Phosphorene nanoribbons

A. Carvalho, A. S. Rodin and A. H. Castro Neto

1 Graphene Research Centre and Department of Physics, National University of Singapore - 117542, Singapore
2 Boston University - 590 Commonwealth Ave., Boston MA 02215, USA

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Abstract – Edge-induced gap states in finite phosphorene layers are examined using analytical models and density functional theory. The nature of such gap states depends on the direction of the cut. Armchair nanoribbons are insulating, whereas nanoribbons cut in the perpendicular direction (with zigzag and cliff-type edges) are metallic, unless they undergo a reconstruction or distortion with cell doubling, which opens a gap. All stable nanoribbons with unsaturated edges have gap states that can be removed by hydrogen passivation. Armchair nanoribbon edge states decay exponentially with the distance to the edge and can be described by a nearly free electron model. This is used to prove the existence of the gap states in the limit of a semi-infinite plane.

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Black phosphorus is a van der Waals-bonded layered material, and the most stable form of this element. Its peculiar orthorhombic structure resembling corrugated cardboard sets it apart from other two-dimensional materials such as graphene and MoS$_2$. Unlike those, it is highly anisotropic with respect to the directions in-plane [1–4], and therefore it offers new terrain to the exploration of physical phenomena in two-dimensional materials.

The isolation of the monolayer form, named phosphorene, has been recently achieved by mechanical cleavage followed by plasma thinning [5] and multi-layered material is now routinely produced by exfoliation [6,7]. Both have been theoretically predicted to be direct-gap or nearly-direct-gap semiconductors with energy gaps ranging from 2 eV to 0.8 eV [8]. However, the band structure is far from typical. The effective masses for carriers are at least one order of magnitude higher along the direction of the to the zigzag ridges (y) than along the perpendicular direction (x). And the electron and hole masses are nearly the same along x, but very asymmetric along y. Further, due to the weak interaction between second and third nearest neighbors, it can easily be deformed under uniaxial compression, becoming, in sequence, an indirect-gap semiconductor, a semimetal, a metal, and ultimately undergoing a transition to a square lattice [1,2,9].

Phosphorene can, in principle, be cut and tailored into derived nanostructures [10–12]. An interesting feature of finite systems is the possibility of edge states. These states are localized at the system-vacuum interface and decay exponentially away from it. The nature of the edge states depends not only on the crystal structure of the system, but also on the way it terminates. Understanding the physics of nanoribbon edges is fundamental for predicting the behavior of real finite systems and for designing more complex nanostructures, such as nanotubes and voids. In this paper, we analyze the electronic states induced by finite edges. We show that armchair edges, cut along the x-direction, invariably lead to localized edge states, decaying exponentially towards the bulk. In contrast, edges cut along the y-direction were found to introduce less localized states for nanoribbons. All the nanoribbons are semiconducting in the large width limit.

We first discuss the origin of these localized gap states. In our previous work [1], we have shown that tight-binding has a limited utility for black phosphorus. Therefore, we will perform our analysis here using the nearly free electron model (NFEM). Before proceeding with the particular phosphorene case, we provide a derivation of the edge states for the reader’s convenience following a standard procedure.

We consider gap states arising from the existence of edges in finite monolayers. Since these states are exponentially decaying, they must have an imaginary momentum component perpendicular to the edge. Assuming that the system is located in the negative half-plane, the wave function inside it is given by

$$\Psi^I = e^{i qu + i(-ip)w} \sum_n A_n e^{i nLw},$$

(1)
where \( u \) is the coordinate along the edge and \( w \) is perpendicular. \( L_w = 2\pi/a_w \) is the smallest reciprocal vector transverse to the edge so that the summation runs over all the reciprocal vectors along \( \hat{w} \), in accordance with the Bloch theorem. To keep the notation as clean as possible, we introduce

\[
E_n^* = \frac{\hbar^2 \left[ q^2 + (nL_w - ip)^2 \right]}{2m} = (E_n^2)^*.
\]

Consider a gap at the \( \Gamma \) point between \( 2N \)-th and \( (2N + 1) \)-th bands. This means that the states in the vicinity of the gap are primarily defined by \( N \) and \( -N \)-th harmonics. Writing down the Bloch equation for only these two harmonics gives

\[
H_N \xi_N = (E - V_0) \xi_N, \quad H_N = \left( \begin{array}{cc} E_N^2 - V_{2N} \ 
2N \\
V_{2N} \ E_N^* \end{array} \right).
\]

In the equation above, \( V_n = V_{-n} = V_G = nL_w \hat{w} \), where \( V_G \) is the Fourier component of the lattice potential. Diagonalizing the Hamiltonian in eq. (3) yields

\[
\frac{E}{F} = N^2 L_w^2 + p^2 + q^2 + \frac{V_0}{F} \pm \sqrt{\frac{V_{2N}^2}{F^2} - 4N^2 L_w^2 p^2},
\]

where \( F = \hbar^2/(2m) \). The eigenvectors are

\[
\xi_N = \left( \begin{array}{c} e^{i\beta_N} \\
e^{-i\gamma_N} \end{array} \right), \quad \delta_N = \frac{1}{2} \arcsin \left( -\frac{2pNL_w F}{V_{2N}} \right)
\]

with

\[
2pNL_w \tan \delta_N = -\frac{V_{2N}}{F} \pm \sqrt{\frac{V_{2N}^2}{F^2} - 4N^2 L_w^2 p^2}.
\]

From this, we get the following form for the internal wave function:

\[
\psi_N \sim e^{i(qw + pNw) \cos (NL_w w + \delta_N)}.
\]

By setting the outside wave function to

\[
\psi^O = e^{i(qw - \sqrt{q^2 - E/F} w)}
\]

one can show that the momentum \( p_N \) is determined from

\[
p_N = NL_w \tan(NL_w W + \delta_N) = -\sqrt{q^2 - \frac{E_N}{F}},
\]

where \( W \) is the coordinate of the boundary. Defining \( \alpha = NL_w W, \ k = p_N/(NL_w), \ U = V_{2N}/(FN^2 L_w^2), \) and \( B = V_0/(FN^2 L_w^2) < 0 \) allows us to combine eqs. (4), (6), and (9) to get

\[
k = \left[ \frac{\tan \alpha \sqrt{1 - \frac{4B}{k^2 U} + 1}}{2k^2 U \tan \alpha} \right] + \left[ -B - U - k^2 - 1 + k \frac{4k/U}{\sqrt{1 - 4\frac{k^2}{U^2} + 1}} \right] = 0.
\]

Realistically \(|B| \gg |U|\). In addition, \( \tan \alpha \) is \( \pi \)-periodic so that \( \alpha \) enters as

\[
\alpha \to (\alpha \mod \pi) - \pi \Theta[(\alpha \mod \pi) - \pi/2].
\]

It is instructive to determine the values of \( \alpha \) that allow positive real-\( k \) solutions of eq. (10) for specified \( B \) and \( U \). This means that we need to find all values of \( \alpha \) for which eq. (10) is satisfied with \( 0 \leq k \leq U/2 \). To do so, we first find the values of \( \alpha \) which have solutions at \( k = 0 \) and \( k = |U|/2 \). For the first case, one gets

\[
\alpha_1 = \arctan \left[ \sqrt{-1 - B - U} \right] > 0.
\]

The second case yields

\[
\alpha_2 = \arctan \left[ \frac{-2 + U + \sqrt{-4 - 4B - U^2} \text{sign}[U]}{(-2 + U) \text{sign}[U] + \sqrt{-4 - 4B - U^2}} \right].
\]

Assuming that \(|B| \) is large enough so that all the square roots are real, it is possible to show by plotting that the \( \alpha \) that results in positive \( k \) is

\[
\frac{\pi}{2} \leq \alpha \leq \alpha_2 \quad \text{if} \quad U < 0, \quad \frac{\pi}{2} \leq \alpha \leq \alpha_1 \quad \text{or} \quad \alpha_1 \leq \frac{\pi}{2} \quad \text{if} \quad U > 0.
\]

This expression will be used to determine whether one should expect edge states in phosphorene.

In order to establish the existence of edge states, we need to find the value of \( U \) and, therefore, \( V_{2N} \). As there are four atoms per unit cell in phosphorene, the Fourier component \( V_G \) of the full potential is given by the product of the transform of a single-atom potential \( V^F \) multiplied by the geometrical structure factor \( Q_G \). Since the atomic potential is attractive, \( V^F_G < 0 \).

The atoms inside the unit cells are located at \( \pm l_1 \) and \( \pm l_2 \), where

\[
l_1 = \left( \frac{a \cos \frac{\theta}{2}}{2} \frac{a \sin \frac{\theta}{2}}{2} \right), \quad \text{and} \quad l_2 = \left( \frac{a \cos \frac{\theta}{2}}{2} + S a, \frac{a \sin \frac{\theta}{2}}{2} \right).
\]

see fig. 1 for details. The corresponding reciprocal vectors are

\[
Q = \left( \frac{\pi}{a} \frac{n_x}{S + \cos \frac{\theta}{2}} \frac{n_y}{\sin \frac{\theta}{2}} \right),
\]

where \( n_x \) and \( n_y \) are integers. If \( Q \) points along one of the principal axes, so that either \( n_x \) or \( n_y \) is zero, one gets the following. For \( n_x = 0 \),

\[
Q_G = 4 \cos \left( \frac{n_y \pi}{2} \right) = \begin{cases} 4, & \text{if} \ n_y = 4, 8, 12 \ldots, \\ 0, & \text{if} \ n_y \text{ is odd}, \\ -4, & \text{if} \ n_y = 2, 6, 10 \ldots. \end{cases}
\]
Thus, in this case $\hat{w} = y$ and $L_w W = \pi/2$. In addition, since phosphorene has 20 electrons per unit cell, the gap is between the tenth and the eleventh bands, and therefore we know that $N = 5$. This means that the geometrical structure factor $Q = -4$, in accordance with eq. (19). Therefore, $U \propto V_G = V_G^A \times Q > 0$ since the Fourier component of the attractive atomic component $V_G^A$ is negative. Finally, $\alpha = NL_w W = N \pi/2 \rightarrow \pi/2$ for odd $N$, per eq. (11). From eq. (15), we conclude that odd-$N$ harmonics have real-$k$ solutions and armchair edges contain edge states.

Next, we address the zigzag and cliff-edges. Here, the cuts are made along the $y$-axis and $L_w = \pi / (Sa + a \cos \frac{\theta}{2})$. Following our earlier assumption that the system extends to negative infinity, the boundaries of zigzag and cliff-edges are at $W = \pm \frac{\pi}{2} \cos \frac{\theta}{2}$, respectively. This yields $\alpha = \pm N \pi \cos(\theta/2)/[2(S + \cos(\theta/2))]$. Unlike the armchair case, here the reduced $\alpha$ is between 0 and $\pi/2$. Therefore, one needs to know the explicit form of the potential to determine $\alpha_1$ and $\alpha_2$ to see whether eqs. (14), (15) hold. Without this information, one cannot make a definite prediction concerning the existence of edge states.

We confirm this by modeling numerically the electronic band structure of infinitely long black-phosphorus nanoribbons [13]. The nanoribbons were modeled within the framework of density-functional theory [14], as implemented in the SIESTA package [15,16]. The generalized gradient approximation of Perdew, Burke and Ernzerhof is used for the exchange-correlation functional [17]. The electronic core is accounted for by using $ab$ initio norm-conserving pseudopotentials with the Troullier-Martins parametrisation [18] in the Kleinman-Bylander form [19]. The charge density was assumed to be independent of spin.

The basis sets for the Kohn-Sham states are linear combinations of numerical atomic orbitals [20,21]. These are a double-zeta polarized basis set for P and a single-zeta polarized basis set for H. The charge density is projected on a real-space grid with an equivalent cutoff energy of 250 Ry to calculate the exchange-correlation and Hartree potentials. A Monkhorst-Pack [22] scheme with at least 10 points along the periodic direction of the nanoribbons is used to sample the Brillouin Zone. The structural parameters of monolayer black phosphorus are given in table 1.

The armchair nanoribbons are constructed by cutting a phosphorene monolayer parallel to the $x$-axis, in such a way that they are symmetric with respect to the $xOz$ plane. As they can also be seen as being composed of $n$ atomic lines, each one parallel to the $x$-axis, we use $n$ to label the nanoribbon width (fig. 2).

The zigzag nanoribbons are constructed by cutting a phosphorene monolayer parallel to the $y$-axis, in such a way that they are symmetric with respect to the $yOz$ plane. We also label those nanoribbons by the number ($n$) of atomic lines parallel to the $y$-axis, where $n$ is an even number not multiple of 4 (fig. 2). Zigzag nanoribbons undergo little structural relaxation with exception of the outermost edge atoms. However, they are sensitive to Peierls distortion [12] with unit cell doubling, which results in a modest energy gain ($< 40$ meV).

Another type of edge along $y$, which we name “cliff”-edge, is found by cutting the layer along the line [EF] in fig. 1. As the cliff-edge leaves the edge P atoms with coordination one, it leads to considerable reconstruction. We found the reconstruction with unit cell doubling proposed in ref. [12] to be the most stable, leading to an energy gain of 0.37 eV/unit cell per edge relative to the reconstruction respecting the original periodicity of the nanoribbon along the $y$-direction. Since the resulting local edge structure is very distinct from the pristine black phosphorus, this

Table 1: Structural parameters of monolayer phosphorene: $a$ is the length of the in-plane bond, $Sa$ is the length of the projection of the out-of-plane bond in the plane, and $h$ is the layer thickness.

|          |      |      |
|----------|------|------|
| $a$ (Å)  | 2.2578 |     |
| $Sa$ (Å) | 0.8014 |     |
| $h$ (Å)  | 2.15  |     |
| $\theta$ (°) | 95.7 |     |
type of nanoribbon cannot be described using our previous analytical arguments. For cliff-edge nanoribbons \( n \) is a multiple of 4.

The presence of the edge boundary gives rise, in all three kinds of nanoribbons, to gap states (fig. 3) [23]. The gap state of armchair nanoribbons is an edge state with predominantly \( p_z \) character (fig. 4), of which the amplitude decays very fast with the distance from the edge (fig. 5).

In contrast, in unreconstructed zigzag and cliff-edge nanoribbons the Fermi level intersects the gap state, making them metallic. However, this is changed by the structure reconstruction/distortion with unit cell doubling. In the case of the zigzag phosphorene nanoribbon, the gap states are originally composed of \( p_x \) and \( p_z \) orbitals. These nanoribbons have a pair of gap states, which are half-occupied. The atoms in the origin of the gap states can be seen as a linear system, and since they have half-occupancy, are sensitive to Peierls distortion with unit cell doubling. The doubled states can be folded back into the smaller Brillouin zone and as a result of the distortion, a small gap is opened at the zone edge. The opening of the gap between occupied and unoccupied states results in a small energy lowering (< 40 meV).

The unoccupied gap state, similar to the original gap state of the zigzag nanoribbon before Peierls distortion,
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Fig. 5: (Color online) Profile of the wave function square of gap states of armchair, zigzag (Peierls-distorted) and cliff (reconstructed) nanoribbons with approximately the same width, integrated over the perpendicular directions. The states shown are the lowest unoccupied state at Γ (armchair, \( n = 13 \) and zigzag, \( n = 22 \)) and for the highest occupied state at Γ (cliff, \( n = 20 \)).

has higher amplitude at the edge, however it approaches that of a bulk state closer to the inner region (fig. 5). It can thus be considered a surface resonance, or a softly confined state \[24\], rather than an edge state. The occupied gap state, however, is an edge state.

The cliff nanoribbon is insulating with a double degenerate edge-related state in the lower gap. There is hybridization of the edge states with the conduction band bottom close to Γ. Thus, for some of the gap states the situation is intermediate between that of a pure edge state and that of a surface resonance. The edge states separated in energy from “crystal-like” states however are localized on the edge atoms (as the one shown in fig. 5).

A difference between phosphorene nanoribbons and their graphene counterpart is that while zigzag graphene nanoribbons give rise to flat bands near the Fermi level and as a result, the non-magnetic phase is unstable, in phosphorene zigzag nanoribbons, the edge-related states have mixed \( p_x \) and \( p_z \) character. They only become localized on the outer fringes for wide nanoribbons (\( n > 20 \)). The respective bands have dispersion in the \( k_y \)-direction (of the order of 1 eV to a few eV, depending on size). Thus, the density of states is smaller. Accordingly, we found no instability towards an antiferromagnetic phase.

To confirm that the gap states originate on the edge, we examined the effect of saturating the edge atoms with hydrogen. Each edge P was passivated with enough H to remove dangling bonds and restore three-fold coordination: 2 hydrogen atoms per edge P for armchair and cliff nanoribbons, and one hydrogen atom per edge P. In all three kinds of nanoribbons, the passivation with hydrogen removes the gap states fig. 6. As expected, the bandgap in this case decreases with increasing system size.

Finally, in order to evaluate whether such edges can be found experimentally, we look into their formation energy, defined as

\[
E_f = E_{\text{tot}} - \sum_i n_i \mu_i, \tag{21}
\]

where \( E_{\text{tot}} \) is the total energy of the unit cell, and \( n_i \) and \( \mu_i \) are the number of atoms of the element \( i = \{\text{P}, \text{H}\} \) and its chemical potential, respectively. The chemical potentials for P and H are obtained assuming an environment rich in \( \text{P}_4 \) and \( \text{PH}_3 \).

As black-phosphorus crystals, phosphorene nanoribbons have a negative formation enthalpy with respect to white phosphorus. In the unsaturated state, all three types of nanoribbons have close formation energies, in a range of about 0.1 eV/atom (fig. 7). With hydrogen saturation, the formation energies follow a similar trend, with the monosaturated zigzag-edge nanoribbons hovering higher in energy by about 0.2 eV. This energy difference might be the energy gain of forming a P-H bond from a P dangling orbital.

Fig. 6: (Color online) Band structures of H-saturated phosphorene nanoribbons. The zero of the energy scale is set to the valence band top.

Fig. 7: (Color online) Formation energy of the unsaturated (top) and saturated (bottom) nanoribbons, in an environment rich in \( \text{P}_4 \) and \( \text{PH}_3 \), as a function of the number of phosphorus atoms. For cliff-type nanoribbons the energies with a minimal unit cell or with unit-cell-doubling reconstruction (rec.) are both shown for comparison.
Figure 7 shows that the formation energy of the pristine nanoribbons increases, in absolute value, with the size, but the opposite is found for H-passivated nanoribbons. This indicates that black-phosphorus monolayers will dissolve in reducing conditions.

In conclusion, phosphorene nanoribbons were found to be semiconducting, in their lowest-energy structure, with edge-induced states in the gap. Armchair edges introduce edge states which decay exponentially with the distance to the edge.

In contrast, zigzag nanoribbons are metallic unless they undergo a Peierls distortion, at low temperature, which opens a gap between occupied and unoccupied bands. The highest occupied and lowest unoccupied bands originate on the edges and can be described as a surface resonance and an edge state, respectively. Since the distortion energy is very small (< 40 meV), it is likely that at high temperatures the edges present domain wall defects or even become metallic.

Cliff-type edges also originate gap states, but those are less localized on the edge atoms than the edge states of armchair nanoribbons.

An analytical description based on a NFEM provides further insight into the origin of edge states, in particular in the case of armchair edges. Since this is simultaneously the type of edge less changed by reconstruction, it is valid to generalise the DFT results obtained for narrow edges using this model. In fact, within the NFEM it is possible to prove the existence of edge states for non-interacting edges in armchair nanoribbons of any length. This applies to a semi-infinite sheet, which cannot be obtained from the DFT model. The amplitude of these edge states was shown to decay exponentially toward the bulk, in agreement with result of the first-principles calculations.

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