Molecular doping of blue phosphorene: a first-principles investigation

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Abstract
Using first-principles calculations, we show that p-doped blue phosphorene can be obtained by molecular doping with 2,3,5,6-tetrafluoro-7,7,8,8-tetracyanoquinodimethane (F4-TCNQ) and 1,3,4,5,7,8-hexafluorotetracyanonaphthoquinodimethane (F6-TNAP), whereas n-doped blue phosphorene can be realized by doping with tetrathiafulvalene (TTF) and cyclooctadecanonaene (CCO). Moreover, the doping gap can be effectively modulated in each case by applying an external perpendicular electric field. The optical absorption of blue phosphorene can be considerably enhanced in a broad spectral range through the adsorption of CCO, F4-TCNQ, and F6-TNAP molecules, suggesting potential of the doped materials in the field of renewable energy.

Keywords: molecular doping, blue phosphorene, doping gap, absorption, first-principles

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monolayer MoS2 is drastically enhanced by adsorption of doping can also be used to tune the optical properties of 2D materials without perturbing the hand structure (in contrast to traditional atomic doping) [37, 54, 56]. While monolayer graphene grown by ambient-pressure chemical vapor deposition shows p-type characteristics, transition to n-type characteristics can be realized by adsorption of piperidine, making it possible to form a graphene-based p-n junction [42]. Molecular doping can also be used to tune the optical properties of 2D materials. For instance, the photoluminescence intensity of monolayer MoS2 is drastically enhanced by adsorption of 2,3,5,6-tetrafluoro-7,7,8,8-tetracyanoquinodimethane (F4-TCNQ) or 7,7,8,8-tetracyanoquinodimethane (TCNQ) [45]. It can be enhanced by a factor of nine by doping with salicylic acid molecules [52]. While these investigations show that molecular doping is a very powerful approach to tune 2D materials and broaden their range of applications, the technique was not applied to BlueP so far.

In the present work, we therefore explore the effects of molecular doping on BlueP, using first-principles calculations. In addition, application of an external electric field is considered as a tool to modify the electronic properties. Four organic molecules (atomic structures illustrated in figure 1, see also figure S1 in the supporting information [stacks.iop.org/JPhysCM/32/055501/mmedia]) are investigated: tetrafluoro-tetrathiafulvalene (TTF) and cyclooctadecanonaene (CCO) as examples of electron donors (low ionization potentials of 6.83 and 7.23 eV, respectively [63, 64]), and F4-TCNQ and 1,3,4,5,7,8-hexafluorotryptacnaphthoquinodimethane (F6-TNAP) as examples of electron acceptors (high electron affinities of 5.24 and 5.37 eV, respectively [65, 66]). We will demonstrate that adsorption of these molecules can yield p- and n-doping without breaking the 2D structure of BlueP. In addition, the ability to absorb sun light can be enhanced significantly, which is critical for the application of BlueP-based materials in energy harvesting.

Computational details

First-principles calculations are performed using the Vienna ab-initio simulation package based on plane-wave density functional theory and the projector-augmented wave method [67]. The energy cutoff of the plane-wave expansion is set to 450 eV and the exchange-correlation functional is treated in the generalized gradient approximation (Perdew–Burke–Ernzerhof form). To describe the long-range interaction between BlueP and organic molecules, the vdW-D3 correction of Grimme is used [68]. The optB86b-vdW functional [69] is found to result in unreasonably large adsorption heights. A large 6 × 6 × 1 supercell is adopted with a vacuum region thicker than 15 Å to eliminate artificial interaction between periodic images. One molecule is added to the supercell, which corresponds to an areal density of 3 × 1013 cm−2. We can neglect substrate effects, because the binding energy between BlueP and the usual substrate, Au(1 1 1), is only 0.11 eV/atom [70]. Brillouin zone sampling on a Monkhorst-Pack 3×3×3 k-point mesh provides good convergence. In the structure optimization the atomic coordinates are relaxed until the Hellmann–Feynman forces are reduced to 0.01 eV/Å. Absorption spectra are calculated using the Heyd–Scuseria–Ernzerhof (HSE06) hybrid functional [71]. The imaginary part of the dielectric tensor is obtained by neglecting local field effects and approximating the macroscopic dielectric function by the head of the microscopic dielectric matrix [72]. The real part of the dielectric tensor then results from Kramers–Kronig transformation.

Results and discussion

We obtain for pristine BlueP an optimized lattice constant of 3.277 Å. Previously, Zhu et al [1] predicted a lattice constant of 3.33 Å, while Bao et al [20] obtained a value of 3.29 Å. Thus, our calculated lattice constant is in satisfactory agreement with previous results, demonstrating reliability of the methods used in the present work. We determine the

| Adsorption site | E_{ad} (eV) | h (Å) | ΔQ (e) | n (10^{12} e/cm^2) |
|----------------|------------|-------|--------|-------------------|
| BlueP-TTF      | 6          | 0.87  | 3.08   | −0.11             | 3.3 |
| BlueP-CCO      | 6          | 1.12  | 3.50   | −0.04             | 1.2 |
| BlueP-F4-TCNQ  | 2          | 0.87  | 3.51   | 0.12              | 3.6 |
| BlueP-F6-TNAP  | 2, 6       | 1.11  | 3.37   | 0.16              | 4.8 |

Table 1. Adsorption energy (E_{ad}), adsorption height (h), electron transfer (ΔQ), and injected carrier concentration (n; ΔQ per area).

Figure 1. Top views of the atomic structures of TTF, CCO, F4-TCNQ, and F6-TNAP.

Figure 2. Top (upper panel) and side (lower panel) views of the favorable adsorption configurations of TTF, CCO, F4-TCNQ, and F6-TNAP molecules on BlueP. The orange and blue sticks represent upper and lower P atoms, respectively. The white, black, red, green, and yellow spheres represent H, C, N, F, and S atoms, respectively.
favorable adsorption configuration for each molecule (table 1, figure 2) by calculating the adsorption energy $E_{\text{ad}} = E_{\text{molecule}} + E_{\text{BlueP}} - E_{\text{BlueP+molecule}}$, where $E_{\text{molecule}}$, $E_{\text{BlueP}}$, and $E_{\text{BlueP+molecule}}$ are the energies of the molecule, pristine BlueP, and BlueP with adsorbed molecule, respectively. For each molecule, several high-symmetry adsorption sites are considered (figure S2 in the supporting information).

The TTF molecule favors alignment parallel to the zigzag direction of BlueP with $E_{\text{ad}} = 0.87 \text{ eV}$ and adsorption height $h = 3.08 \text{ Å}$. The electron-abundant C$_3$S$_2$ rings located above P atoms. The CCO molecule favors a high symmetry configuration with space group C$_{3v}$ ($E_{\text{ad}} = 1.12 \text{ eV}$, $h = 3.50 \text{ Å}$). The aromatic ring of the F$_4$-TCNQ molecule is located directly on top of a P atom to maximize the overlap between its delocalized $\pi$ electrons and the p electrons of the P atom. In addition, two cyano groups cross the P–P bonds, increasing the electron transfer and binding strength ($E_{\text{ad}} = 0.87 \text{ eV}$ and $h = 3.51 \text{ Å}$). Finally, for the F$_6$-TNAP molecule there exist two adsorption sites with $E_{\text{ad}} = 1.11 \text{ eV}$ (2 and 6 in figure S1(d) with $h = 3.37$ and 3.43 Å, respectively). We choose the configuration presented in figure 2 for the following calculations.

Charge transfer in the molecule-doped materials may alter the transport and optical performance of BlueP, as revealed for the F$_4$-TCNQ/MoS$_2$ interface in an earlier study [45]. To investigate this question, figure 3 shows the charge density difference isosurfaces obtained for BlueP with adsorbed TTF, CCO, F$_4$-TCNQ, and F$_6$-TNAP molecules. The pink regions denote accumulation of electrons and the gray regions denote depletion of electrons. In the cases of TTF–BlueP and CCO–BlueP (figures 3(a) and (b)) electrons are transferred from the highest occupied molecular orbital (HOMO) to BlueP.
Figure 5. Doping gaps and electron transfer in (a) TTF–BlueP, (b) CCO–BlueP, (c) F4-TCNQ–BlueP, and (d) F6-TNAP–BlueP as functions of the electric field. The insets show the structure under an electric field of strength ±0.5 V Å⁻¹ and the orange arrows indicate the positive direction of this field.

Figure 6. Projected band structures of (a) TTF–BlueP and (b) CCO–BlueP under an electric field of –0.5 V Å⁻¹, and of (c) F4-TCNQ–BlueP and (d) F6-TNAP–BlueP under an electric field of 0.5 V Å⁻¹. Red color highlights the contributions of the molecules.
(P atoms in the contact region), i.e. the TTF and CCO molecules act as electron donors. Bader analysis [73–75] shows that 0.11 and 0.04 electrons are transferred from the TTF and CCO molecules to BlueP, respectively. In the cases of F4-TCNQ–BlueP and F6-TNAP–BlueP (figures 3(c) and (d)) electrons are transferred from BlueP to the lowest unoccupied molecular orbital (LUMO), i.e. the F4-TCNQ and F6-TNAP molecules act as electron acceptors. The transferred electrons are found mainly in the interlayer region between molecule and BlueP as well as on the cyano groups of the F4-TCNQ and F6-TNAP molecules. Large gray isosurfaces at P atoms near the indirect bandgap of 1.94 eV (figure 4(a)), in agreement with TCNQ than that of TTF (between CBM and HOMO for n-dopants, between LUMO states due to the HOMOs of TTF and CCO. The doping gap is 0.11 and 0.04 electrons are transferred from BlueP to the F4-TCNQ and F6-TNAP molecules, respectively. The electron transfer from/to the TTF, CCO, F4-TCNQ, and F6-TNAP molecules leads to a carrier concentration of \(3.3 \times 10^{12}\), \(1.2 \times 10^{12}\), \(3.6 \times 10^{12}\), and \(4.8 \times 10^{12}\) cm\(^{-2}\) in BlueP.

Figure 4 shows a projected band structure of pristine BlueP in comparison to results for TTF–BlueP, CCO–BlueP, F4-TCNQ–BlueP, and F6-TNAP–BlueP. Pristine BlueP has an indirect bandgap of 1.94 eV (figure 4(a)), in agreement with the findings of previous investigations [2, 3]. The band structures of TTF–BlueP (figure 4(b)) and CCO–BlueP (figure 4(c)) resemble that of pristine BlueP. However, additional flat bands emerge at 0.27 and 0.97 eV, respectively, i.e. below the conduction band minimum (CBM) of BlueP, reflecting n-doping of the host material. The flat bands represent localized electronic states due to the HOMOs of TTF and CCO. The doping gap (between CBM and HOMO for n-dopants, between LUMO and VBM for p-dopants) of TTF–BlueP is significantly smaller than that of TTF–BlackP (0.73 eV) [56]. For F4-TCNQ–BlueP (figure 4(d)) and F6-TNAP–BlueP (figure 4(e)) the LUMOs of F4-TCNQ– and F6-TNAP appear at 0.39 and 0.45 eV, respectively, i.e. above the valence band maximum (VBM) of BlueP, reflecting p-doping of the host material.

The n-dopants TTF and CCO as well as the p-dopants F4-TCNQ and F6-TNAP induce a large doping gap in BlueP, which makes them to ineffective doping molecules. We next investigate whether an external perpendicular electric field can help to overcome this issue. The dependence of the doping gap and electron transfer on the applied electric field is shown in figure 5 (results after re-relaxing the atomic coordinates in the electric field, which, however, induces only minor structural changes). The strength of the electric field ranges from \(-0.5\) to \(0.5\) V Å\(^{-1}\) with the positive direction oriented from the molecule to BlueP, see the insets of figure 5. We note that the change in bandgap of pristine BlueP is negligible under such an electric field [3]. For TTF–BlueP and CCO–BlueP the electron transfer is reduced for increasing electric field, i.e., the HOMOs of the molecules shift toward the CBM of BlueP and the doping gaps increase monotonically. Under an electric field of \(-0.5\) V Å\(^{-1}\) (figures 6(a) and (b)) the doping gaps are as small as 79 and 94 meV, respectively, indicating that shallow acceptor states are formed in both F4-TCNQ–BlueP and F6-TNAP–BlueP, resembling typical p-type semiconductors. We note that the HSE06 functional enhances the doping gaps, while maintaining the general trends described above.

In addition to enhancing the effectiveness of doping, applying an electric field also supports the charge carrier injection and therefore can be beneficial for the device performance. More specifically, according to figure 5, 0.44 electrons are transferred from TTF to BlueP, which leads to an n-type carrier concentration of \(1.31 \times 10^{13}\) cm\(^{-2}\) in BlueP. Transfer of 0.41 and 0.46 electrons from BlueP to F4-TCNQ and F6-TNAP, respectively, leads to p-type carrier concentrations of \(1.22 \times 10^{13}\) and \(1.37 \times 10^{13}\) cm\(^{-2}\) in BlueP.

Absorption spectra are shown in figure 7 together with the incident AM1.5g global standard spectrum. Pristine BlueP does not exhibit strong optical absorption in the visible region and, therefore, is not a good photovoltaic material. Similar results are obtained for TTF–BlueP. On the other hand, CCO–BlueP shows a pronounced absorption peak (intensity \(7.2 \times 10^6\) cm\(^{-1}\)) in the visible region centered at 534 nm. For F4-TCNQ–BlueP we obtain more enhancement of the absorption in the visible region with a broader absorption peak (intensity \(6.2 \times 10^6\) cm\(^{-1}\)) centered at 641 nm. Finally, for F6-TNAP–BlueP realizes a very broad absorption peak (intensity \(6.2 \times 10^6\) cm\(^{-1}\)) centered at 877 nm. Apart from these additional absorption peaks, which are mainly due to molecular transitions, we observe for all doped materials a redshift of the absorption spectrum with respect to pristine BlueP, which enlarges the overlap with the incident solar flux and thus further enhances their ability to absorb sun light. To quantify the performance of the materials under investigation, we calculate the efficiency as

\[
\eta = \int_0^{\infty} a(\lambda) S(\lambda) d\lambda,
\]

where \(\lambda\) is the wavelength, \(a(\lambda)\) is the absorption spectrum, and \(S(\lambda)\) is the incident AM 1.5g global standard spectrum. We obtain for TTF–BlueP,
COC–BlueP, F4-TCNQ–BlueP, and F6-TNAP–BlueP values of 10.6, 37.2, 24.1, and 28.8 W cm\(^{-2}\), respectively. In particular, both COO–BlueP and F6-TNAP–BlueP therefore turn out to be superior to T2-TCNQ–Arsenene (28.0 W cm\(^{-2}\)) [76].

Conclusions

Our investigation of molecular doping of BlueP, based on first-principles calculations, shows that TTF and COO molecules act as electron donors and lead to n-doping of BlueP while F2-TCNQ and F6-TNAP molecules act as electron acceptors and lead to p-doping of BlueP. The amount of electron transfer, which determines the carrier dynamics and thus the device performance, has been quantified. It turns out that an external perpendicular electric field shifts the HOMO of TTF closer to the CBM of BlueP, resulting in a shallow donor state and effective n-doping of BlueP. Similarly, the LUMOs of F2-TCNQ and F6-TNAP can be shifted closer to the VBM of BlueP, resulting in shallow acceptor states and effective p-doping of BlueP. The application of an electric field also enhances the charge transfer and thus the charge carrier injection into BlueP. COO, F4-TCNQ, and F6-TNAP molecules induce redshifts of the absorption spectrum of BlueP and enhance the absorption in the visible and infrared regions, which is beneficial for solar energy harvesting. Due to a lack of experimental studies, our results provide important insights into the electronic and optical properties achievable by molecular doping of BlueP. They offer guidelines for the design of electronic, optoelectronic, and photovoltaic devices.

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Supporting information

Structural properties of TTF, COO, F4-TCNQ, and F6-TNAP. Studied adsorption configurations of TTF, COO, F4-TCNQ, and F6-TNAP molecules on BlueP.

Notes

The authors declare no competing financial interest.

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