Semiconducting Layered Blue Phosphorus: A Computational Study

Zhen Zhu and David Tománek

Physics and Astronomy Department, Michigan State University, East Lansing, Michigan 48824, USA

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We investigate a previously unknown phase of phosphorus that shares its layered structure and high stability with the black phosphorus allotrope. We find the in-plane hexagonal structure and bulk layer stacking of this structure, which we call “blue phosphorus,” to be related to graphite. Unlike graphite and black phosphorus, blue phosphorus displays a wide fundamental band gap. Still, it should exfoliate easily to form quasi-two-dimensional structures suitable for electronic applications. We study a likely transformation pathway from black to blue phosphorus and discuss possible ways to synthesize the new structure.

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Elemental phosphorus is stable in a large number of structures, including the common white, red, violet, and black allotropes [1,2], with the color defined by the fundamental band gap. Most stable among them is black phosphorus, which—besides graphitic carbon—is the only layered structure of an elemental solid we know of. Individual layers of black phosphorus, shown in Fig. 1(a), resemble the honeycomb structure of graphene in terms of connectivity but are nonplanar. We noted that specific dislocations may convert black phosphorus, characterized by armchair ridges in the side view of the layers, to a well-defined structure with zigzag puckering, shown in Fig. 1(b). Assuming that the modified structure is stable, it appears worthwhile to study the equilibrium atomic arrangement in the bulk and the possibility of exfoliating individual layers. Whereas the observed fundamental band gap of 0.3–0.4 eV in black phosphorus [3,4] is rather narrow, it is intriguing to see if the modified phosphorus structure is a semiconductor with a wider gap. If the carrier mobility were also high, few-layer phosphorus in the new phase would become a worthy contender in the emerging field of post-graphene two-dimensional (2D) electronics.

Here, we use ab initio calculations to investigate this previously unknown phase of phosphorus that shares its layered structure with the most stable black phosphorus allotrope. We find this structural phase, which we call “blue phosphorus,” to be nearly as stable as black phosphorus. Whereas the in-plane hexagonal structure and bulk layer stacking of blue phosphorus are closely related to graphite, the main advantage of blue phosphorus is its wide fundamental band gap in excess of 2 eV. Because of the weak interlayer interaction, blue phosphorus should exfoliate easily to form quasi-2D structures for potential electronic applications. We study a likely transformation pathway from black to blue phosphorus and discuss possible ways to synthesize the postulated structure.

Our computational approach to gain insight into the equilibrium structure, stability, and electronic properties of blue phosphorus is based on ab initio density functional theory (DFT) as implemented in the SIESTA [5] and VASP [6] codes. We used periodic boundary conditions throughout the study, with multilayer structures represented by a periodic array of slabs separated by a 15 Å thick vacuum region. We used the Perdew-Burke-Ernzerhof [7] exchange-correlation functional, norm-conserving Troullier-Martins pseudopotentials [8], and a double-ζ basis including polarization orbitals. The reciprocal space was sampled by a fine grid [9] of 8 × 8 × 1 k points in the Brillouin zone of the primitive unit cell. We used a mesh cutoff energy of 180 Ry to

FIG. 1 (color online). The layered structure of (a) black and (b) blue phosphorus in top and side views. Atoms at the top and bottom of the nonplanar layers are distinguished by color and shading, and the Wigner-Seitz cells are shown by the shaded region. (c) Schematic of the conversion of black to blue phosphorus by dislocations, highlighted by the shaded regions and arrows. (d) Equilibrium structure of AB stacked blue phosphorus in side view.
determine the self-consistent charge density, which provided us with a precision in total energy of \( \lesssim \)2 meV/atom. All geometries have been optimized by SIESTA using the conjugate gradient method [10], until none of the residual Hellmann-Feynman forces exceeded \( 10^{-2} \) eV/Å. Our SIESTA results for the optimized geometry, interlayer interactions, and electronic structure were found to be in general agreement with VASP calculations. To verify the stability of the system at elevated temperatures, we performed a canonical molecular dynamics calculation of the monolayer and free-standing flakes using the SIESTA code. We used the Verlet integration algorithm to cover time periods up to 1 ps with the time step of 1 fs and present our results in the Supplemental Material [11].

The optimized reference structure of black phosphorus, shown in Fig. 1(a), is presented next to the proposed blue phosphorus structure in Fig. 1(b). The top view of both structures illustrates their similarity with the honeycomb lattice of graphite, which contains two atoms per layer per unit cell. Both phosphorus allotropes differ from graphite in the nonplanar structure of their layers. In top view, the isotropic structure of blue phosphorus in Fig. 1(b) differs significantly from the anisotropic structure of black phosphorus in Fig. 1(a). As seen in side view in Figs. 1(a) and 1(b), the puckered zigzag structure in the cross section of blue phosphorus differs from the distinct armchair ridges that cause the anisotropy of black phosphorus. The puckering in the blue phosphorus monolayer is similar to the postulated structure of single-wall phosphorus nanotubes [12].

The structural relationship between blue and black phosphorus is illustrated schematically in Fig. 1(c). A dislocation is introduced in a monolayer of black phosphorus by flipping specific P atoms from a “down” to an “up” position without changing the local bond angles, as described below and indicated by the arrows in Fig. 1(c). Subjecting every fourth row of P atoms to this transformation converts a monolayer of black to blue phosphorus. The location of dislocation lines in the monolayer structures is emphasized by the shaded regions in Fig. 1(c).

Atoms in the layers of blue phosphorus are covalently bonded at the equilibrium distance of 2.27 Å, resulting in a large binding energy of 5.19 eV/atom. This value differs from that of black phosphorus by less than 2 meV/atom, suggesting that blue and black phosphorus are equally stable. A weak interlayer interaction (see the Supplemental Material [11]) of 6 meV/atom holds the layered structure together at the interlayer distance \( d = 5.63 \) Å, as seen in Fig. 1(d), which indicates the possibility of easy exfoliation. The \( AB \) hexagonal stacking and the \( ABC \) rhombohedral stacking of layers differ energetically by less than 1 meV/atom.

The optimized hexagonal unit cell of an isolated blue phosphorus monolayer, shown in Fig. 1(b), is spanned by lattice vectors \( \vec{a}_1 \) and \( \vec{a}_2 \), with \( a = |\vec{a}_1| = |\vec{a}_2| = 3.33 \) Å. The influence of the interlayer interaction on the in-layer structure is small, causing only a negligible change from \( a = 3.324 \) Å in the bulk to \( a = 3.326 \) Å in the isolated monolayer. With its higher symmetry, the smaller hexagonal Wigner-Seitz cell of blue phosphorus, which contains two atoms and is shown in Fig. 1(b), differs from the rectangular Wigner-Seitz cell of the anisotropic black phosphorus monolayer with 4 atoms/unit cell, as shown in Fig. 1(a). Still, monolayers of blue and black phosphorus may form an “ideal” in-layer connection by the dislocation illustrated in Fig. 1(c).

One way to compare the stability and structural rigidity of the different phosphorus allotropes is by studying the vibration spectrum. Our results for the vibration spectra of blue and black phosphorus monolayers are presented in Fig. 2. We find the vibration spectra to be rather similar, reflecting a very similar bonding character. Acoustic and optical modes are well separated in both black and blue phosphorus. The harder longitudinal optical modes reflect a higher in-plane rigidity of the blue phosphorus allotrope in comparison to the accordionlike black phosphorus structure. The calculated flexural rigidity value \( D = 0.84 \) eV in blue phosphorus is lower than the \( D = 1.51 \) eV value in black phosphorus. These values provide a quantitative explanation for the dispersion of the flexural acoustic modes \( ZA \) near the \( \Gamma \) point [13] in the two allotropes (see the Supplemental Material [11]). The high rigidity of a free-standing blue phosphorus monolayer was also confirmed by our molecular dynamics calculations at nonzero temperatures, discussed in the Supplemental Material [11].

We next compare the slopes of the longitudinal acoustic branches near \( \Gamma \), which correspond to the speed of sound and reveal the in-plane stiffness. As seen in Fig. 2(a), the speed of sound along the \( \Gamma-Y \) direction in black phosphorus, \( v_s^{\Gamma-Y} \), is significantly higher than the \( v_s^{\Gamma-X} \), is significantly higher than the \( v_s^{\Gamma-X} = 3.8 \) km/s value along the \( \Gamma-X \) direction, reflecting an anisotropy in the elastic constants. The lower rigidity along the \( \Gamma-X \) direction, corresponding to the \( \vec{a}_1 \) direction in Fig. 1(a), reflects the fact that compression along \( \vec{a}_1 \) requires primarily bond bending, which comes at a lower energy cost than bond stretching. In strong contrast to those findings, our results in Fig. 2(b) indicate that the in-plane elastic response
of blue phosphorus is nearly isotropic, with nearly the same value $v_s = 7.7$ km/s for the speed of sound along the $\Gamma$-$M$ and the $\Gamma$-$K$ directions. The predicted finite in-layer compressibility of these nonplanar structures is advantageous when accommodating lattice mismatch during chemical vapor deposition (CVD) growth on a substrate.

Our DFT results for the electronic structure of blue phosphorus are presented in Fig. 3. The calculated band structure and density of states, presented in Figs. 3(a) and 3(b), indicate that a blue phosphorus monolayer is a semiconductor with an indirect band gap $E_{\text{gap}} \approx 2$ eV. Comparison with more proper self-energy calculations based on the $GW$ approach, performed by VASP, indicate that the DFT band gap is underestimated by $\approx 1.0 \pm 0.2$ eV in monolayers of blue P as a common shortcoming of DFT. Still, the electronic structure of the valence and the conduction band region in DFT is believed to closely represent experimental results. Therefore, we expect the charge density associated with states near the top of the valence band, shown in Fig. 3(c), to be represented accurately. These states correspond to the energy range highlighted by the green shading in Figs. 3(a) and 3(b), which extends from midgap to 0.1 eV below the top of the valence band. These states cause the interlayer band dispersion in few-layer systems (Supplemental Material [11]), and their hybrids with electronic states of the contact electrodes will play a crucial role in the carrier injection and quantum transport.

More interesting than the precise value of the fundamental band gap is its dependence on the number of layers $N$ in a multilayer slab, shown in Fig. 3(d). This is a consequence of the interlayer dispersion near the Fermi level in the bulk material, which is discussed in the Supplemental Material [11]. Independent of the type of stacking, we find $E_{\text{gap}}$ to be inversely proportional to $N$ between one monolayer and the bulk structure. We conclude that modifying the slab thickness may change the value of $E_{\text{gap}}$ by up to a factor of 2, which is very important for electronic applications. The $N$-dependent band gap value of $\lesssim 3$ eV lies just above the photon energy of visible blue light. We derive the name blue phosphorus from this absorption edge, which plays a key role in the optical appearance.

As seen in Fig. 3(e), the band gap $E_{\text{gap}}$ depends also sensitively on the applied in-plane strain $\sigma$. In view of the nonplanarity of the structure, the discussed strain range $-10\% < \sigma < +10\%$ can be achieved without a large energy penalty, as discussed in connection with the vibration spectra. The possibility to change the band gap value by $\lesssim 50\%$ in strained epitaxial geometries on different substrates is one more indication that blue phosphorus may find intriguing applications in nanoelectronics.

We next studied the possibility of converting a monolayer of black to blue phosphorus by introducing an array of dislocations, depicted schematically in Fig. 1(c). Our discussion of the conversion process including energy estimates is presented in Fig. 4 and the Supplemental

![FIG. 3 (color online). (a) Electronic band structure and (b) density of states of a blue phosphorus monolayer. (c) Electron density $\rho_{vb}$ in the 0.1 eV wide energy range near the top of the valence band in blue phosphorus, indicated by the green shaded region in (a) and (b). $\rho_{vb}$ is represented at the isosurface value $\rho = 7 \times 10^{-3}$ e/Å$^3$ and superposed with a ball-and-stick model of the structure. Dependence of the fundamental band gap $E_g$ on (d) the number of layers and (e) the in-plane stretching along the $\vec{a}_1$ direction.](176802-3)
Figure 4 (color online). (a) Series of structural snapshots depicting the transition from black to blue phosphorus. (b) Total energy change during the transformation from black to blue phosphorus. The reaction coordinate range 1–2 describes initial in-layer stretching along the horizontal direction, identified as \( \tilde{a}_1 \) in Fig. 1(a), followed by out-of-plane displacement of atoms at a fixed value of \( \tilde{a}_1 \) in the range 2–8.

Material [11]. The starting black phosphorus, the final blue phosphorus, and an intermediate structure are depicted in side view in Fig. 4(a). To estimate changes in the atomic arrangement during the conversion process, we changed and constrained the out-of-plane displacement of specific atoms in the unit cell to follow the “black-to-blue” trajectory and relaxed all other degrees of freedom. This provided us with a sequence of structures “1–8” in-between black and blue phosphorus, which loosely define the reaction coordinate in Fig. 4(b). The change in the unit cell from black to blue phosphorus has been imposed between steps 1 and 2 by deforming it from the initial size and shape in black phosphorus to the final rectangular supercell in blue phosphorus, followed by atomic relaxation. Our results for the relative energy with respect to the black phosphorus structure 1 again illustrate our finding that the blue phosphorus structure 8 is equally stable as 1. The activation barrier for the conversion process is likely overestimated due to the constraints imposed on the intermediate structures and may further be lowered by stretching black phosphorus layers. Thus, the true value should be below the already low value of 0.47 eV/atom, indicating the relative ease of mechanical conversion from black to blue phosphorus.

As suggested earlier, the weak interlayer interaction should allow the mechanical exfoliation of blue phosphorus in analogy to the black allotrope [14]. As a matter of fact, mechanically depositing a monolayer of black phosphorus onto a stepped substrate may cause the formation of dislocation lines and, thus, the formation of narrow domains of blue phosphorus in the monolayer structure along the step edges of the substrate. In analogy to graphene [15,16] and silicene [17], layered phosphorus structures may also be grown by CVD on specific substrates. As suggested in this study, blue and black phosphorus should be equally stable. The preferential phase should, thus, be determined by the lattice constant and the symmetry of the substrate in order to maximize the adsorption energy. Consequently, we expect blue phosphorus to form preferentially on substrates with hexagonal symmetry and a matching lattice constant, such as MoS\(_2\), or the (0001) surfaces of Zr and Sc, whereas black phosphorus should start growing on substrates with a rectangular lattice. Because of the low energy of forming a dislocation that connects blue and black phosphorus, both allotropes could coexist on particular substrates, including stepped surfaces, to optimize the adlayer-substrate interaction.

One of the main reasons for the interest in 2D semiconductors including graphene for electronic applications is the observed high mobility of carriers. Related quasi-2D systems, including MoS\(_2\), bring the benefit of a nonzero band gap but display lower intrinsic carrier mobility due to enhanced electron-phonon coupling, primarily caused by the presence of heavy elements such as Mo [18]. It appears likely that the blue phosphorus structure, similar to black phosphorus [14], may exhibit a higher carrier mobility than MoS\(_2\). The combination of a significant band gap and high carrier mobility would turn blue phosphorus into an excellent contender for a new generation of nano electronic devices.

In conclusion, we have conducted ab initio calculations to investigate a previously unknown layered phase of phosphorus, which we call blue phosphorus. We find blue phosphorus to be nearly as stable as black phosphorus, the most stable phosphorus allotrope. While sharing the atomic connectivity with the honeycomb lattice of graphene, layers of blue phosphorus are nonplanar. Whereas the bulk layer stacking of blue phosphorus is closely related to graphite, the main advantage of this allotrope is its wide fundamental band gap in excess of 2 eV. Because of the weak interlayer interaction, blue phosphorus should exfoliate easily to form quasi-2D structures for potential electronic applications. We have investigated a likely transformation pathway from black to blue phosphorus and show that the postulated structure may form spontaneously by CVD on a lattice-matched substrate or may result by stretching black phosphorus. Monolayers of blue phosphorus may form a structurally ideal connection to monolayers of black phosphorus with a different electronic structure.

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Note added in the proof.—We have learned that blue phosphorus is structurally related to the postulated [19] and observed [20] A7 phase of phosphorus.

\[^{*}\]tomanek@pa.msu.edu