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# Halide-free synthesis of metastable graphitic BC<sub>3</sub>†

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Layered BC $_3$ , a metastable phase within the binary boron–carbon system that is composed of graphite-like sheets with hexagonally symmetric  $C_6B_6$  units, has never been successfully crystallized. Instead, poorly-crystalline BC $_3$ -like materials with significant stacking disorder have been isolated, based on the copyrolysis of a boron trihalide precursor with benzene at around 800 °C. The halide leaving group (–X) is a significant driving force of these reactions, but the subsequent evolution of gaseous HX species at such high temperatures hampers their scaling up and also prohibits their further use in the presence of hard-casting templates such as ordered silicates. Herein, we report a novel halide-free synthesis route to turbostratic BC $_3$  with long-range in-plane ordering, as evidenced by multi-wavelength Raman spectroscopy. Judicious pairing of the two molecular precursors is crucial to achieving B–C bond formation and preventing phase-segregation into the thermodynamically favored products. A simple computational method used herein to evaluate the compatibility of bottom-up molecular precursors can be generalized to guide the future synthesis of other metastable materials beyond the boron–carbon system.

Boron and carbon each exhibit uniquely complex elemental chemistry.1 In particular, boron has been crystallized in a plethora of allotropic forms owing to the rich chemistry that results from its lack of electrons compared to its number of ground state atomic orbitals.2 The vast number of carbon allotropes and their exceptional properties are well-known.<sup>3</sup> When combined, boron and carbon give rise to a phase space that is both enormous and often difficult to accurately characterize, owing to the nearly identical size and hybridization states of the two elements. The single thermodynamically-stable binary phase at ambient conditions is boron carbide (nominally B<sub>4</sub>C), a structure comprised of the dominant B<sub>12</sub> icosahedral unit prevalent in boron's elemental phases. However, as might be expected, this phase exhibits wide compositional variability and there remains disagreement as to its formal description as B<sub>12</sub>C<sub>3</sub> or B<sub>13</sub>C<sub>2</sub> in its most carbon-rich form.<sup>4,5</sup>

Numerous theoretical studies<sup>6-9</sup> support growing experimental evidence<sup>10-12</sup> for the existence of a high symmetry graphite-like phase of composition  $BC_3$  (Fig. 1). Indeed, the equilibrium solubility of boron in graphite is established to be

relatively high: 2.35 at% at 2375 °C.<sup>13</sup> The difference in formation energies between BC<sub>3</sub> and equivalent quantities of graphite and boron carbide is not currently known, but is likely small compared to the thermal energies present at the temperatures characteristic of graphite and carbide synthesis. Hence, many researchers have reported strategies to isolating metastable BC<sub>3</sub> at ambient conditions, <sup>12,14–20</sup> an effort that has been intensified owing to the promising theoretical applications of BC<sub>3</sub> for energy storage applications. <sup>21–26</sup>

Our previous work<sup>12</sup> culminated in the synthesis of a bulk (free-standing) graphite-like material of composition BC3 via the bottom-up reaction of two molecular precursors: boron tribromide (BBr<sub>3</sub>) and benzene (C<sub>6</sub>H<sub>6</sub>). At that time, previous efforts had only succeeded to produce either thin-films (led by Kouvetakis, Bartlett and coworkers)10,15 or bulk flakes via the laborious synthesis and subsequent pyrolysis of 1,3-bis-(dibromoboryl) benzene (led by King, Wright, and coworkers)11. While the earlier approach benefits from the direct pyrolysis of highpurity, commercially available precursors, only small quantities of BC3 could be produced. The latter approach produces bulk BC3-like material yet it suffers in terms of complexity and from the presence of residual contaminants due to the complex, multi-step organic synthesis of the boronated benzene; our approach combined these two strategies by leveraging the highpurity nature of combining only two commercially-available precursors (similar to the BCl<sub>3</sub> and benzene chosen for the thin-film synthesis) and the liquid miscibility associated with a Br-terminated boron precursor (as inspired by the 1,3-bis-(dibromoboryl)benzene precursor chosen to produce the first

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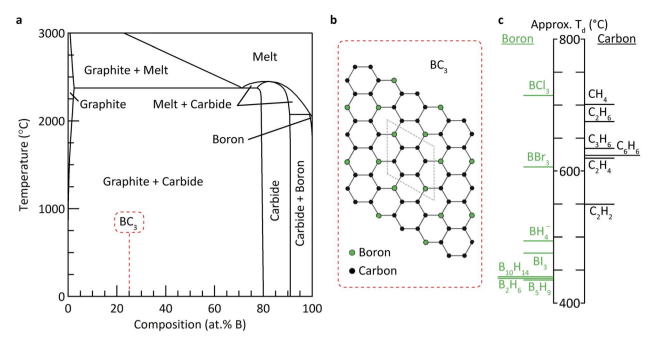


Fig. 1 (a) Binary phase diagram of the boron–carbon system.<sup>27</sup> The region of interest for the formation of metastable BC<sub>3</sub> is highlighted. (b) The proposed in-plane structure and unit cell of BC<sub>3</sub>.<sup>7,10</sup> (c) Thermochemical ranking of boron and carbon precursors based on their estimated temperature of decomposition ( $T_d$ ).

free-standing BC<sub>3</sub>-like flakes). The crystalline quality of the resulting products is low, especially with respect to ordering in the stacking direction, but our direct synthesis approach has been shown to yield long-range ordered in-plane structures with a vibrational structure that is consistent with BC<sub>3</sub>.<sup>20</sup>

The investigation of directly-synthesized  $BC_3$ -like materials for hydrogen storage and for electrochemical energy storage applications has remained severely limited by its inherent release of HBr at elevated temperatures, typically 800 °C, in closed-reactor conditions, preventing the use of stainless steel reaction vessels or their equivalent. Therefore, two important synthetic goals of research in this area became (1) to achieve higher crystallinity of bulk  $BC_3$  and (2) to devise halide-free routes to the synthesis of bulk  $BC_3$  in order to prevent the release of corrosive HX or  $X_2$  gases (X = halogen) at elevated temperatures. Our natural hypothesis was that diborane ( $B_2H_6$ ), the smallest borane species and a reactive gas at ambient pressure,§ would be well-suited to reactions with benzene to produce bulk  $BC_3$  with only the release of  $H_2$  as a side-product.

The resulting studies of the reaction between diborane and benzene within a custom-designed apparatus comprising a closed, stainless steel reaction vessel at up to 800 °C (see ESI† for details) consistently revealed a heterogeneous mixture of products, a hodgepodge compounded by the presence of metal particle-catalyzed graphitic fibers which are known to be characteristic of carbon soots produced under similar conditions. Control reactions (e.g., of liquid mixtures of condensed diborane and benzene) in the absence of catalytic effects from the metal-walled reactor were not safe enough to conduct within the scope of this work. Meanwhile, benzene remained the optimal hydrocarbon precursor owing mainly to

the desire to preserve its ring structure within the planar layers of BC<sub>3</sub> (Fig. 1b), and hence we set out to identify a compatible, halide-free boron precursor to pair it with in our efforts to achieve the goals listed above.

Two additional halide-free molecular boron precursors were then explored as candidates toward elucidation of the synthesis of bulk BC<sub>3</sub>: decaborane (B<sub>10</sub>H<sub>14</sub>) and a bespoke polyaromatic carborane species (1,8-naphthalenediyl-bridged diborane, NDB).30 The former, a readily available commercial compound, is obtainable as a high-purity crystalline solid at ambient conditions and is miscible in benzene up to the desired B:C ratio of 1:3. However, reactions between decaborane and benzene at up to 800 °C always generated carbon-rich flakes with a colorful (iridescent) thin surface layer of amorphous boron, and with no discernible content of BC<sub>3</sub> (see ESI†). The latter precursor, NDB, itself requires a multi-step organic synthesis that, even under strict air-free handling and after judicious purification, inevitably leads to the introduction of impurities such as oxygen, chlorine, and aluminum in the eventual high-temperature pyrolysis product (see ESI†). Scanning electron microscopy (SEM) studies of this product revealed the existence of a rippled, carbide-like film on the surface of carbon-rich flakes. The overall maximum boron content of such products is in any case limited to ~17 at% owing to its stoichiometry of  $B_2C_{10}H_{10}$  (B: C of 1:5), preventing the realization of BC<sub>3</sub>. No other planar carboranes exist with a higher content of boron, and hence a halide-free single precursor "tiling" route to bulk BC<sub>3</sub> was also abandoned. The products of the attempted syntheses of bulk BC3 from diborane(II), decaborane(III), and NDB(IV) are compared to that from boron tribromide(I), the previously established route, in Fig. 2. In summary, a systematic

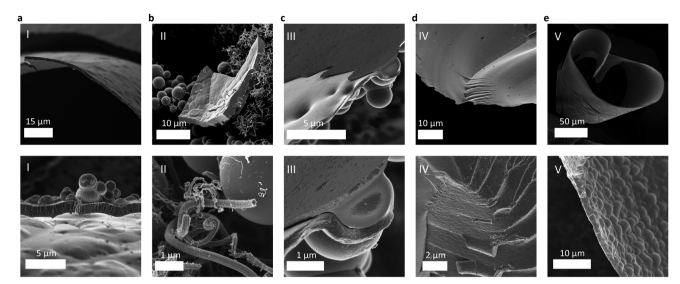


Fig. 2 SEM images of the graphitic  $BC_x$  pyrolysis products resulting from five different synthesis routes: (a) route I via boron tribromide, (b) route II via diborane, (c) route III via decaborane, (d) route IV via NDB, and (e) route V via sodium borohydride.

approach to identifying a compatible halide-free boron precursor for reactions with benzene at near 800 °C was needed.

The heterogeneity of each of the products of the aforementioned reactions can likely be explained by a mismatch in the decomposition or cracking temperatures of the boron and carbon species within each reaction. In order to realize a homogeneous metastable phase within a binary system, the thermodynamically favored phases (enriched in one or the other of the two elements, in this case graphite and boron carbide) must be prevented from forming, which requires the simultaneous decomposition of the corresponding precursors: i.e., co-pyrolysis. We undertook to estimate the relative decomposition temperatures  $(T_d)$  of an expansive library of boron and carbon precursors using a simple density functional theory (DFT) method, described in detail in the ESI.† In short, the change in energy of simplified decomposition reactions were calculated at the MN15/def2-QZVPP31,32 level (recently benchmarked for studies of small molecular systems of composed of boron, carbon, and hydrogen).33,34 The normalized change in electronic energy,  $\Delta E^{\rm SCF}$ , where the  $\Delta$  refers to decomposition into B or C and the relevant diatomic gases, was found to be highly correlated with the experimental decomposition temperature of each precursor (Fig. S24†). The results revealed that among all of the precursors investigated, borohydride (BH<sub>4</sub><sup>-</sup>) exhibited the closest energetic match to benzene, without the presence of borohalide (B-X) bonds (Fig. 1c).

Borohydride salts are widely available in high purity, and their decomposition temperatures are known to be directly related to the electronegativity of the counter ion  $(M^+)$ .<sup>35</sup> However, no previous studies of even common borohydrides (M = Li, Na, or Mg) with simple hydrocarbon precursors such as benzene have been reported. Many of the less-stable borohydrides are known to emit diborane at modest temperatures  $(e.g., \text{ as low as } \sim 61 \, ^{\circ}\text{C}$  for  $\text{Al}(\text{BH}_4)_3$ ) during decomposition.<sup>35,36</sup> On the

other hand, the stable borohydrides such as Li(BH<sub>4</sub>) do not produce detectable diborane upon decomposition, leading to polymeric B–B bond formation and the eventual formation of higher boranes such as  $B_{12}H_{12}^{2-}$ .<sup>37</sup> The similar decomposition temperatures of NaBH<sub>4</sub> ( $\sim 600 \, ^{\circ}\text{C}$ )<sup>35</sup> and benzene ( $\sim 750 \, ^{\circ}\text{C}$ )<sup>38</sup> led us to predict that reactions between these two precursors could result in the formation of BC<sub>3</sub>.

A stoichiometric mixture of benzene and sodium borohydride (with a B:C ratio of 1:3) was charged into a quartz ampule under argon, solidified by submersion in liquid nitrogen, and then flame-sealed within the ampule under active vacuum (referred to herein as "synthesis route V," see ESI†). Pyrolysis was carried out at 800 °C for 1 h, and the sample was then collected in air at room temperature. To prevent the formation of sodium metal and/or reactions of the sodium with the walls of the quartz reactor, a sodium-ion-trapping agent should also be added to the mixture (e.g., I2, Se, or Te) to form a water-soluble salt that can be easily separated during workup of the product. Dark lustrous flakes were collected, showing very little tendency to adhere to the inner walls of the ampule, a characteristic feature of bulk polycrystalline BC<sub>3</sub> derived from BBr<sub>3</sub> (referred to herein as "synthesis route I"). <sup>12</sup> SEM imaging further supports that sodium borohydride-derived BC<sub>3</sub> (route V) exhibits a homogeneous, flake-like morphology (~2 μm in width) with the same "cratered" texture (Fig. 2e) as that derived from boron tribromide (Fig. 2a).

Further materials characterization (Fig. 3) reveals borohydride-derived bulk BC<sub>3</sub> (route V) to be similar in structure and composition to that derived from boron tribromide (route I). The underlying structure is exclusively graphitic, as evidenced by powder X-ray diffraction (XRD), with significant misalignment (turbostratic disorder) along the stacking direction as shown by the broad reflection at  $2\theta=25.4^{\circ}$  corresponding to an interlayer spacing of 3.5 Å. This broadness is in agreement with theoretical predictions for crystalline BC<sub>3</sub> which

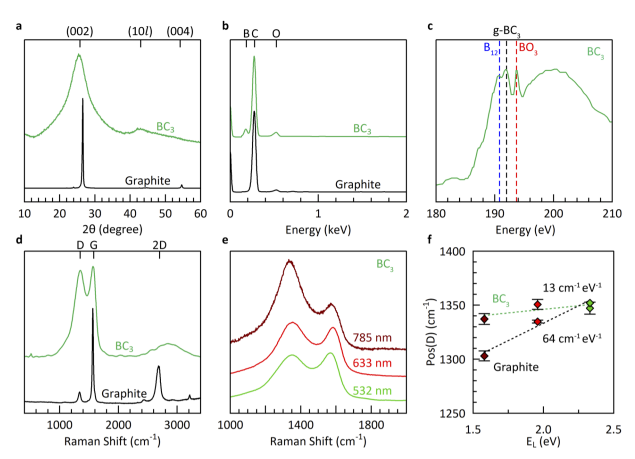


Fig. 3 Materials characterization of borohydride-derived  $BC_3$  compared to graphite: (a) XRD patterns, (b) EDX spectra, (c) XAS spectra at the BKedge, and (d) Raman spectra at 532 nm. (e and f) Dispersion relation of the D peak as revealed by multi-wavelength Raman spectroscopy.

show that its various high-symmetry stacking sequences all lie within a narrow range of formation energies (±0.011 eV per atom),<sup>7</sup> indicating the strong likelihood of stacking misalignment. The in-plane structure is revealed to be more ordered, with an average domain size of 42 Å according to a Scherrer analysis of the (10*l*) peak at  $2\theta = \sim 42^{\circ}$ . The elemental composition measured using energy dispersive X-ray spectroscopy (EDX) is estimated to be  $\sim$ BC<sub>5</sub>, identical to equivalent measurements of the products of synthesis route I. It must be noted that past work has shown EDX to underestimate the boron content compared to more sensitive methods such as elastic recoil detection analysis (ERDA).12

The boron chemical environments present in borohydridederived BC3 were investigated by a combination of X-ray absorption spectroscopy (XAS) and multi-wavelength Raman spectroscopy (Fig. 3c-f). The presence of planar g-BC<sub>3</sub> units in long-range ordered, hexagonal symmetry is evidenced by a dominant absorption feature in the B K-edge at 192.0 eV.39 Some icosahedral B<sub>12</sub> environments are also detected at 190.8 eV, likely due to the small degree of mismatch between the decomposition temperatures of benzene and BH<sub>4</sub> which favors the slightly earlier formation of boron rich phases prior to the desired reaction to form BC<sub>3</sub>. Most importantly, the Raman spectrum exhibits the characteristic features of bulk BC<sub>3</sub>: a broad and intense D peak at Pos(D) =  $\sim$ 1350 cm<sup>-1</sup>,

exhibiting almost no dispersion as a function of the energy of irradiation  $(E_L)$ , and a broad, intense, and modulated 2D region associated with two-phonon intervalley scattering. 40 Previous work revealed that the dispersion relation of the D peak, in particular, is highly sensitive to the presence of hexagonally ordered flower-like B<sub>6</sub>C<sub>6</sub> units, originating from the flattening of the highest optical phonon branches around the K point in BC<sub>3</sub>.20 The non-dispersive nature of the D peak of borohydridederived BC3 coupled with the presence of an intense feature in the XAS spectrum at 192 eV is the ultimate evidence for the structure of BC<sub>3</sub> (in the absence of long-range crystalline order).20

In conclusion, we report a novel borohalide-free route to turbostratic BC3 with exceptional in-plane ordering via the reaction of benzene with sodium borohydride at 800 °C. The computational method developed herein can be extended to identify candidate precursors for other bottom-up synthesis efforts toward binary metastable phases. The limited set of experimental conditions explored herein compels future work to tune the decomposition profile of the borohydride precursor by varying the counter ion and/or matching a given salt with a hydrocarbon of similar stability, opening an abundance of possible chemical routes to more crystalline forms of graphitelike BC<sub>3</sub>.

### Data availability

The associated experimental and computational data are available in the ESI† and further details are available by request from the corresponding author.

#### Author contributions

The manuscript was written through contributions of all authors. D. M. developed the hypothesis, performed the primary experiments, analyzed all of the data, and wrote the first draft of the manuscript. C. M. and M. A. M. performed computational experiments and analysis. C. M. and D. L. assisted with pyrolysis experiments. D. M. and Y. H. performed XAS measurements. N. P. S. and M. T. M. supervised the experimental work and M. A. M. supervised the computational work.

#### Conflicts of interest

There are no conflicts to declare.

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#### Notes and references

§ Important safety precautions associated with the handling of gaseous diborane at high pressures are detailed in the ESI.†

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