Twisted bilayer blue phosphorene: A direct band gap semiconductor

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Abstract

We report that two rotated layers of blue phosphorene behave as a direct band gap semiconductor. The optical spectrum shows absorption peaks in the visible region of the spectrum and in addition the energy of these peaks can be tuned with the rotational angle. These findings makes twisted bilayer blue phosphorene a strong candidate as a solar cell or photodetection device. Our results are based on ab initio calculations of several rotated blue phosphorene layers.

Keywords: blue phosphorene, bilayer, rotated, ab initio

1. Introduction

Starting with the synthesis of graphene[1] a new era of two dimensional (2D) materials have emerged. Among them we find hexagonal boron nitride, molybdenum disulphide, dichalcogenides, layered oxides[2], silicene and more recently one layer of black phosphorous (phosphorene)[3].

These materials, in general, have diverse properties, but some of them are semiconductors, a dream of stacking them in a certain order has become a reality resulting in structures with desired properties[4]. It has been possible not only to stack them in certain order but also rotate layers as well to exploit certain properties that twisted structures posses[5]. For instance in twisted bilayer graphene a plethora of interesting angle dependent properties have been found [6, 7, 8, 9, 10, 11, 12, 13].

Single layer phosphorene is a direct band gap semiconductor (2 eV). The addition of more layers reduce the value of the gap, allowing in principle to control its size which goes inverse with the layer number down to the value of the bulk (0.3 eV)[3, 14]. Nonetheless the main optical absorption peak of monolayer phosphorene is in 1.2 eV due to the presence of excitons[15].

On the other hand it has been shown in Ref.[16] that a displacement of some atoms in crystal phosphorene results in a honeycomb structure like graphene. The authors studied its stability by first principle methods and found that it can
be as stable as phosphorene; this new 2D allotrope, named blue phosphorene, has an unit cell composed of two sublattices, like in honeycomb lattice but with the difference that the two atoms are not in the same horizontal plane, forming a periodically buckled topology. The blue phosphorene is a semiconductor with an indirect band gap of $\sim 2 \text{ eV}$.

Along this line a theoretical work based on first principles calculation proposes the use of blue phosphorene nanoribbons\cite{17} to modulate the band gap, in this case the gap decreases for narrower nanoribbons. Nonetheless the system remains an indirect band gap semiconductor for either armchair or zigzag edges.

Another way to modulate the physical properties of blue phosphorene is changing the stacking. Two blue phosphorene layers stacked in AA or AB fashion have also shown a decrease in the band gap compared with the blue phosphorene monolayer. Unfortunately the band gap is also indirect\cite{18} which is a real limitation to use this 2D allotrope in the solar cell industry due to the poor efficiency that characterizes indirect band gap semiconductors\cite{19}.

In this article we show that twisted bilayer blue phosphorene (TBBP) is a direct band gap semiconductor at least in three of the four structures studied. The relative rotation angle (RRA) between layers modulates the value of the gap. The band gap goes from 0.95 eV for a RRA of 7.3° to a value of 1.6 eV for an angle of 21.8°. Unlike stacked AA and AB bilayer blue phosphorene, TBBP has absorption peaks in the visible range of the absorption spectra, and additionally the presence of flat valence bands in TBBP structures could improve the optical absorption probability. These findings makes TBBP a good candidate for a solar cell device, since an ideal solar cell device would be composed of a battery of stacked direct band gap semiconductors, each of them with a different value of the band gap and hopefully around 1.5 eV to take advantage of the solar spectrum\cite{20, 21}.

2. Method

2.1. Geometry of the Unit cell

Crystalline blue phosphorene is composed of an hexagonal lattice with a basis of two P atoms like the honeycomb structure of graphene. The results of relaxation processes determined by \textit{ab initio} methods shows that the two atoms are not in the same plane. In fact, there is a vertical shift between them of 1.24 Å\cite{18}. On the other hand, one of the ways of stacking two layers is in the Bernal or AB stacking. In bilayer graphene it is the most stable configuration. The unit cell of stacked AB blue phosphorene will have four atoms, two in each layer. One of the atoms at the top layer will have directly below it an atom of the bottom layer whereas the second atom of the upper layer will be located exactly above the center of a hexagon of the layer beneath, as shown in Fig 1 (a).

We consider a system composed of two coupled blue phosphorene layers when there is a RRA between them. The unit cell of the rotated structure was constructed following a procedure exposed in several previous works\cite{8, 22}. 

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Starting from a stacked AB bilayer one rotates one layer from a point in the lattice to an equivalent location. This can be done by taking a vector \( \vec{r} = m\vec{a}_1 + n\vec{a}_2 \) and rotating it around a line perpendicular to the layers passing through the origin to an equivalent position at \( \vec{t} = n\vec{a}_1 + m\vec{a}_2 \), where \( \vec{a}_1 \) and \( \vec{a}_2 \) are the monolayer lattice vectors; \( n, m \) are integers. This way the rotational angle and all the relevant geometric information of the bilayer twisted cell can be expressed in terms of \( n \) and \( m \).

In Figure 1(a) we show the stacked AB blue phosphorene, in panel (b) a lateral view of the two layers where the buckled structure can be observed, and in panel (c) the twisted unit cell obtained doing a commensurate rotation. In this case we selected a hollow site of the stacked AB bilayer as a center of rotation, which is also the origin of the twisted unit cell. The hollow site is the point where an atom of the bottom layer have directly above a center of an hexagon of the top layer.

2.2. DFT calculations

The calculations were performed within the density functional theory (DFT) approach, as implemented in the SIESTA code\[23\]. The Kohn-Sham orbitals were expanded in a localized orbitals basis set (double-\( \xi \), single polarized in the present work). The twisted blue phosphorene bilayers were simulated considering cells with 28 (TBBP-28), 76 (TBBP-76), 148 (TBBP-148) and 244 (TBBP-244) atoms, corresponding to RRA (\( \theta \)) of 21.8°, 13.2°, 9.4°, and 7.3° respectively, and with a vacuum region of 20 Å. For the exchange-correlation potential we employed vdw-KBM approach\[24\], that includes the van der Waals (vdW) interactions within the self-consistent process. All structures were fully relaxed considering a force convergence tolerance of 0.01 eV/Å per atom. The Brillouin zone (BZ) was sampled by a \( 3 \times 3 \times 1 \) Monkhorst-Pack mesh that ensure the total energy convergence. The optical absorption spectrum is obtained through the imaginary part of the dielectric function, implemented in the SIESTA code using the dipolar approximation. This approximation does not take into account the many body effects as GW corrections and the electron-hole interaction. These calculations are prohibitive for such a big systems as TBBPs. So that, the dipolar approach is certainly a good first order approximation to understand the physical properties of TBBPs, in particular the optical absorption spectrum. All optical spectra were calculated with a Gaussian smearing of 0.06 eV. To ensure the convergence of the optical spectrum and the total density of states we employed a k-point grids of \( 51 \times 51 \times 1 \) for AA and AB stacking, \( 20 \times 20 \times 1 \) for RRAs of 21.8° and 13.2°, and \( 11 \times 11 \times 1 \) for RRAs of 9.4° and 7.3° respectively.

3. Results

We have calculated the geometrical structure, the band structure, the total density of states, and the optical absorption of monolayer BP, bilayers stacked AA and AB of BP and TBBP with different RRA. The main geometric parameters and the band gap for each system are shown in the table 1. For the
Table 1: Geometric parameters, band gap, and binding energy of several blue phosphorene configurations. \(a\) - lattice constant, \(d\) - maximal distance between layers and \(E_b\) - binding energy by atom.

| System       | \(a (\text{Å})\) | Gap (eV) | \(d (\text{Å})\) | \(E_b (\text{eV/atom})\) |
|--------------|-----------------|---------|-----------------|-----------------|
| Monolayer    | 3.37            | 2.098   |                 |                 |
| Bilayer AA   | 3.38            | 0.976   | 3.185           | −0.027          |
| Bilayer AB   | 3.41            | 0.645   | 2.993           | −0.017          |
| Twisted 21.8°| 8.91            | 1.330   | 3.188           | −0.021          |
| Twisted 13.2°| 14.671          | 1.183   | 2.996           | −0.022          |
| Twisted 9.4° | 20.489          | 1.068   | 2.917           | −0.022          |
| Twisted 7.3° | 26.320          | 0.949   | 2.933           | −0.022          |

BP monolayer the results obtained by us for the lattice constant, and the band gap are consistent with a previous SIESTA calculations\[^{[16]}\]. In the case of BP stacked AA and AB the results are in agreement with a previous calculation\[^{[18]}\], where PBE and DFT-D2 corrections were employed to incorporate a vdW interaction. There, the authors obtained a distance of 3.23 and 3.20 Å for AA and AB stacking, respectively. Our results for the binding energy of all structures show that the AA staking is more favourable than the AB and TBBP with a binding energy per atom of −0.027 eV. In the case of TBBP, the RRA does not make any perceptible variation in the binding energy per atom, the value obtained for all structures were −0.022 eV/atom. This results suggest that the TBBPs are more stable than the AB staking regardless of the RRA.

In Figure 2 we show the results of the DFT calculations of the band structure for monolayer and bilayer staked AA and AB of blue phosphorene, the red arrow indicates the indirect band gap. The three structures are indirect band gap semiconductors, the size of the gap varies from \(\sim 0.6\) to 2 eV.

The density of states (DOS) of the same three structures (Figure 3(a)) shows clearly the evolution of the size of the band gap. The top valence band in the two bilayer systems is almost flat around the \(\Gamma\) point of the BZ, and the vHs below the Fermi level can be associated with this valence band. A strong absorption of a material at a given energy is usually associated with the presence of vHs in the DOS. The absorption spectra in Figure 3(b)) show strong absorptions peaks for energies around \(\sim 4\) eV, but below this energy and above the absorption edges there is not any strong absorption peak whatsoever. This fact is certainly similar to what is observed in other hexagonal flat structures as graphene which is transparent in the visible spectrum range.

We were interested to see what happens when we rotate the layers. According to our previous experience, many fascinating angle dependent properties may appear. We have calculated the band structure of four twisted structure by DFT, and the results are shown in the Figure 4. It is possible to observe that the rotation angle changes the electronic properties of the system. in fact, for certain values of the relative rotation angle, the bilayer blue phosphorene becomes a direct band gap semiconductor. At least in three of the four structures
the minimum distance between top valence band an lower conduction band lies at the same k-point. However the top valence band is almost flat throughout the whole BZ and, as long as the angle is smaller this effect is reinforced. One might explain such a behaviour as a result of the shrinking of the BZ. The flat region around the Γ point observed in stacked AB bilayer have become relatively larger in the reduced zone of the twisted structures. As a result of this band flatness, optical absorption peaks will appear near the band gap value with amplitudes larger than those appearing in the stacked AB and AA structures.

The value of the band gap changes with the angle. Likely, more results are needed in order to establish the full variation pattern but in accordance with the four cases investigated here it is possible to advance the decrease of the band gap as a function of the rotation angle. In table 1 the values of the band gap for the angles studied are listed. It has a value of 1.33 eV for the larger angle, 21.8° and goes to 0.95 eV for a RRA of 7.3°. The distance between layers in the relaxed structures diminishes slightly as the angle gets lower. The latter is expected since our starting point was a stacked AB bilayer.

The DOS of the twisted structures, Figure 5(a), express in another format the facts we were mentioning; that is, the slight dependence of the band gap with the angle and the presence of vHs just in the band edge of the valence bands. The absorption spectra in Figure 5(b) reveal the presence of absorption peaks below 2 eV, the position of the peaks changes with the angle and follows in principle the same behaviour of the band gap. The intensity of the peaks diminishes also for lower angles.

It is important to notice that we have not included many-electron effects in our calculations. The presence of these flat bands which is in many cases related with the appearance of many body effects as well as the difference of the experimental optical band gap with the theoretical (single electron calculations) in its cousin bilayer black phosphorene lead us to suggest that further studies of these structure should be made considering many body effects. However the task is complicated due to the large size of the unit cell.

4. Summary and conclusions

We have studied by DFT calculations the electronic structure of two coupled layers of blue phosphorene and found that a relative rotation between layers changes the electronic properties of the system. Bilayer stacked AA or AB blue phosphorene are indirect band gap semiconductor while some of the twisted structures are direct band gap semiconductor and moreover the size of the band gap depends on the RRA. The absorption spectra show peaks in the visible range of the spectra which makes twisted bilayer blue phosphorene another candidate for solar cell applications. The dependence of the band gap with the angle suggests that stacking several of these structures together will increase the efficiency in solar cell devices as it will cover a wider range in energy of the solar spectrum as well as a larger interface.
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Figure 1: (a) Two stacked AB blue phosphorene layers. (b) Lateral view of the two layers in AB stacking. (c) Twisted structure for a relative rotating angle of $9.34^\circ$. 
Figure 2: Bands structure for three different blue phosphorene structures (a) monolayer, (b) stacking AA and (c) stacking AB. The red arrow shows the indirect band gap.
Figure 3: Total density of states for three different blue phosphorene structures (a) monolayer, (b) stacking AA and (c) stacking AB. Panel (d) is for the imaginary part of the dielectric function for monolayer, stacking AA, and stacking AB of blue phosphorene, the insert in the panel (d) shows the imaginary part of the dielectric function in different incident photon energy range.
Figure 4: Bands structure of four twisted bilayer blue phosphorene structures, in each panel a different rotating angle (a) $\theta=21.8^\circ$, (b) $\theta=13.2^\circ$, (c) $\theta=9.4^\circ$ and (d) $\theta=7.3^\circ$. The (red vertical) arrows show the band gap.
Figure 5: Total density of states of four twisted bilayer blue phosphorene structures, in each panel a different rotating angle (a) $\theta=21.8^\circ$, (b) $\theta=13.2^\circ$, (c) $\theta=9.4^\circ$ and (d) $\theta=7.3^\circ$. Panel (e) is for the imaginary part of the dielectric of four twisted bilayer blue phosphorene structures, the insert in the panel (e) shows the imaginary part of the dielectric function in different incident photon energy range.