear distances, and the configuration when the inter-nuclear distance is reduced to zero and the participating nuclei coalesce [22] [28]. The understanding of molecular structure derived from such correlation diagrams parallels that of atomic structure afforded by the periodic table [28].

In the case of metal hexaborides, two united atom conditions can be envisaged: 1) as the atoms move away from the boron octahedral clusters at the vertex and meet at the middle points of the cube sides; and 2) as the atoms coalesce at the cube vertex. The first condition results in a dodecahedral coordination of coalesced B₂ clusters, while the second results in a cubic coordination of coalesced B₆ clusters. In this latter case, the B₆ cluster is equivalent to the super atom approach described above. Note that for metal hexaborides, if the metal is to have d character, the crystal or ligand field splitting goes in the same direction for dodecahedral and for cubic coordinations (see Figure 2.7 in reference [29]). Therefore, the effects of ligands on the metal in a metal hexaboride can be considered as a crystal or ligand field condition intermediate between these two united atom conditions.

4. Earlier Band Structure Calculations

As mentioned above, both Longuet-Higgins and de V. Roberts [16] and Yamazaki [17], use tight binding approximations to describe metal hexaboride band structures. Later, Perkins et al. [30] attribute the difference between their results and those of Yamazaki [17] to inclusion of overlap between Bloch functions of the same symmetry [30]. Furthermore, Perkins et al. [30] found that in three metal hexaboride solids (i.e. Ca, Sr and Ba), the metal atoms transfer 0.9, 1.0 and 0.0 electrons to the B₆ cage in each unit cell, respectively. Thus, they argue that there is no absolute requirement for each B₆ cage to acquire two electrons for stability in the solid-state [30]. According to Perkins et al. [30], in all hexaboride metallic systems, the metal s states partake in bonding that broadens the lowest valence band along the Λ, Σ and Δ directions. These directions are equivalent to GR, GM and GX, respectively, in CASTEP calculations [3]. However, Perkins et al. [30] hold that no significant s orbital contribution is found at the Fermi surface.

In later work, Hasegawa and Yanase [31] use a non-relativistic, symmetrized augmented plane wave (APW) method to calculate the band structure of LaB₆, constructing a one electron potential on the basis of Slater’s Xα method. An interesting point mentioned by Hasegawa and Yanase [31], and also attributed to Longuet-Higgins and de V. Roberts [16], is that the six s states of the B atoms in an octahedron, as well as the six p states of these B atoms, form together d-like orbitals retaining some of the symmetry about the B₆ octahedral centre. These d-like orbitals and the La d state have nearly equal energies in LaB₆, and therefore, strong hybridization occurs. Similarly, Walch et al. [32] examine the predictions of the Hartree-Fock-Slater (HFS) model for the occupied levels and low-energy excitations of LaB₆, adopting the simplest, one-electron direct-transition interpretation of the excitation spectrum. Two possible neutral atom configurations for La, i.e. La: 5d⁰6s² and La: 5d²6s¹ are compared. This work re-emphasizes the importance of the La-B bond, in contrast to earlier work that attributes a predominant, or an unduly large, role for electronic conduction to the La-La metal bond.

For CaB₆, Hasewaga and Yanase [33] use a self-consistent augmented plane wave (APW) method with the local-spin density approximation (LSDA), taking into account relativistic effects in a fashion that eliminates spin-orbit interaction terms from the Hamiltonian. They compare muffin-tin and non-muffin tin corrections and show that these corrections can have a large effect on the value of the semiconducting gap. They determine a gap of 0.3 eV for CaB₆, and suggest this is in good agreement with an experimental result of 0.4 eV from Johnson and Daane [33].

A renewed interest in CaB₆ is triggered by experimental observation of ferromagnetism at high temperature in La-doped forms of CaB₆ [34]-[40]. Many studies on CaB₆ explore whether this hexaboride is a semiconductor or a semimetal [35] [37] [41] [42].

A wide array of experimental techniques is utilized to determine the band gap of a material. In addition to temperature dependence of resistivity by Johnson and Daane [33], angle resolved photoemission spectroscopy (ARPES) [35] [36], resonant inelastic X-ray scattering (RIXS) [35], electron energy loss spectroscopy (EELS) and transport [38], optical absorption and tunneling conductance [43] are reported for metal hexaborides. Table
lists the experimental and theoretical results in the literature for determination of the CaB$_6$ band gap.

To date, few theoretical models for this simple CaB$_6$ structure have been systematically compared with each other or with similar parameter attributes. Thus, it is difficult to evaluate the actual value for the CaB$_6$ band gap, particularly with the wide range of outcomes listed in Table 1. The estimated band gap values show significant variations whether determined experimentally or through DFT calculations.

A systematic comparison of CASTEP models including initial conditions and choice of functional parameters allows identification of potential, non-obvious sources of disparity in theoretical results and effective comparison with experimental data.

A wide range of metal hexaboride band structure calculations within the framework of DFT use the local density approximation (LDA) and the generalized gradient approximation (GGA) [3] [7] [9] [48] [49]. The LDA is exact in the uniform electron density limit and the GGA includes information on the spatial variations in electron density, using functions which are only valid for slowly varying densities [50] [51]. These DFT calculations show similar results on the shapes of the bands and emphasize the GM join as a region in reciprocal space near the Fermi surface of particular interest [3] [9].

The accuracy of the DFT calculations encompasses both a mathematical and a physical context [50] [51]. Views on which functional provides the most accurate results appear to be material system and property specific, evolving with time and undergoing revision on a regular basis [15] [50]-[61]. The transition point where the prescribed accuracy of the functional refers to a molecular orbital or to an extended solid-state material is not always clear.

Recently, Lejaeghere et al. [62] have reviewed the intrinsic and numerical errors for DFT predictions. This statistically based evaluation provides guidelines for the assignment of percentage errors and benchmark comparison of calculated and experimental results for many elements of the periodic table. While this work focuses on elemental crystalline solids, verification of the methodology on multicomponent solids suggests that intrinsic errors for prediction of materials properties are low [62].

5. DFT Model Results

The physical properties (e.g. superconductivity) for which there are unequivocal experimental data on metal hexaborides occur at low temperatures (<10 K). The overarching effect of temperature, particularly with theoretical models based on room temperature structural data such as lattice parameters, places limitations but also reality checks, on the viability of specific models. In this work, we compare model calculations on the basis of

| Year | Experimental Method | Theoretical Model | Estimated Band Gap [eV] | Reference |
|------|---------------------|-------------------|-------------------------|-----------|
| 1963 | T dependent resistivity | - | 0.4 | [33] |
| 1979 | - | LSDA | 0.3 | [44] |
| 2001 | - | GW | 0.8 ± 0.1 | [45] |
| 2002 | ARPES & RIXS | - | >1.0 | [35] |
| 2002 | - | GW | 0.0 | [37] |
| 2003 | ARPES | - | 1.0 | [36] |
| 2004 | EELS | LDA** | 0 | [38] |
| 2004 | e-transport, tunneling | - | 0.2 and 1.0 | [43] |
| 2005 | - | sX-LDA | >1.2 | [46] |
| 2008 | - | QMC | 1.3 | [39] |
| 2008 | - | LSDA | 0.2 | [40] |
| 2009 | - | GGA-HCTH | 0.85 | [47] |

**LDA shows that the valence and conduction bands touch at the X point. However, the instrumental resolution of 1.6 - 1.8 eV in this study may not resolve a gap around 1 eV.
structural parameters determined from room temperature refinements [3]. Experimental data [63] [64] show that structural parameters of hexaborides adjust at low temperatures, and in limited cases [65], can show non-linear step changes in cell dimension at low temperatures. Nevertheless, for the purposes of these model evaluations, comparisons are performed using structural parameters determined at room temperature as this is a reliable reference point.

As noted earlier, the viability and fidelity of specific computational models is evaluated through comparison with well defined, precise experimental data on structural parameters [3] for metal hexaborides. The models used, and the functionals that comprise each model are listed in Tables 2 and 3. The relative fidelity of calculated parameters compared with experimental values (shown as the parameter, \( \delta_0 \) or \( \delta_3 \)) is shown in Tables 2 and 3.

Table 2 lists ten different models utilizing 34 different combinations of functional for the YB\(_6\) structure. Table 2 shows that few combinations of DFT models calculate a cell dimension for YB\(_6\) equivalent to the experimentally determined value. Only the LDA-CA-PZ combination and the B3LYP model generate a cell dimension similar to the experimental value primarily because the cell dimension is not optimized nor refined during the calculation.

For YB\(_6\), within the LDA-CA-PZ set of comparisons, the non-metal, spin polarized option produces structural parameters closest to experimental data with full geometry optimization. If the lattice parameter is not optimized, a similar outcome occurs with selection of a metal configuration and unpolarized spins. Full geometry optimization in the GGA-PBE approximation produces similar outcomes when the YB\(_6\) system is modeled as a non-metal. Models using a non-metal, spin polarized selection provides calculated structural parameters similar to experimentally-determined values.

In all other cases, except for the LDA + U GGA-PBE model without metal configuration, the calculated cell dimension is underestimated compared with the experimental value. Calculation of bond length (B-B\(_{\text{out}}\)) is consistent with experimental values where the parameter is not refined or optimised and, in general, is overestimated for LDA-CA-PZ configurations of DFT models.

In general, not selecting the “metal” option provides closer approximations to experimental values. Calculations with occupancy fixed, not unexpectedly, provide values closer to the experimentally determined cell dimensions. If the calculation includes selection of the “metal” option, the capacity to calculate a band structure with fixed occupancy is not allowed.

Table 3 lists five different models utilizing 31 different combinations of functional for CaB\(_6\). For CaB\(_6\), optimization of the unit cell for the LDA-CA-PZ approximation is sufficient to cause overlap between the conduction and valence bands and to destroy any indication of a band gap. This outcome is due to a reduction in both the lattice parameter and the B-B\(_{\text{out}}\) distance relative to input values. If the input values are not optimized, a small gap of 0.016 eV results. Optimizing the B-B\(_{\text{out}}\) distance without changing the lattice parameter produces a slight increase in the B-B\(_{\text{out}}\) distance relative to the input value, and results in a slightly larger gap of 0.048 eV.

Using the GGA-PBE approximation, the resulting lattice parameter and the B-B\(_{\text{out}}\) distance for CaB\(_6\) do not deviate significantly from the input values. In this case, the band gap is comparable to the largest values of the LDA-CA-PZ approximation (0.039 - 0.048 eV). The use of these functions suggests a modest correlation between the B-B\(_{\text{out}}\) distance and energy gap as noted by Schmitt et al. [66]. Table 3 lists other calculations with band gaps ranging from 0.077 eV to 0.149 eV and between 0.196 eV and 0.394 eV. A significant correlation between the magnitude of the energy gap and B-B\(_{\text{out}}\) value is observed in these calculations.

For CaB\(_6\), Table 3 shows that choice of a spin polarized calculation is not significant for band gap estimation compared with the spin unpolarized calculation with less accurate functionals (e.g., the LDA-CA-PZ, GGA-PBE and sX-LDA-CA-PZ choices in Table 3). With a more accurate functional, such as the HSE06 [13] [14], a difference of ~0.070 eV occurs between the polarized and non-polarized calculations. This suggests that spin polarization effects are of the same order of energy as the energy of characteristic phonons of metal hexaborides [67] [68]. Hence, coupling between electron spin polarization and phonons in this system is probable.

A lack of, or limited, spin polarization dependence largely originates from the fact that CaB\(_6\) contains an even number of electrons. The computational protocol for CASTEP calculations is that the number of orbitals is determined as the number of electrons divided by two, with sequential pairs of electrons within available orbitals [13], so that no unpaired electron remains in a system with an even number of electrons.

The above computational protocol is confirmed by similar calculations using YB\(_6\) super cells (data not shown). With an even number of super cells, such as 2 \( \times \) 2 \( \times \) 2 or 4 \( \times \) 4 \( \times \) 4, spin polarization dependence is lost, while
The table below shows the model conditions used for electronic band structure calculations for YB₆ (input experimental parameters \( a = 4.1000 \) Å and B-B_{out} = 1.6300 Å [3]).

| Selected Function | Metal (Y/N) | Spin Polar’n (Y/N) | SCF Minimizer (Occ: Y/N) | Optimize (Y/N) | Calculated Parameters [Å] |
|-------------------|-------------|--------------------|--------------------------|----------------|---------------------------|
|                   |             |                    |                          |                | B-B_{out} | \( \delta_{B} \) | \( a \) | \( \delta_{a} \) |
| LDA-CA-PZ         | Yes         | Yes                | DM—Yes                   | Yes            | 1.6022  | 0.0278  | 4.0334 | 0.0666 |
| LDA-CA-PZ         | Yes         | Yes                | AB—No                    | Yes            | 1.6020  | 0.0280  | 4.0304 | 0.0696 |
| LDA-CA-PZ         | No          | No                 | DM—Yes                   | Yes            | 1.6180  | 0.0120  | 4.0549 | 0.0451 |
| LDA-CA-PZ         | No          | Yes                | DM—Yes                   | Yes            | 1.6210  | 0.0090  | 4.0571 | 0.0429 |
| LDA-CA-PZ         | No          | Yes                | AB—Yes                   | Yes            | 1.6210  | 0.0090  | 4.0571 | 0.0429 |
| LDA-CA-PZ         | No          | Yes                | AB—No                    | Yes            | 1.6022  | 0.0278  | 4.0307 | 0.0693 |
| LDA-CA-PZ         | No          | No                 | DM—Yes                   | Yes            | 1.6180  | 0.0120  | 4.0549 | 0.0451 |
| LDA-CA-PZ         | No          | Yes                | AB—Yes                   | No cell        | 1.6424  | −0.0124 | 4.1000 | 0.0000 |
| LDA-CA-PZ         | No          | No                 | AB—Yes                   | No cell        | 1.6359  | −0.0059 | 4.1000 | 0.0000 |
| LDA-CA-PZ         | No          | Yes                | AB—Yes                   | No cell        | 1.6401  | −0.0101 | 4.1000 | 0.0000 |
| LDA-CA-PZ         | No          | Yes                | AB—Yes                   | No cell        | 1.6425  | −0.0125 | 4.1000 | 0.0000 |
| LDA-CA-PZ         | Yes         | Yes                | AB—No                    | No at all      | 1.6302  | −0.0002 | 4.1000 | 0.0000 |
| LDA-CA-PZ         | Yes         | No                 | AB—No                    | No at all      | 1.6300  | 0.0000  | 4.1000 | 0.0000 |
| LDA-CA-PZ         | No          | No                 | AB—Yes                   | No at all      | 1.6300  | 0.0000  | 4.1000 | 0.0000 |
| LDA-CA-PZ         | No          | Yes                | AB—Yes                   | No at all      | 1.6302  | −0.0002 | 4.1000 | 0.0000 |
| GGA-PBE           | Yes         | No                 | DM—Yes                   | Yes            | 1.6151  | 0.0149  | 4.0634 | 0.0366 |
| GGA-PBE           | No          | No                 | DM—Yes                   | Yes            | 1.6151  | 0.0149  | 4.0634 | 0.0366 |
| GGA-PBE           | No          | Yes                | DM—Yes                   | No cell        | 1.6302  | −0.0002 | 4.0871 | 0.0129 |
| LDA + U-CA-PZ     | No          | Yes                | DM—Yes                   | Yes            | 1.6307  | −0.0007 | 4.0756 | 0.0244 |
| LDA + U-CA-PZ     | Yes         | Yes                | DM—Yes                   | Yes            | 1.6068  | 0.0232  | 4.0392 | 0.0608 |
| LDA + U GGA-PBE   | No          | Yes                | DM—Yes                   | Yes            | 1.6484  | −0.0184 | 4.1157 | -0.0157 |
| LDA + U GGA-PBE   | No          | Yes                | DM—No                    | Yes            | 1.6204  | 0.0096  | 4.0793 | 0.0207 |
| LDA + U GGA-PBE   | Yes         | Yes                | AB—No                    | Yes            | 1.6280  | 0.0020  | 4.0870 | 0.0130 |
| LDA + U GGA-PBE   | No          | Yes                | AB—No                    | No cell        | 1.6204  | 0.0096  | 4.0794 | 0.0206 |
| HF-LDA-CA-PZ      | No          | No                 | AB—Yes                   | No cell        | 1.6559  | −0.0259 | 4.1000 | 0.0000 |
| HF-LDA-CA-PZ      | No          | Yes                | AB—Yes                   | No cell        | 1.6566  | −0.0266 | 4.1000 | 0.0000 |
| sX-LDA-CA-PZ      | No          | No                 | AB—Yes                   | Yes            | 1.6291  | 0.0009  | 4.0791 | 0.0209 |
| sX-LDA-CA-PZ      | No          | Yes                | AB—Yes                   | Yes            | 1.6352  | −0.0052 | 4.0845 | 0.0155 |
| B3LYP             | No          | No                 | AB—Yes                   | No cell        | 1.6404  | −0.0104 | 4.1000 | 0.0000 |
| B3LYP             | No          | Yes                | AB—Yes                   | No cell        | 1.6446  | −0.0146 | 4.1000 | 0.0000 |
| B3LYP             | No          | No                 | AB—Yes                   | No at all      | 1.6302  | −0.0002 | 4.1000 | 0.0000 |
| HSE06             | No          | No                 | AB—Yes                   | Yes            | 1.6215  | 0.0085  | 4.0569 | 0.0431 |
| HSE06             | No          | Yes                | AB—Yes                   | Yes            | 1.6360  | −0.0060 | 4.0830 | 0.0170 |
Table 3. Model conditions used for band structure calculations for CaB₆ (input lattice parameter: \( a = 4.1514 \text{ Å} \) and B-Bout = 1.6760 Å [3]).

| Selected Function | Metal (Y/N) | Spin Polariz'n (Y/N) | Band Gap (eV) | Parameter Optimiz'n (Y/N) | Calculated Parameters [Å] |
|-------------------|-------------|----------------------|---------------|--------------------------|--------------------------|
|                   |             |                      |               |                          | B-Bout \( \delta_B \) a \( \delta_a \) |
| LDA-CA-PZ         | Yes         | No                   | 0             | Yes                      | 1.6465 0.0295 4.0811 0.0703 |
| LDA-CA-PZ         | No          | No                   | 0             | Yes                      | 1.6453 0.0307 4.0783 0.0731 |
| LDA-CA-PZ         | No          | Yes                  | 0.048         | No cell                  | 1.6829 −0.0069 4.1514 0.0000 |
| LDA-CA-PZ         | No          | Yes                  | 0.048         | No cell                  | 1.6829 −0.0069 4.1514 0.0000 |
| LDA-CA-PZ         | No          | No                   | 0.016         | No at all                | 1.6760 0.0000 4.1514 0.0000 |
| LDA-CA-PZ         | No          | Yes                  | 0.016         | No at all                | 1.6760 0.0000 4.1514 0.0000 |
| LDA-CA-PZ         | No          | No                   | 0.048         | No cell                  | 1.6829 −0.0069 4.1514 0.0000 |
| LDA-CA-PZ         | No          | Yes                  | 0.048         | No cell                  | 1.6829 −0.0069 4.1514 0.0000 |
| GGA-PBE           | Yes         | No                   | 0.043         | Yes                      | 1.6749 0.0011 4.1469 0.0045 |
| GGA-PBE           | No          | No                   | 0.043         | Yes                      | 1.6730 0.0030 4.1459 0.0055 |
| GGA-PBE           | No          | Yes                  | 0.048         | Yes                      | 1.6730 0.0030 4.1459 0.0055 |
| GGA-PBE           | No          | No                   | 0.043         | No cell                  | 1.6756 0.0004 4.1514 0.0000 |
| GGA-PBE           | No          | Yes                  | 0.041         | No cell                  | 1.6759 0.0001 4.1514 0.0000 |
| GGA-PBE           | No          | No                   | 0.039         | No at all                | 1.6760 0.0000 4.1514 0.0000 |
| GGA-PBE           | No          | Yes                  | 0.039         | No at all                | 1.6760 0.0000 4.1514 0.0000 |
| sX-LDA-CAPZ       | No          | No                   | 0.077         | Yes                      | 1.6637 0.0123 4.1184 0.0330 |
| sX-LDA-CAPZ       | No          | Yes                  | 0.078         | Yes                      | 1.6635 0.0125 4.1184 0.0330 |
| sX-LDA-CAPZ       | No          | No                   | 0.149         | No cell                  | 1.6803 −0.0043 4.1514 0.0000 |
| sX-LDA-CAPZ       | No          | Yes                  | 0.149         | No cell                  | 1.6803 −0.0043 4.1514 0.0000 |
| sX-LDA-CAPZ       | No          | No                   | 0.107         | No at all                | 1.6760 0.0000 4.1514 0.0000 |
| sX-LDA-CAPZ       | No          | Yes                  | 0.107         | No at all                | 1.6760 0.0000 4.1514 0.0000 |
| HSE06             | No          | No                   | 0.196         | Yes                      | 1.6381 0.0379 4.0632 0.0882 |
| HSE06-01          | No          | Yes                  | 0.274         | Yes                      | 1.6414 0.0346 4.0735 0.0779 |
| HSE06-02          | No          | Yes                  | 0.266         | Yes                      | 1.6409 0.0351 4.0737 0.0777 |
| HSE06             | No          | No                   | 0.394         | No cell                  | 1.6833 −0.0073 4.1514 0.0000 |
| HSE06             | No          | Yes                  | 0.476         | No cell                  | 1.6842 −0.0082 4.1514 0.0000 |
| HSE06             | No          | No                   | 0.324         | No at all                | 1.6760 0.0000 4.1514 0.0000 |
| HSE06             | No          | Yes                  | 0.394         | No at all                | 1.6760 0.0000 4.1514 0.0000 |
| B3LYP*            | No          | No                   | 0.871         | No cell                  | 1.6807 −0.0047 4.1514 0.0000 |
| B3LYP*            | No          | Yes                  | 0.827         | No cell                  | 1.6806 −0.0046 4.1514 0.0000 |
| B3LYP*            | No          | No                   | 0.827         | No at all                | 1.6760 0.0000 4.1514 0.0000 |
| B3LYP*            | No          | Yes                  | 0.871         | No at all                | 1.6760 0.0000 4.1514 0.0000 |
spin polarization dependence is apparent with an odd number of super cells such as $3 \times 3 \times 3$. Other evaluations of spin polarized super cell calculations (e.g. via sX-LDA and HSE06), particularly with an ultra-fine grid, did not converge.

5.1. Metals or Insulators—Electron Smearing

CASTEP calculations differentiate between metals and insulators on the basis of occupied bands at different k-points in the Brillouin zone. For insulators, the number of occupied bands is calculated as one half the total number of valence electrons. As noted above, these options for CASTEP calculations introduce the concept of electron energy smearing which is computationally possible between 0.01 eV and 10 eV. These energy values correspond to temperatures of $\sim 116^\circ$K and $\sim 116,000^\circ$K, respectively (using $\text{Energy} \approx k_B T$, where $k_B$ is Boltzmann’s constant). In terms of temperature equivalence, these energies are excessively large for phenomena that may be dominated by low temperature interactions and couplings active at a few degrees Kelvin. For instance, the superconducting transition temperature for YB$_6$ is 7.1 K [69] [70].

The smearing default values in CASTEP are 0.1 eV and 0.2 eV, respectively, for orbital occupancy and for display of the density of states (DOS). All default values in CASTEP are a reasonable average choice for common modeling conditions, although caution is advised with respect to the selection of smearing values [13]. However, comparable values around 0.1 eV to 0.2 eV are the preferred choice when the modeling intent is to remove potential local mathematical instabilities at the Fermi energy. Unfortunately, it is not always apparent that such choices may force the modeled system to undergo an unrealistic, non-physical, equivalent temperature condition for the property of interest.

For example, in a system close to a Mott metal-insulator transition, small energy pseudo gaps of tens of meV (or even smaller gaps), may play an important role [71]. Thus, smearing of orbitals with a 0.1 eV or 0.2 eV value may sacrifice, unnecessarily and detrimentally, important physical information about the system. Furthermore, all equivalent temperature effects from smearing are likely to introduce a non-directional random character, which may not always correspond to the limited and most intense phonon modes. These phonon modes will dominate system behavior at low temperatures. To minimize this issue, we have chosen a fine interpolation for the display of DOS.

Similar effects are evident for model calculations with the metal option selected in CASTEP calculations. Typically, the standard default value of 0.1 eV overrides spin polarized results in metal hexaboride systems with an odd number of electrons. This outcome indicates that the energy differences related to spin dependence are smaller than 0.1 eV. Recovery of spin polarization is apparent for the lower limit of smearing at 0.01 eV. However, as noted above, this value may be too high for physically meaningful temperature equivalence. If the “no metal” option is selected without fixed occupancy while using the SCF electronic minimizer setup, the outcome is similar to selection of a metal due to the introduction of electron energy smearing.

Figure 1 shows the effects on band structure calculations for YB$_6$ with different model options chosen for “metal” or “no metal” and for choice of fixed occupancy. A shift of about 0.25 eV relative to the Fermi energy in all bands and DOS is clearly evident for these two model choices. The shift in band structure using the LDA-CA-PZ function illustrates the importance of appropriate model selection.

The recent review article by Lejaeghere et al. [62] emphasizes that, in general, comparison of DFT calculations with experimental results requires that the experimental results should refer to 0 K and that measurements are corrected for zero-point vibration effects, which are not present in standard DFT calculations. Alternatively, DFT calculations should incorporate interactions and phenomena that appear and vary as temperature changes.

5.2. Spin Polarization

Calculation trends from this work on hexaborides indicate that inclusion of spin polarization produces more accurate lattice parameter and B-B_{out} values. When the spin polarized option is chosen for CASTEP calculations, different wave functions for different spins are used [13]. If a calculation without spin polarization is chosen, the same orbitals for alpha and beta spins are used. Thus, spin polarization effectively results in an increased number of parameters for the calculation, which may lead to a better mathematical approximation, and to a better quantum mechanical approximation provided assumptions about spins in the system reflect physical reality.

Spin-polarized CASTEP calculations are normally carried out with a variable magnetic moment [13]. How-
Figure 1. Calculated band structures for YB$_6$ using the LDA-CA-PZ functional; assumes no geometrical optimization and no spin polarization with (a) “metal” option and (b) “non-metal” option. Note that the energy bands adjacent to the Fermi surface along the GM join and the DOS are shifted by about 0.25 eV in (b) compared to (a).

However, this procedure does not guarantee that the lowest energy state is determined as DFT solutions can converge to a variety of local minima related to metastable states. The most common solutions correspond to high spin and low spin states. The state generated by CASTEP calculations depends on the initial magnetic configuration, since the solution is likely to converge to the nearest local minimum rather than to a global minimum. Initial magnetic moments on atoms are treated correctly by CASTEP only when the density mixing (DM) scheme is used for electronic minimization [64].

However, our comparison of the DM and the all bands (AB) schemes for electronic minimization revealed
similar results for the spin polarized bands of YB\textsubscript{6}. When there is one unpaired electron in the unit cell, the risk of convergence to multiple local minima is reduced. This outcome may not occur if more than one electron is unpaired as may be for CaB\textsubscript{6} if, for example, two electrons are assumed unpaired.

**Figure 2** shows examples of YB\textsubscript{6} band structure calculations that include spin polarization using the B3LYP functional. The band structures in Figure 3 for YB\textsubscript{6} are calculated using the HSE06 functional. Both of these functionals are considered accurate choices \cite{13} \cite{14} \cite{50}-\cite{61} for quantum mechanical calculations. **Figure 3(a)** also compares the calculation for unpolarized spins with the HSE06 functional.

Although both functionals lead to somewhat different lattice parameters and B-B\textsubscript{out} distances, the qualitative effects on the band structures adjacent to the Fermi energy are quite similar. The band splits at the RG and GM sections, and results in a strong alpha component close to the Fermi energy with another component moving to higher energies in sections adjacent to G. The same alpha band also develops sections along the R direction much closer to the Fermi energy.

Although the amounts of band splitting or the proximity to the Fermi energy along the R direction are not always identical, this phenomenon is qualitatively the same for many non-metal, spin polarized calculations with different functionals. This reproducibility of trends for the bands and DOS combined with optimized parameters close to the experimental input values suggest that spin polarization is a physical reality for YB\textsubscript{6} at low temperatures. We expect that in a single electron approximation for YB\textsubscript{6}, the compound requires an amount of spin in order to participate in coupling for superconductivity and to produce perfect diamagnetic Meissner effects.

In order to extract additional physical information from this spin polarization dependence, we display various DOS components for YB\textsubscript{6} and LaB\textsubscript{6}. **Figures 4** and **5** show the p and d spin components and the alpha and beta components calculated for YB\textsubscript{6} and LaB\textsubscript{6}, respectively. YB\textsubscript{6} displays more pronounced differences adjacent to the Fermi energy than LaB\textsubscript{6}. These differences are indicative of a requirement for stronger coupling in YB\textsubscript{6} compared to that in LaB\textsubscript{6}, likely relating to superconductivity, and the correspondingly higher $T_c$ for YB\textsubscript{6} compared to LaB\textsubscript{6}.

While there may be minor inadequacies with assumptions in DFT calculations, the accuracy of results is improved using spin polarization. When this increased accuracy of model calculation is combined with a physical interpretation of a magnetic contribution to the hexaboride structure, differentiation of YB\textsubscript{6} and LaB\textsubscript{6} properties is evident. In this case, we indicate that spin polarization may be crucial to the correct determination and interpretation of electronic band structure for superconducting (and other magnetic) materials, particularly at low temperatures.

### 6. Conclusions

The electronic band structures of metal hexaborides are calculated for a range of models using a variety of density functionals. This systematic evaluation of band structure calculations for metal hexaborides identifies factors that account for the disparity of previously reported theoretical values.
Figure 3. Calculated band structures and DOS using HSE06 function, assumes “no metal” with geometrical optimization with (a) no spin polarization and (b) spin polarization for YB₆. Both calculations use the same structural input parameters. (c) Calculated band structures and DOS for LaB₆ using the B3LYP function, assumes “no metal” with geometric optimization and spin polarization.

The mathematical procedure used to avoid discontinuities when an energy band crosses the Fermi energy during SCF minimization gives rise to flexible orbital occupation accompanied by smearing of energy levels in model calculations. Such smearing is equivalent to unrealistically high temperatures which may mask fine band and DOS detail related to physical phenomena at very low temperature (i.e. near 0°K). For instance, effects related to coupling for low-temperature superconductivity or a Mott metal-insulator transition may not be detected for materials with gaps or pseudo gaps of a few meVs.

Consideration of spin polarization improves the numerical accuracy and the physical interpretation of associated phenomena. Calculated spin polarization effects for metal hexaborides are typically concentrated adjacent to the Fermi energy and thus, affect electronic properties. For YB₆ and LaB₆, unpolarized bands are split into (a) a large alpha component that approaches the Fermi energy in large regions of reciprocal space, and (b) another
Figure 4. Partial DOS for YB$_6$ using the HSE06 function, for “non-metal” with geometric optimization and spin polarization: (a) p (blue line) and d spin components, (b) alpha (blue line) and beta components. Note that the d spin projection is larger in magnitude than the p spin projection near the Fermi energy. The asymmetry in (b) around the Fermi energy is responsible for a net spin.

Figure 5. Partial DOS for LaB$_6$ using the B3LYP function, assuming “non-metal” with geometric optimization and spin polarization: (a) p (blue line) and d spin components, (b) alpha (blue line) and beta components. Note that the p spin and d projections are similar for LaB$_6$ compared with data for YB$_6$ shown in Figure 4.
alpha component that trends away from the Fermi energy particularly near the G section. Slight differences in alpha and beta projection in the DOS for YB₆ and LaB₆ may correlate to different superconducting transition temperatures.

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