Novel precursors for boron nanotubes: the competition of two-center and three-center bonding

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We present a new class of boron sheets, composed of triangular and hexagonal motifs, that are more stable than structures considered to date and thus are more likely to be the precursors of boron nanotubes. We describe a simple and clear picture of electronic bonding in boron sheets and highlight the importance of three-center bonding and its competition with two-center bonding, which can also explain the stability of recently discovered boron fullerenes. Our findings call for reconsideration of the literature on boron sheets, nanotubes and clusters.

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All boron nanotubes (BNT), regardless of diameter or chirality, are predicted to be metallic and have large densities of states (DOS) at their Fermi energies ($E_F$) \cite{footnote1}. In contrast, carbon nanotubes (CNT) can be semiconductors or metals with small DOS at their $E_F$. Metallic CNT are used widely to study one-dimensional (1D) electronics \cite{footnote2, footnote3} and are superconducting at low temperatures \cite{footnote4, footnote5}. Due to the larger DOS, BNT should be better metallic systems for 1D electronics and may have higher superconducting temperatures than CNT.

Recent experiments have fabricated boron nanotubular structures both as small clusters \cite{footnote6} and long, 1D geometries \cite{footnote7}. Understanding the properties of BNT is crucial for realizing their applications. For CNT, it has been fruitful to study two-dimensional (2D) graphene: e.g., many properties of CNT are derived directly from graphene \cite{footnote8, footnote9}. For boron, however, no 2D planar structure exists in its crystals which are built from $B_{12}$ icosahedra \cite{footnote10}. Researchers have proposed several 2D boron sheets (BS). The hexagonal graphitic BS was found to be unstable \cite{footnote11, footnote18}. Based on extensive theoretical studies of boron clusters \cite{footnote11, footnote12, footnote13, footnote14, footnote15}, an Aufbau principle was proposed whereby the most stable structures should be composed of buckled triangular motifs \cite{footnote12}. Experiments on small clusters of 10-15 atoms support this view \cite{footnote16}. A recent study of many possible sheet structures found, again, the buckled triangular arrangement to be most favorable \cite{footnote17}. Hence, 2D triangular BS have been studied and used to construct BNT \cite{footnote18, footnote19, footnote20}.

In this Letter, we present a class of boron sheets that are more stable than the currently accepted ones. We describe their structures, energetics, electronic states, and provide a clear picture of the nature of their bonding that clarifies their stability. We also show that clusters with these structures are competitive with or more favorable than those considered to date. Our findings have important consequences for understanding and interpreting the properties of these systems. For example, the unusual stability of $B_{80}$ fullerenes \cite{footnote21} can be explained by our bonding picture. Hence, in our view, it is necessary to reconsider previous work in this general field.

We use Density Functional Theory \cite{footnote22, footnote23} within the ab initio supercell planewave pseudopotential total energy approach \cite{footnote24}. Calculations are done by PARATEC \cite{footnote25}. We use both the local density approximation (LDA) \cite{footnote26, footnote27} and the generalized gradient approximation (GGA) \cite{footnote28} for exchange and correlation. Most results below employ the LDA and key results are checked by the GGA. The LDA and GGA yield same qualitative results with minor quantitative variances. The planewave basis has a 32 Ryd cutoff energy. K-point samplings for each case converge total energies to better than 1 meV/atom. Norm-conserving pseudopotentials have cutoff radii $r_s^p=1.7$ and $r_s^p=2.1$ a.u.. The BS are extended in x-y directions while supercells have periodic copies along z where a separation of 7.4 Å is sufficient for convergence. For all structures, relaxations are preformed until the atomic Hellmann-Feynman forces are less than 1 meV/Å and all in-plane stresses are less than 5 MPa.

Table I shows our results for four different sheets: the flat and buckled triangular sheets \cite{footnote18, footnote19}, the hexagonal sheet, and one of our sheets ($\alpha$ in Figure 1). The hexagonal sheet is unstable with respect to in-plane shear, so we obtain the values in Table I by maintaining hexagonal symmetry while optimizing the bond length. The

![FIG. 1: (A, B) Two examples of our BS (top view). Red solid lines show the unit cells. (C) Four boron clusters: $B_{24}(a)$ and $B_{12}(a)$ are clusters with hexagonal holes; $B_{24}(b)$ and $B_{12}(b)$ are the double-ring clusters from refs. \cite{footnote14, footnote15}. Gray balls are boron atoms, and gray “bonds” are drawn between nearest neighbors.](image-url)
The buckled triangular sheet is more stable than the flat sheet by 0.12 eV/atom. The triangular sheet has ε_{\text{BNT}} made from triangular sheets \[18, 19\]. Buckled direction \[19\]. We also can reproduce previous results on BNT made from triangular sheets \[18, 19\].

### Table I: Binding energies $E_b$ (in eV/atom) and geometric parameters (in Å) of four BS: the flat and buckled triangular sheets, the hexagonal sheet, and one of our sheets ($\alpha$ in Figure 1).

|                     | Flat triangular |          | Buckled triangular |          |
|---------------------|-----------------|----------|--------------------|----------|
|                     | $E_b$           | $d^{\text{flat}}$ | $E_b$           | $d^{\text{buckled}}$ |
|                     | $d^{\text{hex}}$ | $\Delta z$ | $d^{\text{diag}}$ | $\Delta z$ |
| LDA                 | 6.58            | 1.68     | 6.74              | 1.59     | 1.80  | 0.81  |
| previous LDA \[19\] | 6.76$^a$        | 1.69     | 6.94$^b$          | 1.60     | 1.82  | 0.82  |
| previous LDA \[19\]| 6.53            | -        | 6.79              | -        | -     | -     |
| GGA                 | 5.79            | 1.70     | 6.00              | 1.60     | 1.86  | 0.88  |
| previous GGA \[17\] | 5.48$^b$        | 1.71     | 5.70$^b$          | 1.61     | 1.89  | -     |

|                     | Hexagonal |          | Sheet $\alpha$ |          |
|---------------------|-----------|----------|----------------|----------|
|                     | $E_b$     | $d^{\text{hex}}$ | $E_b$     | $d^{\text{diag}}$ |
|                     | $d^{\text{hex}}$ | $\Delta z$ | $d^{\text{diag}}$ | $\Delta z$ |
| LDA                 | 5.82      | 1.65     | 6.86            | 1.64-1.67|
| GGA                 | 5.25      | 1.67     | 6.11            | 1.66-1.69|
| previous GGA \[17\]| 4.96$^b$  | 1.68     | -               | -        |

$^a$Boron’s atomic spin-polarization energy of 0.26 eV/atom explains the $E_b$ differences between \[19\] and our work \[18\].

$b$While the absolute $E_b$ from \[17\] do not match our GGA results, $E_b$ differences among the sheets match very well.

The triangular sheet has $\eta=0$, the hexagonal $\eta=1/3$, and sheets $\alpha$ and $\beta$ have $\eta$ of 1/9 and 1/7, respectively.

$\textbf{A priori}$, the energies of these sheets can depend on both $\eta$ and the pattern of hexagons. This results in a huge phase space of hexagonal patterns for a given $\eta$. The most stable structures occur when the hexagons are distributed as evenly as possible at fixed $\eta$. Figure 2 shows the LDA binding energies $E_b$ versus $\eta$ for this class of structures. $E_b$ reaches a maximum of 0.86 eV/atom at $\eta=1/9$ (sheet $\alpha$). We also have investigated the other extreme where hexagons form lines (e.g., sheet $\beta$). These “linear” structures are more stable than the buckled triangular sheet for $\eta \approx 1/9$ but are less stable than the “evenly-distributed” class described above.

To explain the stability of these structures, we describe the nature of their electronic bonding. Generally, in-plane bonds formed from overlapping $sp^2$ hybrids are stronger than out-of-plane $\pi$-bonds derived from $p_z$ orbitals, so a structure that optimally fills in-plane bonding states should be most preferable. Guided by this principle, Figure 3 shows projected densities of states (PDOS) for five BS with separate in-plane (the sum of $s, p_x$ and $p_y$) and out-of-plane ($p_z$) projections.

We begin with the hexagonal sheet, a textbook $sp^2$ bonded system. All $sp^2$ hybrids are oriented along nearest neighbor vectors so that overlapping hybrids produce canonical two-center bonds. A large splitting ensues between in-plane bonding and anti-bonding states. The $p_z$ orbitals form their own manifold of bonding and anti-bonding states. The $p_z$ PDOS vanishes at the transition point between the two. In the case of graphene, the four valence electrons per atom completely fill the $sp^2$ and the $p_z$ bonding states, leading to a highly stable structure. However, a boron atom has only three valence electrons. As shown in Figure 3 some of the strong in-plane $sp^2$ bonding states are unoccupied, explaining the instability of this sheet. For our discussion below, this sheet is highly prone to accepting electrons to increase its stability they should be available from another source.

Next, we consider the flat triangular sheet. Each atom has six nearest neighbors but only three valence electrons. No two-center bonding scheme leads to a proper description. Previous work has noted qualitatively that a three-center bonding scheme exists here \[19\]. We now present a detailed model of the three-center bonding with crucial implications for the stability of our sheets. Figure 4 shows a choice of orientations for the $sp^2$ hybrids.
where three hybrids overlap within an equilateral triangle formed by three neighboring atoms. For an isolated such triangle, we have a simple $3 \times 3$ tight-binding problem with $D_3$ symmetry. Its eigenstates are dictated by group theory: one low-energy symmetric bonding orbital $b$ and two degenerate high-energy anti-bonding orbitals $a^*$. (This is “closed” three-center bonding; details on this and other types of bonds are found in standard references [28].) These orbitals then broaden into bands due to inter-triangle couplings. Separately, the $p_z$ orbitals also broaden into a single band (not shown). In Figure 3 the in-plane PDOS becomes zero at the energy separating in-plane bonding and anti-bonding states. Ideally this sheet would be most stable if: (i) two electrons per atom would completely fill the $b$-derived in-plane bonding bands, (ii) the anti-bonding $a^*$-derived bands were empty, and (iii) the remaining electron per atom would half fill the low-energy (bonding) portion of the $p_z$-derived band. This would mean that the $E_F$ would be at the zero point of the in-plane PDOS in Figure 3. Clearly, this picture is a valid zeroth-order description. However, $E_F$ lies slightly above the ideal position and makes some electrons occupy in-plane anti-bonding states. In other words, this sheet prefers to donate these high-energy electrons which has critical implications below. (Although we seem to break symmetry by making half of the triangles filled and half empty, filling the entire $b$-derived in-plane bonding band makes all hybrids equally occupied. This restores full in-plane symmetry: i.e., the two possible initial orientations of hybrids give the same final state.)

The flat triangular sheet, however, buckles under small perturbations along $z$ [18]. The buckling mixes in-plane and out-of-plane states and can be thought of as a symmetry reducing distortion that enhances binding. As shown in Figure 3 some states move below the $E_F$ as indicated by the small peak immediately below the $E_F$.

Finally, we turn to the new structures. The above discussion has shown that the hexagonal sheet should be able to lower its energy by accepting electrons while the flat triangular structure has a surplus of electrons in anti-bonding states. From a doping perspective, the three-center flat triangular regions should act as donors while the two-center hexagonal regions act as acceptors. Thus if the system is able to turn into a mixture of these two phases in the right proportion, it should be able to benefit from the added stability of both subsystems. Specifically, the hexagon-triangle mixture with the highest stability should be the one that places the $E_F$ precisely at the zero-point of in-plane PDOS, filling all available in-plane bonding states and none of the anti-bonding ones. The remaining electrons will fill the low-energy $p_z$-derived states, leading to a metallic system. These expectations are born out clearly in Figure 3 as well as in the energetic stability of the various structures (Figure 2). In fact, the most stable sheet, $\alpha$, satisfies this condition precisely while the less stable sheet, $\beta$, has a slight shift of $E_F$ from the ideal position.

These findings have ramifications for boron clusters. Our structures and bonding picture can explain that the extreme stability of $B_{80}$ fullerenes composed of triangular motifs with pentagonal holes [21] is due to the well

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**FIG. 3:** Projected Densities of States (PDOS) for four BS. DOS are projected in-plane (sum of $s$, $p_x$, and $p_y$) and out-of-plane orbitals ($p_z$). Red solid lines show in-plane and blue dashed lines show out-of-plane projections; thick vertical solid lines show $E_F$. All curves are broadened using Gaussians with a width of 0.3 eV. The vertical scale is arbitrary.

**FIG. 4:** Three-center bonding scheme in flat triangular sheets. Left: orientation of $sp^2$ hybrids. Center and right: overlapping hybrids within a triangle ($D_3$ symmetry) yield one bonding ($b$) and two anti-bonding ($a^*$) orbitals. These then broaden into bands due to inter-triangle interactions.
balance of three-center and two-center bonds. Also the α sheet can be seen as the precursor of B$_{80}$ just as graphene is the precursor of carbon fullerenes. We also have studied some clusters. Figure 1 shows the double-ring structures for B$_{24}$ and B$_{32}$ along with clusters constructed by us. The new B$_{24}$ cluster with a hexagon hole is less favorable by 0.08 eV/atom while the B$_{32}$ is more favorable by 0.03 eV/atom than the corresponding double-ring. The stability of our sheets, of B$_{80}$, and our clusters with hexagonal holes suggests that for boron systems with more than 20-30 atoms, the Aufbau principle breaks down and a more general structural rule is required.

In brief, we demonstrate a novel bonding mechanism in pure boron compounds arising from the competition between two- and three-center bonding. This explains the stability of our boron sheets as well as larger boron clusters. Our results have relevant implications on the stability and structure of boron clusters, boron nanotubes, and other boron systems.

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