Effect of Net Charge on the Relative Stability of 2D Boron Allotropes

Dan Liu* and David Tománek*

Physics and Astronomy Department, Michigan State University, East Lansing, Michigan 48824, United States

Supporting Information

ABSTRACT: We study the effect of electron doping on the bonding character and stability of two-dimensional (2D) structures of elemental boron, called borophene, which is known to form many stable allotropes. Our ab initio calculations for the neutral system reveal previously unknown stable 2D ε-B and o-B structures. We find that the chemical bonding characteristic in this and other boron structures is strongly affected by extra charge. Beyond a critical degree of electron doping, the most stable allotrope changes from ε-B to a buckled honeycomb structure. Additional electron doping, mimicking a transformation of boron to carbon, causes a gradual decrease in the degree of buckling of the honeycomb lattice that can be interpreted as piezoelectric response. Net electron doping can be achieved by placing borophene in direct contact with layered electrides such as Ca₂N. We find that electron doping can be doubled by changing from the B/Ca₂N bilayer to the Ca₂N/B/Ca₂N sandwich geometry.

KEYWORDS: Ab initio calculation, electronic structure, doping, boron, structural stability, 2D

Identifying the most stable allotrope of a given compound is one of the key problems in physics and chemistry. Whereas charge-neutral systems have attracted the most attention, notable exceptions are low-dimensional systems that can be charged by doping in specific bulk geometries. Electron doping induced by Li intercalation does not affect the honeycomb structure of 2D graphene layers in graphite intercalation compounds (GICs) but does change the equilibrium structure of MoS₂ monolayers from the 2H to the 1T allotrope. Even though the effect of excess charge on chemical bonding and equilibrium geometry should be general, we expect the most drastic changes to occur in structures of elemental boron, which is notorious for its many stable allotropes. What appears as frustrated bonding in mostly metallic 2D structures of elemental boron, called borophene, reflects the inability of the element to follow the octet rule of bonding due to its electron deficiency. We speculate that both the electron deficiency and thus the chemical bonding characteristic may be modified by placing a nonzero charge on B atoms. In that case, the most stable borophene structure may differ from a puckered triangular lattice with monatomic vacancies or equally stable irregular structures identified in the neutral system. Indeed, placing a net charge of ≲1 e per atom on borophene, provided by an Al(111) substrate or intercalating Mg ions in between borophene layers in the MgB₂ compound, changes the most stable allotrope to a very different honeycomb lattice. Structural changes induced by charging may significantly modify the electronic structure, turning semiconducting 2H-MoS₂ to metallic 1T’-MoS₂ locally and undoped borophene to honeycomb lattices in diborides including MgB₂, which displays superconducting behavior.

In this study, we explore the effect of net charge on the bonding character and structural stability of 2D allotropes of boron. Our ab initio calculations for the neutral system reveal a previously unknown stable 2D ε-B structure with a 0.2 eV wide fundamental band gap. We find that the chemical bonding characteristic in this and other boron structures is strongly affected by extra charge, including a 23% lattice constant change in ε-B, induced by changing the net charge from 0.25 holes to 0.25 electrons per B atom. Beyond a critical degree of doping near 0.5 electrons/atom, the most stable allotrope changes from ε-B to a buckled honeycomb structure. Additional electron doping, mimicking a transformation of boron to carbon, causes a gradual decrease in the degree of buckling of the honeycomb lattice that can be interpreted as piezoelectric response. We propose that net electron doping can easily be achieved by placing borophene in direct contact with layered electrides such as Ca₂N. In this system, we find that electron doping of borophene can be doubled by changing from the B/Ca₂N bilayer to the Ca₂N/B/Ca₂N sandwich geometry.

As mentioned above, the vast number of stable neutral 2D boron allotropes, including α-B, β-B, γ-B, δ-B, and η-B structures, reflect a frustrated bonding character of electron-deficient boron. This element may engage its three valence electrons in only three covalent bonds, resulting in an electron sextet instead of the desirable octet noble-gas configuration. To partly compensate for the lack of electrons in the octet configuration, boron atoms often prefer to increase their coordination to six nearest neighbors in the triangular lattice.
This is essentially equivalent to adding electrons\textsuperscript{19,20} to the individual atoms that are held together by pure three-center bonding. On the contrary, the stability of the triangular lattice is often enhanced by removing atoms and forming hexagon-shaped monatomic vacancies. This process somehow mimics subtracting electrons while the structure locally converts to a honeycomb lattice, where atoms are held together by two-center bonding. The competition between two-center and three-center bonding has been used to identify the optimum concentration of hexagonal vacancies in the neutral triangular lattice\textsuperscript{16,21,22} but likely also controls less common neutral structures with four- and five-fold coordinated boron atoms.\textsuperscript{10,11} Biasing this competition by net charge has been shown to affect the fraction of hexagonal vacancies in triangular borophene lattices.\textsuperscript{23} Even though doping of B layers has been linked to the unexpected superconducting behavior of MgB\textsubscript{2} and found to be useful to modulate CO\textsubscript{2} capture,\textsuperscript{24} no systematic attention has been paid to the possibility of deliberately changing the bonding and the equilibrium structure of boron by excess charge.

A viable possibility to significantly dope 2D structures of elemental boron by electrons is to place them in direct contact with electrides including Ca\textsubscript{3}N\textsuperscript{25,26} and Y\textsubscript{2}C.\textsuperscript{27} In these highly electronegative systems with a layered structure, regions of large electron density are found in between the layers. The charge density in this electron layer amounts to one electron per formula unit in Ca\textsubscript{3}N and two electrons per formula unit in Y\textsubscript{2}C. The possibility of exfoliation down to a monolayer\textsuperscript{25} has been demonstrated in Ca\textsubscript{3}N, so that assembly of various vertical heterostructures is possible.

Results. Allotropes of Neutral and Doped Borophene. Inspired by the honeycomb structure of the negatively doped boron sublattice found in diborides including MgB\textsubscript{2}, we started our investigation of electron-doped borophene structures with the honeycomb lattice. To provide substantial configuration freedom for the lattice structure, we consider a superlattice with 32 atoms per unit cell and subject all atoms in the supercell to random distortion. To study the effect of doping on the equilibrium structure and the chemical nature of bonding, we changed the degree of doping gradually and optimized each system using the conjugate gradient (CG) optimization method.

Starting with no excess charge, we found the neutral honeycomb structure to be unstable and to convert to rather stable allotropes that have not been reported previously. The first allotrope, shown in Figure 1a and called $\epsilon$-B, is only 7 meV/atom less stable than the most stable $\alpha'$-B phase\textsuperscript{1} and displays a very uncommon morphology with triangles and pentagons, quite distinct from the well-documented class of triangular lattices with vacancies. The structure shown in Figure 1b, which we call $\omega$-B, has a very similar morphology to known allotropes containing triangles and hexagons only\textsuperscript{1,3,5,6} but is 26 meV/atom less stable than $\alpha'$-B. $\epsilon$-B and $\omega$-B add to the large number of known allotropes, and we expect many more to follow.

Both $\epsilon$-B and $\omega$-B are buckled. Still, we could locate a locally stable, flat counterpart of $\omega$-B, which is 7 meV/atom less stable than $\omega$-B. The cohesive energies of neutral borophene allotropes are compared in Table 1.

Next, we added extra 5, 10, 13, 16, 24, and 32 electrons to the 32-atom unit cell with an initial honeycomb arrangements and optimized the geometry. We found the optimum geometries not to depend on initial deformations imposed on the starting structure, which may not be reachable in molecular dynamics trajectories. The optimum boron structures labeled $\epsilon_1$ and $\epsilon_0$ and their average doping levels, ($\Delta Q$), are displayed in Figure 2a–f. Optimum structures for different doping levels are discussed in the Supporting Information (SI). We find all of these structures to differ significantly from the neutral $\epsilon$-B and $\omega$-B structures in Figure 1, which have been optimized in the same way.

The structure $\epsilon_1$-B in Figure 2a, which contains triangles, pentagons, and hexagons, is reminiscent of $\epsilon$-B but is completely flat. With increasing electron doping, the density of pentagons gradually diminishes and eventually vanishes in $\epsilon_3$-B in Figure 2c, representing the buckled $\lambda_1$ phase.\textsuperscript{3} At the same time, the density of hexagons increases from $\epsilon_1$-B to $\epsilon_4$-B until all other polygons are eliminated in $\epsilon_7$-B shown in Figure 2d–f, as the doping level exceeds |$\Delta Q|$ $\geq$ 0.5 e/atom. The buckled honeycomb lattice of $\epsilon_7$-B gradually flattens to the graphene-like structure of $\epsilon_\infty$-B with increasing electron doping.

The interpretation of these structural changes is rather straightforward. With one extra electron per atom, boron behaves as sp\textsuperscript{2}-bonded carbon with four valence electrons, with atoms forming the 2D graphene honeycomb lattice. This structure is rather robust with respect to electron and hole doping, as evidenced in GIC structures. Similarly, also, doped boron should keep its optimum honeycomb lattice structure even if the net charge may be smaller or larger than one extra.

Table 1. Cohesive Energy, $E_{\text{coh}}$, of Selected Neutral Borophene Allotropes, Obtained Using DFT-PBE Calculations\textsuperscript{4,6}

<table>
<thead>
<tr>
<th>allotrope</th>
<th>$E_{\text{coh}}$ (eV/atom)</th>
<th>n</th>
<th>Z</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\epsilon$-B</td>
<td>5.699 \textsuperscript{b}</td>
<td>8</td>
<td>4</td>
</tr>
<tr>
<td>$\omega$-B</td>
<td>5.680 \textsuperscript{b}</td>
<td>32</td>
<td>4, 5, 6</td>
</tr>
<tr>
<td>$\alpha'$-B</td>
<td>5.706 \textsuperscript{b}</td>
<td>8</td>
<td>5, 6</td>
</tr>
<tr>
<td>$\beta_1$-B</td>
<td>5.762 \textsuperscript{c}</td>
<td>8</td>
<td>5, 6</td>
</tr>
<tr>
<td>$\beta_2$-B</td>
<td>5.712 \textsuperscript{c}</td>
<td>5</td>
<td>3, 5, 6</td>
</tr>
<tr>
<td>$\delta_1$-B</td>
<td>6.23 \textsuperscript{d}</td>
<td>5</td>
<td>3, 5, 6</td>
</tr>
<tr>
<td>$\delta_2$-B</td>
<td>5.662 \textsuperscript{c}</td>
<td>4</td>
<td>6</td>
</tr>
<tr>
<td>$\gamma_1$-B</td>
<td>5.723 \textsuperscript{c}</td>
<td>4</td>
<td>3, 5</td>
</tr>
<tr>
<td>$\gamma_2$-B</td>
<td>6.19 \textsuperscript{e}</td>
<td>4</td>
<td>3, 5</td>
</tr>
</tbody>
</table>

\(n\) is the number of boron atoms per unit cell and Z is the coordination number of individual atoms in the unit cell.\textsuperscript{3} Present work.\textsuperscript{3} Ref 3. \textsuperscript{3} Ref 8.
electron per atom. As seen in Figure 2d–f, the honeycomb structure of borophene, with different degree of buckling, is preferred for the excess charge \( \langle \Delta Q \rangle \) ranging between \(-0.5\) and \(-1.0\) e/atom.

To understand the change in the electronic structure that caused a profound structural change from neutral \( e^-\)-B in Figure 1a to negatively charged \( e^-\)B” in Figure 2f, we first probed the charge density changes, \( \Delta \rho \), associated with charging. Our results for \( \Delta \rho \) caused by placing artificially extra electrons on free-standing borophene structures \( e_1\)-B and \( e_2\)-B are shown in Figure 3a,b. \( \Delta \rho \) can be viewed as the crystal counterpart of the Fukui function, and our results indicate that the extra electrons are accommodated in \( p_z \) (or \( \pi \)) states normal to the layer, very similar to graphene layers.

A realistic way to provide a high degree of electron doping, we suggest, is to place borophene layers in contact with the \( \text{Ca}_2\text{N} \) electrode. The optimum lattice constant of the triangular lattice of \( \text{Ca}_2\text{N} \) is \( a = 3.97 \) Å. This layered system has the nominal configuration \( [\text{Ca}_2\text{N}]^+ \). \( \text{e}^- \), with layers of \( \text{Ca}_2\text{N} \) separated by layers of excess electrons, and can be exfoliated down to a monolayer chemically.\(^{25}\) It is to be expected that borophene will be electron-doped when placed in the region of excess electrons outside a \( \text{Ca}_2\text{N} \) monolayer. Because we focus on general trends rather than minute details, we studied the charge redistribution using only two prototype structures of doped borophene, namely, \( e_1\)-B at low- and \( e_2\)-B at high-level doping, in contact with \( \text{Ca}_2\text{N} \). To determine the degree of electron doping in borophene caused by a contact to \( \text{Ca}_2\text{N} \), we inspected the charge redistribution when placing \( e_1\)-B and \( e_2\)-B on top of a \( \{\text{Ca}_2\text{N}\}_2 \) bilayer, and our results indicate that the extra electrons, in contact with \( \text{Ca}_2\text{N} \). To determine the degree of doped borophene, namely, \( \langle \Delta Q \rangle \). Additional results for \( e_3\)-B and \( e_4\)-B on top of a \( \{\text{Ca}_2\text{N}\}_2 \) bilayer are presented in the SI.

We should point out that incommensurate vertical heterostructures formed of doped borophene and \( \text{Ca}_2\text{N} \) layers cannot be represented accurately in a periodic structure used in our computational approach. In our calculation, we used the optimum interlayer distance \( d = 4.0 \) Å in agreement with our results in Figure 4a,b. We furthermore matched \( 1 \times 3 \ e_1\)-B supercells with \( 3 \times 7 \) \( \text{Ca}_2\text{N} \) supercells in the \( e_1\)-B/\( \text{Ca}_2\text{N} \) superstructure and primitive unit cells of \( e_2\)-B with \( 3 \times 3 \) \( \text{Ca}_2\text{N} \) in the \( e_2\)-B/\( \text{Ca}_2\text{N} \) superstructure. The remaining lattice mismatch of \( \lesssim 2% \) was accommodated by averaging the lattice constants of borophene and \( \text{Ca}_2\text{N} \). Because of this minor lattice distortion, results presented in Figure 3c–f may differ to a small degree from the charge redistribution in an incommensurate structure. Comparing results for the bilayer in Figure 3c,d with those in the sandwich structure in Figure 3e,f, we see clearly that borophene receives twice the number of electrons in the sandwich in comparison with the bilayer structure. Specifically, \( \langle \Delta Q \rangle \) in \( e_1\)-B almost doubles from \(-0.16\) e/atom in Figure 3c to \(-0.31\) e/atom in Figure 3f. Similarly, \( \langle \Delta Q \rangle \) in \( e_2\)-B almost doubles from \(-0.21\) e/atom in Figure 3d to \(-0.41\) e/atom in Figure 3f.

We have evaluated the dependence of the total energy on the interlayer distance, \( d \), in the \( \text{Ca}_2\text{N}/\text{B}/\text{Ca}_2\text{N} \) sandwich structure for the \( e_1\)-B and \( e_2\)-B allotropes and present our results in Figure 4a,b. In both structures, the optimum interlayer distance \( d \approx 4.0 \) Å.

There is a significant electron accumulation with a maximum at \( d \approx 3 \) Å outside a free-standing \( \text{Ca}_2\text{N} \) electrode layer,\(^{25}\) which is accommodated by an adjacent borophene layer. In this case, the net charge on the borophene layer may be changed by changing the interlayer distance, \( d \). Our results for \( \langle \Delta Q \rangle \) as a function of \( d \) are presented in Figure 4c for \( e_1\)-B and in Figure 4d for \( e_2\)-B sandwiched in between two \( \text{Ca}_2\text{N} \) layers. We find it interesting that \( \langle \Delta Q \rangle \) decreases almost linearly with increasing interlayer distance.

At the equilibrium interlayer distance, we find \( \langle \Delta Q \rangle \approx -0.31\) e/atom in the system with \( e_1\)-B and \( \langle \Delta Q \rangle \approx -0.41\) e/atom in the system with \( e_2\)-B. The higher value of \( \langle \Delta Q \rangle \) in \( e_2\)-B is associated with the better ability of this structure to accept electrons.

It is well known that in GICs the net charge transferred from dopant atoms to the graphene layers changes the lattice constant. This change is relatively small, amounting to \( \approx 0.8% \) in \( \text{KCa} \) with \( \langle \Delta Q \rangle \approx -0.125\) e/C atom. Our corresponding results for the effect of doping on the lattice constants in the stable \( e^-\)B allotrope are shown in Figure 5a. The lattice changes are anisotropic and larger than found in graphene. We find that electron doping expands the lattice more along the softer \( \bar{a}_1 \) direction than along the harder \( \bar{a}_2 \) direction. At the electron-doping levels \( \langle \Delta Q \rangle \approx -0.3 \) to \(-0.5\) e/atom discussed above for the \( \text{Ca}_2\text{N}/\text{B}/\text{Ca}_2\text{N} \) heterostructures, the lattice expansion exceeds 10%.

The in-layer stiffness of neutral and electron- and hole-doped \( e^-\)-B is addressed in Figure 5b. We used the shorthand notation \( 0 \) for the neutral system, \(- \) for \( \langle \Delta Q \rangle = -0.125\) e/B, and \( + \) for \( \langle \Delta Q \rangle = +0.125\) e/B doping. Irrespective of doping, the energy change due to in-layer strain, \( \sigma \), is much larger along...
the harder \( \mathbf{a}_2 \) direction than along the softer \( \mathbf{a}_1 \) direction. In terms of the 2D elastic constants, we find for the softer \( \mathbf{a}_1 \) direction \( c_{11}(-) = 67.99 \text{ N/m}, c_{11}(0) = 18.70 \text{ N/m}, \) and \( c_{11}(+) = 29.98 \text{ N/m} \). Along the harder \( \mathbf{a}_2 \) direction, we find \( c_{22}(-) = 446.79 \text{ N/m}, c_{22}(0) = 360.52 \text{ N/m}, \) and \( c_{22}(+) = 338.58 \text{ N/m} \).

Electronic Structure of Borophene Allotropes. The electronic band structure of selected boron 2D allotropes discussed in this study is shown in Figure 6. We should note that density functional theory (DFT) calculations used in this study do not represent the true quasi-particle band structure and typically underestimate band gaps. With this fact in mind, we find that according to the Perdew–Burke–Ernzerhof (PBE) results in Figure 6a, the neutral \( \epsilon\)-B allotrope of Figure 1a is a semiconductor with a small indirect band gap of \( E_g = 0.2 \text{ eV} \). Whereas stretching along the soft \( \mathbf{a}_1 \) direction by 4% turns \( \epsilon\)-B into a direct-gap semiconductor, compressing by 4% along \( \mathbf{a}_1 \) causes gap closure. Stretching \( \epsilon\)-B by 1% along the hard \( \mathbf{a}_2 \) direction turns this allotrope metallic, whereas compression by 1% changes its indirect gap to a direct gap. More details of the electronic structure of strained \( \epsilon\)-B are provided in the SI.
We find all doped borophene allotropes to be metallic or semimetallic, as shown in Figure 6b–f. Inspection of the band structure reveals the formation of a Dirac cone at $K_1$ and $K_2$ in the honeycomb structures in $\epsilon_1$-B in Figure 6e and in $\epsilon_3$-B in Figure 6f. The Dirac cone appears ~1.4 eV above $E_F$ in $\epsilon_4$-B at the doping level $|\Delta Q| = 0.500$ e/B and at $E_F$ in $\epsilon_5$-B at $|\Delta Q| = −1.000$ e/B, which mimics the structure and valence charge of graphitic carbon.

**Discussion.** The majority of the reported stable structures of neutral 2D borophene were triangular lattices containing arrays of monatomic vacancies, reflecting the frustrated bonding character of electron-deficient boron. When considering the effect of excess charge on the bonding geometry, we observed a transition driven by increasing net negative charge from structures containing triangles and higher polygons, depicted in Figures 1a and 2a–c, for $|\Delta Q| < 0.5$ e/B to all-hexagon structures, depicted in Figure 2c–f, for $|\Delta Q| > 0.5$ e/B. The highest coordinations number of six that may be achieved in a network of triangles reflects to some degree the vain attempt of neutral boron atoms to satisfy the octet rule. Excess negative charge, with the maximum value $|\Delta Q| = 1.0$ e/B considered here, offers the ability to satisfy this rule in the honeycomb network of $\epsilon_5$-B in Figure 2f.

To understand the chemical origin of these structural changes, we need to inspect the charge redistribution caused by additional doping, corresponding to the Fukui function for a crystal. Corresponding results for $\Delta \rho$ in selected electron-doped borophene structures are presented in the SI. Complementing our results presented in Figure 3 for borophene interacting with Ca$_2$N, these results suggest that additional electrons are first accommodated in $\pi$ (or $\pi$) states normal to the borophene layer, which cause buckling, and then in $\sigma$ states, which reduce the amount of buckling. Starting with the lightly electron-doped $\epsilon_1$-B containing triangles, we observe an increasing degree of buckling with increasing electron doping up to $|\Delta Q| \leq 0.5$ e/B. At that point, the structure changes to the honeycomb structure mimicking hole-doped graphene. For $|\Delta Q| > 0.5$ e/B, additional excess charge gets increasingly accommodated in $\sigma$ states, which are much closer to $E_F$ in borophene than in graphene, causing a reduction of buckling down to zero for $|\Delta Q| = −1.0$ e/B.

Assuming that the Ca$_2$N electrode can transfer up to one electron per formula unit to a borophene layer, we can expect the maximum average charge $|\Delta Q|$ per boron atom in B/Ca$_2$N bilayers to range from $−0.22$ e in $\epsilon_1$-B/Ca$_2$N to $−0.28$ e/atom in $\epsilon_5$-B/Ca$_2$N. The transferred charge may be up to twice as large in the Ca$_2$N/B/Ca$_2$N sandwich structures. These values are slightly larger but close to those found in the actual heterostructures, reported in Figure 3. We compared the charge transfer between monolayers and multilayers of the Ca$_2$N electrode in contact with borophene and found essentially no difference, as seen in the SI. Thus the number of Ca$_2$N layers does not affect the maximum value of $|\Delta Q|$.

It is not easy to achieve the doping level $|\Delta Q| = −1.0$ e/B by contacting an electronegative material. According to our results for the Ca$_2$N/B/Ca$_2$N sandwich structure in Figure 4d, the Ca$_2$N electrode can provide only up to ~0.4 electrons per boron atom, much less than the desired doping level of 1 e/B.

**Figure 6.** Electronic band structure of (a) neutral $\epsilon_1$-B, (b) $\epsilon_4$-B with $|\Delta Q| = −0.156$ e/B, (c) $\epsilon_1$-B with $|\Delta Q| = −0.406$ e/B, (d) $\epsilon_4$-B with $|\Delta Q| = −0.500$ e/B, (e) $\epsilon_5$-B with $|\Delta Q| = −0.750$ e/B, and (f) $\epsilon_6$-B with $|\Delta Q| = −1.000$ e/B, calculated using the DFT-PBE functional. The indirect fundamental band gap is indicated by the red arrow in panel a.
An alternative to Ca$_3$N is Y$_2$C, which can supply twice as many electrons as Ca$_3$N but is hard to exfoliate. Assuming maximum charge transfer from Y$_2$C to borophene, B could receive up to 0.8 electrons in the Y$_2$C/B/Y$_2$C heterostructure. Such a large electron transfer should further augment the Coulomb attraction between borophene and Y$_2$C, thus further deducing the interlayer distance, as seen in Figure 3a,b, and increase the amount of electron transfer, possibly up to 1.0 e/B.

We have mentioned electronic structure parallels between the honeycomb structure of ϵ-B carrying one extra electron per atom and graphene. Even though the system of π electrons near $E_F$ and the Dirac cone in the corner of the Brillouin zone (BZ) occurs in both systems, there are notable differences between the systems. In graphene, the top of the σ band lies >3 eV below $E_F$, whereas this energy difference is only 0.2 eV in ϵ-B. Apparently, the lower core charge of elemental boron is the main reason why the σ and π bands are energetically closer than in graphene. As mentioned before, this results in an increased role of σ states in electron-doped borophene structures.

According to our results in Figure 5a, changing the doping level (ΔQ) from −0.25 e to +0.25 e results in a 23% increase in the borophene lattice constant $a_{\text{t}}$. Conversely, we may speculate that changing the lattice constant should modulate the electron transfer from the electrode to the borophene layer, thus changing the dipole moment normal to the interface. In that case, an in-plane vibration of the heterostructure will cause this dipole to oscillate and to emit an electromagnetic signal of the same frequency.

Clearly, the most drastic effect of electron doping is seen in the hexagonal borophene structure stabilized in MgB$_2$, which is the cause of superconductivity in this system. Electron doping may also be used as a way to change structures in a predictable way. As an illustration, we find the $\chi_3$-B structure, which has been synthesized at 680 K on Ag(111), to be very similar to the electron-doped, buckled $\epsilon_3$-B structure. In that case, a simpler way to fabricate $\chi_3$-B may consist of bringing borophene in contact with an electrode, allowing it to relax to $\epsilon_3$-B while electron-doped. The final flat structure of $\chi_3$-B may evolve after the electrode has been removed. These two examples illustrate new possibilities of using doping to change the structure of boron. Even though doping may have a lesser effect on the equilibrium structure of other systems, our findings about the interplay between structure and excess charge have a general validity.

In summary, we have studied the effect of electron doping on the bonding character and stability of 2D structures of elemental boron, called borophene, which is known to form many stable allotropes. Our ab initio calculations for the neutral system have revealed previously unknown stable 2D $\epsilon$-B and $\omega$-B structures in addition to previously reported triangular lattices with monatomic vacancies. We found that the chemical bonding characteristic in this and other boron structures is strongly affected by extra charge, which is first accommodated in the π and subsequently in the σ network of electrons. Beyond a critical degree of electron doping, the most stable allotrope was found to change from $\epsilon$-B containing triangles and higher polygons to a buckled honeycomb structure. Additional electron doping, mimicking a transformation of boron to carbon, causes a gradual decrease in the degree of buckling of the honeycomb lattice that can be interpreted as a piezoelectric response. We also found that net electron doping can be achieved by placing borophene in direct contact with layered electrodes such as Ca$_3$N. We found that electron doping can be doubled to ~0.4 e/B atom by changing from the B/Ca$_3$N bilayer to the Ca$_3$N/B/Ca$_3$N sandwich geometry.

Computational Techniques. Our calculations of the stability, equilibrium structure, and electronic structure have been performed using DFT as implemented in the SIESTA code. Periodic boundary conditions have been used throughout the study, with monolayers represented by a periodic array of slabs separated by a 30 Å thick vacuum region. We used the PBE exchange-correlation functional. The SIESTA calculations used norm-conserving Troullier-Martins pseudopotentials, a double-ζ basis including polarization orbitals, and a mesh cutoff energy of 250 Ry to determine the self-consistent charge density, which provided us with a precision in total energy of ≤2 meV/atom. The reciprocal space has been sampled by a fine grid of $4 \times 4$ k-points in the 2D BZs of the primitive unit cells of neutral and doped borophene containing 32 atoms, $3 \times 3$ k-points in the BZ of supercell of heterostructure of Ca$_3$N and $\epsilon_0$, and $3 \times 1$ k-points in the BZ of supercell of heterostructure of Ca$_3$N and $\epsilon_3$. Geometries have been optimized using the CG method until none of the residual Hellmann–Feynman forces exceeded $10^{-2}$ eV Å$^{-1}$.

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.nanolett.8b04968.

Additional results are presented for the characterization of borophene layers under conditions not covered in the main manuscript. These include optimum geometries and charge distribution in free-standing electron-doped borophene as well as in vertical heterostructures of borophene and Ca$_3$N bilayers. Also presented are electronic band structures of systems described in the main manuscript, results characterizing the effect of strain on the electronic band structure, and charge density differences corresponding to the Fukui function for selected doped borophene allotropes (PDF).

**AUTHOR INFORMATION**

**Corresponding Author**

E-mail: tomanek@pa.msu.edu.

**ORCID**

Dan Liu: 0000-0002-5211-4865

David Tománek: 0000-0003-1131-4788

**Notes**

The authors declare no competing financial interest.

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**REFERENCES**


