BN White Graphene with “Colorful” Edges: The Energies and Morphology

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ABSTRACT: Interfaces play a key role in low dimensional materials like graphene or its boron nitrogen analog, white graphene. The edge energy of hexagonal boron nitride (h-BN) has not been determined as its lower symmetry makes it difficult to separate the opposite B-rich and N-rich zigzag sides. We report unambiguous energy values for arbitrary edges of BN, including the dependence on the elemental chemical potentials of B and N species. A useful manifestation of the additional Gibb’s degree of freedom in the binary system, this dependence offers a way to control the morphology of pure BN or its carbon inclusions and to engineer their electronic and magnetic properties.

KEYWORDS: Boron-nitride, graphene, edges, interfaces, bandgap, magnetism

Current attention to the atomic monolayers of carbon—graphene, and especially the need to introduce an electronic gap in this gapless sheet, has brought to the spotlight its sibling two-dimensional (2D) material of hexagonal boron nitride, h-BN. Its bulk phase (often called white graphite) with weak interlayer bonding permits separation into individual sheets of white graphene. Furthermore, analogous to carbon nanotubes, tubular BN structures have been theoretically predicted, and consequently synthesized, and display a number of properties of fundamental and practical importance.

Morphologically alike to the honeycomb graphene and even with quite similar bond length (instead of 2.46 Å in C, lattice parameter in BN is 2.51 Å, henceforth used as a unit of length), the chemical alternation of B and N atoms causes the ionic nature of this distinctly different, insulating crystal. This makes BN interesting not only on its own, but especially as a counterpart to carbon graphene. Recent work has advanced the connection between the white and black graphene beyond the sheer analogy toward producing actual layers of hybridized BN and graphene domains. These experiments suggest the intriguing possibility of interfacing the white graphene (BN) with the “black” (C) within the same monolayer plane.

Energy analysis suggests that BNC mixtures should separate into immiscible BN and C. If indeed the distinct phases of BN and C coexist as the phase-separate epitaxial domains, their shape must be guided by the thermodynamic interface preferences. The corresponding quantity is the interface energy between the white and black graphene domains. This is of course closely related with a more basic property, the energy of pristine BN edges, γ(χ). Here we investigate how these interface energies can be calculated despite low symmetry of the lattice and reveal their variability with chemical potential of constituents. We further define the equilibrium shape of either free BN-clusters or of the inclusions of C in BN and vice versa and discuss rather briefly what effects they have on their electronic and magnetic properties.

The BN/C interfaces and the free BN edges can be of different kinds, depending on the edge-cut direction. In analogy to graphene, the basic edges are either along the zigzag (Z) atomic motif or along the armchair (A) pattern, in between, a variety of chiral edges can be cut at some angle χ. At this however the similarity ends, as the distinctly different physics of BN edges is caused by the lack of inversion symmetry in the lattice and its binary composition. We will see that the former complicates the definition of the edge energies, while the latter may offer an advantage: having two chemical constituents instead of one adds extra degree of freedom in the Gibbs phase rule, that is a freedom to balance the chemical potential between B and N, which permits control of the shapes, properties, and can be of practical interest.

Despite these fundamental differences, we can initially use the energy decomposition ansatz to represent the edge energy as a sum over the atoms of A and Z types. For BN though, an immediate generalization must be made to account for the fact that its zigzag edges are “chemically unbalanced” and expose either all B or all N atoms, henceforth ZB or ZN. Consequently, the nontrivial chiral angles range doubles to 60° instead of 30° for graphene or carbon tubes. Choosing the armchair BN-bond direction as reference for χ = 0, we obtain

\[ \gamma(\chi) = |\gamma| \cos(\chi + \pi) \]  

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where $|\gamma| = 2(\gamma_A^2 + \gamma_{ZA}^2 - \sqrt{3}\gamma_A\gamma_{ZA})^{1/2}$ and $C = \text{sgn}(\gamma_A) \cdot \text{arctan}(\sqrt{3} - 2\gamma_{ZA}/\gamma_A)$ with the subscript $x = N$ at $-30^\circ < \chi < 0$ or $x = B$ at $0 < \chi < 30^\circ$. This equation determines the energy of arbitrary edge, as long as the basic energies $\gamma_A$, $\gamma_{ZB}$, and $\gamma_{ZN}$ are known (that is for $\chi = 0$ and $\pm 30^\circ$). We therefore proceed finding these important quantities.

A common way to find a surface or edge energy is to compute the total slab energy,\textsuperscript{12,13} subtract the energy of equivalent material in its bulk form, and assign the rest to the edges.\textsuperscript{14} For BN ribbons with the basic edges of length $L$ (in units of $l$) one has (Figure 1a,b)

$$\gamma_A = \frac{E_a - M_{BN}\mu_{BN}}{2L}$$

and

$$\gamma_Z = \frac{E_Z + \gamma_{ZN}}{2} = \frac{(E_a - M_{BN}\mu_{BN})}{2L}$$

(2)

Here $E_a$ is the total energy of a ribbon (subscript “a”) oriented along either A or Z direction, $M_{BN}$ is a number of constituent BN-pairs, and $\mu_{BN} = \mu_B + \mu_N$ is their energy in the BN sheet. It can be chosen as zero level, while the chemical potentials for individual species can vary as $\mu_B, N = 1/2\mu_{BN} \pm \mu$. This approach, straightforward in case of graphene, gets however only “half job” done for BN. Although computations do yield $\gamma_A = 1.91$ eV and $\gamma_Z = 3$ eV, the latter is just a nominal average of the B- and N-rich edges (Figure 1b), while the true physical values remain elusive: only found is the sum of the opposite sides, but they seem to remain inesparable. This general issue with materials of lower symmetries, especially nonvariant upon inversion like BN, has been underscored by Cahn.\textsuperscript{15} Having such problem encountered earlier for 3D semiconductors, we have overcome it by considering the polyhedra with identical faces.\textsuperscript{13} For a simpler 2D case of BN, one can consider the triangles surrounded by either all-ZB or all-ZN (Figure 1c,d). Subtracting the material “cost” from the total energy of such triangle and omitting the insignificant contribution from the corners (which does not scale with size $L$), one recovers their true edge energies. The triangle inversion alters the excess of B to the excess of N around the perimeter $L$.

$$\frac{E_a - M_{BN}\mu_{BN} - L\mu_B}{3L} = \gamma_{ZB}(\mu) = \gamma_{ZB}^{\text{BN}} - \frac{\mu}{3}$$

(3a)

$$\frac{E_a - M_{BN}\mu_{BN} - L\mu_N}{3L} = \gamma_{ZN}(\mu) = \gamma_{ZN}^{\text{BN}} + \frac{\mu}{3}$$

(3b)

For a triangle of size $L$, direct counting yields $M_{BN} = 1/2L(L + 3)$, one extra N omitted as a corner correction, and $L$ extra B atoms. The latter is important as it means that the edge energy depends linearly on the chemical potential, as the right-hand side specifies. For the inverted triangle, B and N are interchanged; see eq 3b. Direct energy calculations (see Supporting Information) for a series of B-rich and N-rich triangular clusters of increasing size yield the data in Figure 1e. Clearly linear plots show that the size is sufficient, and their slopes determine the edge energies. By choosing the elemental chemical potentials as equal, $\mu_B = \mu_N = 1/2\mu_{BN}$, that is $\mu = 0$, we determine the values $\gamma_{ZB}^{\text{BN}} = 3.26$ eV and $\gamma_{ZN}^{\text{BN}} = 2.72$ eV. The average $[\gamma_{ZB}(\mu) + \gamma_{ZN}(\mu)]/2$ = 2.99 eV is independent of $\mu$ and agrees with $\gamma_Z$ of eq 2 to 0.1%, showing excellent consistency. What one gained is now well-defined energies for the basic edges of BN, at arbitrary chemical conditions

$$\gamma_A = 1.9, \gamma_{ZB} = 3.3 - \frac{\mu}{3}, \gamma_{ZN} = 2.7 + \frac{\mu}{3}$$

(4)

all in eV. Together with eq 1, eq 4 describes the BN edge of arbitrary direction, including the basic A, ZB, ZN, and the chiral types, at any chosen chemical potential $\mu$. This in turn enables the use of Wulff construction\textsuperscript{12} to easily determine the equilibrium shapes of pristine BN clusters at different conditions. We note that unlike in case of monoelemental graphene,\textsuperscript{10} the shapes can vary broadly with the chemical potential $\mu$. If BN edges reconstruct,
similar to graphene, their energies can also be obtained, in principle, through either ribbon or triangle models as described above; examples are shown in Supporting Information.

More important than the isolated clusters are possible epitaxial BN inclusions into C-graphene, or vice versa, the inclusions of black-graphene C into white-graphene BN matrix, in their hybrid monolayers.7,8 For these systems of emerging interest, we go on black-graphene C into white-graphene BN matrix, in their hybrid BN inclusions into C-graphene, or vice versa, the inclusions of above; examples are shown in Supporting Information.

control of domain patterns. This poses an important question of what equilibrium shapes to expect, as determined by Wulff construction, based in turn on the interface energy $\Gamma(\gamma)$. Figure 3 shows the basic interface energies along with the computed Wulff constructions for a range of chemical conditions quantified by the value of $\mu$. Modulation of the latter should allow one to broadly vary the shapes of inclusions, in striking contrast to pure graphene where the equilibrium islands are nearly hexagonal or more rounded polygons. Notably, nonintuitive shapes such as sharp triangles become preferred at N-rich (left) or B-rich (right) conditions, Figure 3. In the middle, the Wulff construct is hexagon with truncated-rounded polygons at the transient $\mu$ values.

Along with the shape, other properties such as electronic gap and especially magnetism also change. The magnetism originates from $\pi$-electrons of carbon and is localized at the Z-edge. Computations show (see Supporting Information) that triangle graphene domains are ferromagnetic with the total spin equal to the half of excess number of B or N atoms around the perimeter, obeying the Lieb’s theorem.17 The thin arrows in Figure 3b–f show the cumulative spin around the borders of different domains: triangles have the largest magnetic moment, reduced as the shapes get truncated, and then vanishing for nonmagnetic armchair-edge hexagon. The calculated change of magnetic moment per perimeter unit $|\mu|$ is summarized in Figure 3a by a dotted line. While the details of magnetism and electronics of the emerging islands deserve a separate study, beyond the scope of this report, we mention a few basic features. As example, we computed

$\Gamma_{BN/C} = \chi_{BN} + \chi_{C} - \chi_{BN/C}$

For clarity the material-subscripts are included here, while elsewhere the blank $\chi$ or $\gamma$ refer to the BN/C interface or BN pristine edge. It is opportune that the binding energy, as just a difference between the joint and detached counter-sides (similar to the work of adhesion or cleavage energy) does not depend on $\mu$, and its calculation is unobstructed by the low symmetries. Then, since $\chi_{BN}$ depends on chemical potential, so does the $\Gamma_{BN/C}$. We also include as example the hydrogen-passivated edges BN-H, Figure 2b. The data in Figure 2c show that the termination of the dangling bonds significantly reduces the energy in both cases, compared to pristine edge, as expected. Still, the chemical balance between B and N $(\mu$ value) controls the interface energy in the same way as for pristine edge.

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the spin density for the ferromagnetic triangles, either B-rich (Figure 4a) or N-rich (Figure 4b) at the borders with the densities of opposite spin in opposed shades of gray. If these mutually inverted triangles were to coexist in the same lattice, their magnetic moments would be antiparallel.

Because of quantum confinement, the intrinsically semimetallic graphene isles gain the characteristics of a quantum dots (QD, similar to vacancy clusters in graphane18); Figure 4 c,d shows the computed flat, dispersionless bands with significant HOMO–LUMO distances. The band gap \( E_g \) scales with the size \( M \) of the isle-QD as \( E_g \sim 1/\sqrt{M} \), following the trend for confined Dirac fermions.\(^{18,19}\) Further, in contrast to nonlinear hexagonal QD, a magnetic triangular isle displays distinctly different energy gaps for the spin majority and minority bands. Calculations show that the B-terminated QD have larger band gaps for spin majority than minority, while in the N-terminated QD the band gaps are in the reverse order, smaller for majority-spin bands and greater for minority.

A systematic way discussed above allows one to determine the energies \( \gamma(\chi) \) of arbitrary h-BN edges, for the basic directions given in eq 4. The dependence on chemical potential difference \( \mu \) between B and N species suggests that the equilibrium nanoparticles should be triangular (a counterintuitive shape of larger perimeter) with zigzag edges, except the narrow middle range of chemical conditions where more compact hexagons with armchair edge dominate. In fact, experimental observations\(^{20–22}\) and particularly the shape evolution between hexagonal and triangular nanoplates\(^{23}\) strongly support the notion of morphology control discussed above. Making further connection to BN nanotubes, their chiral distribution is not detailed, but believed in recent report\(^5,24\) to be mostly zigzag; this contrasts to carbon tubes, and can be explained by the preference of zigzag edge (Figure 2c) in either boron-rich or nitrogen-rich conditions. For graphene isles in BN, the BN/C interface energies are also unambiguously computed, as \( \Gamma_A = 0.56 \) and \( \Gamma_{BN} = (0.95 - \mu/3) \), \( \Gamma_{2N} = (0.38 + \mu/3) \), Figure 3a. Their variability suggests a way to selectively synthesize the inclusions of desirable shapes, with triangular quantum-dots islands displaying largest magnetic moment. Further work can clear the ways to use the balance between the B, N and C, and possibly the electronic chemical potential as additional controlling parameter,\(^5,25\) for design and growth of their hybridized domains with desirable physical properties. A concluding disclaimer can hardly be more precise than a quote from Conyers Herring’s classics:\(^{12,13}\) “Although the interpretation of experiments in such fields as the shapes of small particles and the thermal etching of surfaces usually involves problems of kinetics rather than mere equilibrium considerations, it is suggested that a knowledge of the relative free energies of different shapes or [edge] configurations may provide a useful perspective.”

**ASSOCIATED CONTENT**

Supporting Information. Additional information and figures. This material is available free of charge via the Internet at http://pubs.acs.org.

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