

Boron-Doped Graphite

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Theory and Practice: Bulk Synthesis of C₃B and its H₂- and Li-Storage Capacity**

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In memory of Kenneth Wade

Abstract: Previous theoretical studies of C_3B have suggested that boron-doped graphite is a promising H_2 - and Li-storage material, with large maximum capacities. These characteristics could lead to exciting applications as a lightweight H₂-storage material for automotive engines and as an anode in a new generation of batteries. However, for these applications to be realized a synthetic route to bulk C_3B must be developed. Here we show the thermolysis of a single-source precursor (1,3- $(BBr_2)_2C_6H_4$) to produce graphitic C_3B , thus allowing the characteristics of this elusive material to be tested for the first time. C₃B was found to be compositionally uniform but turbostratically disordered. Contrary to theoretical expectations, the H₂- and Li-storage capacities are lower than anticipated, results that can partially be explained by the disordered nature of the material. This work suggests that to model the properties of graphitic materials more realistically, the possibility of disorder must be considered.

he development of practical and efficient chemical and electrical power storage systems is particularly important with regard to mobile applications such as sustainable automotive transport and consumer electronics.^[1-3] Safe and recyclable H₂-[4] and Li-storage^[5] systems which have high energy-toweight ratios are at the cutting edge of materials and engineering design. The boron carbide C₃B (Scheme 1) has been of great interest to materials scientists and physicists since it was first reported over a quarter of a century ago, [6] owing to the presence of electron-poor, substitutional boron in the parent lattice of graphite, which results in increased

BBr₂ 800°C -HBr BBr₂ 1,3-bis-(dibromoboryl) benzene C₃B

Scheme 1. The synthetic route to the formation of idealized C₂B. The structure shown here is that proposed by Bartlett and co-workers. [6]

conductivity.^[7] A recent resurgence of interest in this material has resulted from theoretical evidence that bulk samples of C₃B may be a viable, high-capacity H₂-storage material at moderate temperatures and pressures^[8,9] (up to 6.1 wt %, over the US Department of Energy storage target). [10] In addition, calculations have shown that C₃B may have a Li capacity of 857 mAhg⁻¹, over twice that of graphite, the anode material commercially used in batteries (372 mA h g⁻¹).^[11,12] Crucially, however, the projected H₂-storage properties of C₃B are reliant on the presence of multilayered, bulk samples. Calculations reveal that the most likely mechanism of H₂ absorption involves chemisorption of H₂ progressing by

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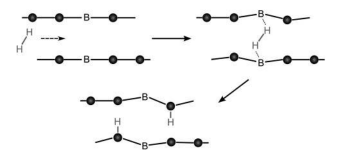
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Scheme 2. Process of H_2 intercalation followed by boron-assisted activation and C-H bond formation. Black spheres represent carbon.

homolytic, boron-assisted cleavage of the H–H bonds with the ultimate formation of two C–H bonds (Scheme 2) with a low kinetic barrier for the initial H₂ intercalation. Significantly, because the pathway involves interlayer activation of H₂ molecules, pure graphite and discrete C₃B monolayers are predicted to show no hydrogen sorption. [8] In addition, the theoretical Li capacity of bulk C₃B (as C₃BLi_{1.5}) is predicted to be higher than that of the monolayer. [12]

To date only microgram quantities of bulk C₃B have ever been prepared because of the inherent slowness and poor scalability of the existing chemical method used for its preparation: chemical vapor deposition (CVD) of BCl₃ and C₆H₆. ^[13,14] While a standard chemical route involving the Wurtz coupling reaction of hexachlorobenzene (C₆Cl₆) or tetrachloroethylene (Cl₂C=CCl₂) and BCl₃ with sodium metal in high boiling-point solvents has been used to produce boron-enriched graphite on the gram scale, the level of boron inclusion in this material (ca. 7.3 at.%) falls well short of that found in C₃B (25 at.%); in addition, high levels of oxidized species are present in the material. ^[15] It should be noted that the maximum boron content that can be obtained because of the solubility of boron in graphite is a mere 2.3%, and requires temperatures in excess of 2300 °C. ^[16]

We show here that the thermolysis of the simple aromatic boron compound 1,3-bis(dibromoboryl)benzene (1,3-(BBr₂)₂C₆H₄, 1; Scheme 1),^[17] which contains the correct 3:1 C/B stoichiometry, produces bulk samples of C₃B quantitatively, a reaction that can be scaled up to the gram scale. This "tiling" synthetic approach is a strategy which has previously shown potential in the synthesis of nitrogen-doped graphitic systems,^[18] but is the first known single-source approach to C₃B. This has enabled the first experimental studies of the H₂-and Li-storage capacity of C₃B and a fundamentally important test of theory.

A previous report on the carbonization of o-phenylenediamine $(1,2\text{-}(NH_2)_2C_6H_4)$ at $800\,^{\circ}\text{C}$ to produce the related graphitic material (C_3N) had shown the formation of two distinct materials, flakes and multiwalled graphite microspheres. The formation of similar graphite microspheres was noted in our initial investigations on the thermolysis of 1 under these conditions (see Section S3 in the Supporting Information). However, optimization of the heating profile enabled the selective synthesis of C_3B flakes. The surface morphology of flakes of C_3B is seen in Figure 1 a, which shows a scanning electron microscopy (SEM) image of flakes which

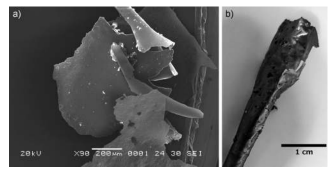


Figure 1. a) SEM image of C_3B typical flakes. b) Photograph of the C_3B material.

have typical thicknesses of 1 to 3 μ m and homogeneous sample make-up (see Section S4 in the Supporting Information). Macroscopically, these flakes of C₃B have a metallic luster (Figure 1b). The curling of the flakes has been put down to their formation through the deposition of graphitic material onto the inside walls of a curved reaction vessel, thereby leading to a curl when the flakes detach from the walls.

The material was further characterized using Raman spectroscopy, powder X-ray diffraction (pXRD), X-ray photoelectron spectroscopy (XPS), solid-state ¹¹B NMR spectroscopy, and laser-desorption/ionization time of flight mass spectrometry (LDI-TOF MS). These techniques show that boron substitution of the samples of C₃B is compositionally uniform within the bulk but that the graphitic structure is turbostratically disordered.

The graphitic nature of the material was shown by pXRD and Raman spectroscopy. The pXRD pattern (Figure 2) shows two main, broad reflections at $\theta = 26^{\circ}$ (002) and 43°

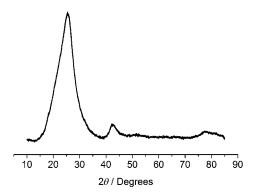


Figure 2. XRD of C₃B flakes. The broad peaks at 26,° 42°, and 78° correspond to the (002), (100), and (110) planes of graphite.

(100), as well as a weaker reflection at 78° (110), thus indicating that the interlayer spacing and mean C–C/C–B bond length in C_3B are similar to the corresponding parameters in graphite. The broadness of the peaks is attributed to the turbostratic nature of the material, in which the layers do not exhibit periodic ordering with respect to one another; the asymmetry of the (002) reflection is ascribed to fluctuations in the interlayer spacing due to curvature in C_3B sheets as



prepared. The presence of disorder is consistent with previous DFT calculations on the packing of C_3B , which indicate that the different packing alternatives of the layers have very similar energies (no more than 2.1 kJ mol^{-1} between them). [19,20] Figure 3 shows the Raman spectrum of C_3B and

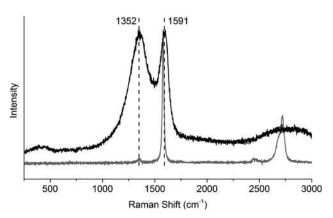


Figure 3. Raman spectrum of C_3B (black) and graphite (gray). The D, G, and 2D bands are seen at 1352, 1591, and 2720 cm⁻¹, respectively

a sample of pristine graphite for comparison. The decrease in the $I_{\rm 2D}/I_{\rm G}$ ratio and increase in intensity of the D band (i.e. an increased $I_{\rm D}/I_{\rm G}$ ratio) are both consistent with high dopant levels as well as the turbostratic nature of the sample.

Compositional analysis was performed using XPS. As XPS is a surface analysis technique, Ar ion beam etching was utilized to enable analysis through the depth of the material. Figure 4 shows how the elemental composition changes with

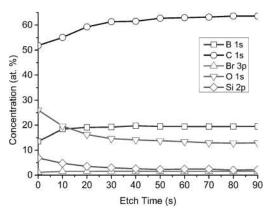


Figure 4. Elemental composition of C₃B flakes as a function of depth (achieved by increasing Ar ion etch time) determined by XPS analysis.

depth. The presence of Si can be attributed to the surface contamination with silica, resulting from the quartz tubes used in the synthesis. Consistent with this conclusion, the levels of Si and O fall off sharply with increased depth. XPS analysis of samples treated with HF show that this contamination is completely removed. The apparently consistently high level of oxygen present in samples, even at depth, is attributed to adventitious (background) contamination (see

the Supporting Information for further discussion). ^[21] The most important observation from XPS measurements is that after etching below the surface layers of the sample (with a surface C/B ratio of 3.8:1), the C/B ratio is maintained between 3.0:1 and 3.3:1. This illustrates that the bulk of the sample of C_3B is compositionally uniform.

MALDI-TOF mass spectrometry assisted further in characterization and provided further evidence of the homogeneity of the samples. Signals corresponding to C_xB_y are present in the spectra, with signals associated with $C_{(x-1)}B_1$ and $C_{(x-2)}B_2$ being more intense than the corresponding C_x signals (see Figure S4 in the Supporting Information). The mass spectra, therefore, support the conclusion that B is covalently bound and distributed homogenously throughout the structure, rather than being present in discrete boron-containing regions. A further important observation is seen from Figure 5, which shows a comparison between the

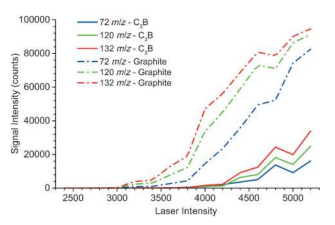


Figure 5. Signal intensity as a function of MALDI-TOF MS power, showing how a much greater laser power is required to achieve flight for fragments from C₃B compared to graphite.

observed signal intensity for a selection of mass fragments with respect to laser power for both a C₃B and graphite substrate. Despite the chosen mass fragments being based solely on carbon, higher laser powers are required for their generation in C₃B, which implies incorporation of a heteroatom (in this case boron), which must first be broken away, in the underlying structure.

Solid-state ¹¹B NMR spectroscopy provides fundamental support for the substitutional incorporation of boron into the structure of C_3B (shown in Scheme 1, right). First principles calculations of the ¹¹B NMR parameters (the quadrupolar coupling parameter C_Q and the anisotropy parameter η_Q) was performed for boron sites in model C_3B structures to enable comparison to experimental data. For all structures analyzed, the calculated values remain similar with $C_Q \approx 5.3$ MHz and $\eta_Q \approx 0$ (see the Supporting Information for the calculated values and structures). The major component (blue line) in the ¹¹B echo magic angle spinning (MAS) experiment (Figure 6) at 43 ppm has C_Q and η_Q values similar to that calculated for C_3B . The minor components (red lines) are ascribed to boron oxide (see Section S7 in the Supporting Information) impurities. The presence of the dominant



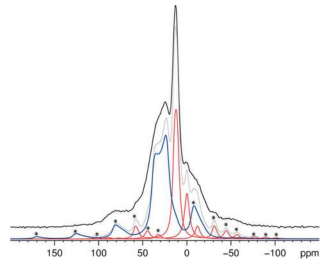


Figure 6. ¹¹B echo ssNMR spectrum acquired at 16.4 T at 10 kHz MAS. The red lines correspond to signals seen in the MQMAS experiment. The blue line was fitted to account for the additional intensity in the spectrum and is assigned to C₃B based on its calculated quadrupolar parameters (further details are given in the Supporting Information). Spinning side bands are marked with an asterisk.

resonance at 43 ppm suggests that boron is present in a substitutional manner in the graphitic lattice.

With well-characterized bulk samples of C_3B in hand, we were next able to assess its H_2 - and Li-storage characteristics. Hydrogen-storage measurements were performed using a modified Sieverts apparatus. Disappointingly, no significant H_2 uptake was observed at 77 K or at 295 K between 0 and 20 bar. To test C_3B as a potential lithium ion battery electrode material, coin cells were produced using powdered C_3B . Using a constant current set up, the capacity of the cell was tested over multiple cycles. The capacity achieved during the first charge exceeds $700 \, \text{mAh g}^{-1}$ and on subsequent cycles a reversible capacity of $374 \, \text{mAh g}^{-1}$ was observed (Figure 7). The reversible capacity closely matches that of

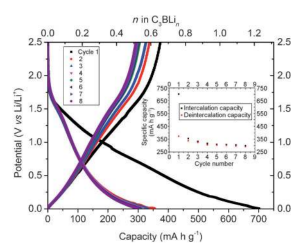


Figure 7. Change in potential during the initial discharge and charge cycles. The high capacity seen during the first discharge is irreversible. The insert shows the capacity versus cycle number.

crystalline graphite (C₃BLi_{0.65} cf. C₆Li in graphite). The capacity of the first discharge approaches the theoretical maximum value (corresponding to a formula of C₃BLi_{1.25} versus the maximum of C₃BLi_{1.5}). One possibility is, therefore, that the cell is cycling reversibly between C₃BLi_{0.65} and C₃BLi_{1.25}. Alternatively, however, the irreversible capacity in the first cycle may merely correspond to the formation of a solid-electrolyte interphase layer (SEI).

In conclusion we have reported the first bulk synthesis of the heavily doped graphite C₃B. This has been fully characterized, both compositionally and structurally, as a turbostratic graphitic phase in which boron has substitutionally replaced carbon. With the production of bulk samples we have been able, for the first time, to test previous theoretical predictions of the materials properties of C₃B. Contrary to theory, we find that negligible H₂ storage occurs. This, and the lower than expected operational Li-storage capacity in a battery, can be traced to a combination of the disorder of the real structure and potentially the adsorption kinetics associated with uptake in micrometer-sized particles. In regard to H₂ adsorption (see Scheme 2) for example, adjacent boron atoms in a perfect AB stacked structure are required for H₂ activation. In addition, the maximum theoretical capacity for lithium intercalation into C₃B requires both a highly ordered AB stacking and an ability to convert into the AA stack. The real material instead has been shown to contain randomly orientated layers. Our study suggests that to model the properties of graphitic materials more realistically, the possibility of disorder in particular must be considered.

Keywords: boron \cdot graphite \cdot H $_2$ storage \cdot Li battery \cdot synthetic methods

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- [21] This was confirmed by XPS control experiments in which commercial (Aldrich) graphite was analyzed and similar levels of oxygen were observed.

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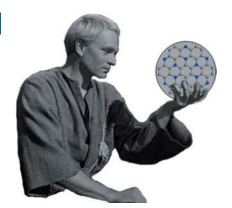
Communications



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Theory and Practice: Bulk Synthesis of C_3B and its H_2 - and Li-Storage Capacity



C₃B or not C₃B that is the question—

Previous theoretical predictions suggest that the stoichiometrically doped graphite C_3B should be a promising H_2 - and Listorage material, with potentially important applications in power storage. The first experimental measurements of the real material indicate, however, that disorder is a limiting factor for the use of C_3B as an energy storage material.