Two-dimensional (2D) crystals exhibit unique and exceptional properties and show promise for various applications. In this work, we systematically studied the structures of a 2D boronphosphide (BP) monolayer with different stoichiometric ratios (BP$_x$, $x = 1, 2, 3, 4, 5, 6$ and $7$) and observed that each compound had a stable 2D structure with metallic or semiconducting electronic properties. Surprisingly, for the BP$_5$ compounds, we discovered a rare penta-graphene-like 2D structure with a tetragonal lattice. This monolayer was a semiconductor with a quasi-direct band gap of 2.68 eV. More importantly, investigation of the strain effect revealed that small uniaxial strain can trigger the band gap of the penta-BP$_5$ monolayer to transition from a quasi-direct to direct band gap, whereas moderate biaxial strain can cause the penta-BP$_5$ to transform from a semiconductor into a metal, indicating the great potential of this material for nanoelectronic device applications based on strain-engineering techniques. The wide and tuneable band gap of monolayer penta-BP$_5$ makes it more advantageous for high-frequency-response optoelectronic materials than the currently popular 2D systems, such as transition metal dichalcogenides and black phosphorus. These unique structural and electronic properties of 2D BP sheets make them promising for many potential applications in future nanodevices.
Another approach to further open the band gap of a structure involves modifying its basic configuration because the properties are closely related to the structural configurations. Graphene generally has a hexacyclic configuration. Surprisingly, Zhang et al. recently reported a very rare penta-graphene starting from the pure pentacyclic configuration. This penta-graphene is not only dynamically and mechanically stable but also has a wide band gap (3.25 eV), unusual negative Poisson’s ratio, and ultrahigh ideal strength. Subsequently, more attention has been paid to this approach, which has been extended to other systems, such as CN, B–N, Si–H, B2C and AlN. A corresponding stable penta-2D structure with a wide band gap above 2.0 eV in each compound was observed. Therefore, it is necessary to find a suitable penta-2D material to obtain a large-band-gap semiconductor.

As a typical semiconducting material, boronphosphide (BP) has also attracted considerable attention. Experimentally, bulk BP is the only known stable compound in the B–P system. Bulk BP exhibits many outstanding semiconducting properties with an indirect wide band gap, resulting in its wide applications in solid-state neutron detectors. BP films have been synthesised on silicon carbide using chemical vapour deposition. Theoretically, all the previously predicted 2D BP structures have been constructed from hexagonal configurations, and all of them have a band gap in the range of 1–1.8 eV. To date, no 2D material with a band gap larger than 2.0 eV has been reported in the B–P system. Therefore, determining whether a 2D single-layer structure composed of pure pentagons with a larger band gap exists in the B–P system is of interest. Moreover, current experimental and theoretical studies have focused on the 1:1 ratio of B to P, and the search for a 2D structure for other compounds remains lacking. Hence, whether stable 2D monolayer structures with excellent properties exist in other ratios of B and P remains to be determined.

To address these issues, we systematically studied the structures of 2D BP monolayers with different stoichiometries, including BP, BP2, BP5, BP6, and BP7, using particle swarm optimisation (PSO) combined with ab initio molecular dynamics (MD) calculations. The simulation results indicate that each compound has a stable 2D structure. Surprisingly, for BP5, we observed a rare penta-graphene-like structure with a quasi-direct band gap of 2.68 eV. Moreover, the 2D materials with ratios of 1:3, 1:6, and 1:7 exhibited semiconducting properties with indirect band gaps of 0.8–2 eV, whereas the 2D materials with 1:2 and 1:4 ratios exhibited metallic properties.

Results and Discussion

Structural properties. Six different compositions of 2D BP compounds were considered using CALYPSO, including 1:1, 1:2, 1:3, 1:4, 1:5, 1:6 and 1:7. The simulation results indicated that each compound contains a series of 2D structures. Here, we focus on the 1:5 compound.

Figure 1 shows the optimised 2D structures of the 1:5 compound. This penta-2D structure has a tetragonal lattice with a lattice parameter of a = b = 4.54361 Å and space group P-421c (No. 114). This structure is similar to that of the previously reported penta-graphene structure, which possesses P-42m symmetry (space group No. 113). As observed in the top view of penta-BP5 in Fig. 1a, the structure is constructed of pure pentagons, where each pentagon includes one B atom and four P atoms, forming the famous Cairo pentagonal tiling. In the side view of penta-BP5 (Fig. 1c), buckling is observed. The “thickness” of this sheet is 2.50 Å, which is the vertical coordinate difference between P atoms in the top and bottom layers. The “thickness” of penta-BP5 is larger than...
that of penta-graphene (1.20 Å) and other penta-2D materials. The lattice parameters and atomic positions are listed in Table S1.

In Fig. 1(b–d), the electron localisation function (ELF) was calculated using the Perdew–Burke–Ernzerhof (PBE) method. In the structure, all the B atoms are equivalent, forming a four-coordinated sp³ hybrid. There are two equivalent positions for the P atom, including the formation of four and three coordinates. Analysis of the ELF reveals that all the P atoms are also sp³ hybrid; however, there is a lone pair of electrons for the three-coordinate case. It is an all-sp³ electronic structure that leads to the semiconducting properties.

Dynamical stability. To examine the dynamical stability of the 2D structure, we calculated the phonon spectra, as shown in Fig. 2. No imaginary modes are observed in the first Brillouin zone for penta-BP₅, indicating its dynamical stability.

For the kinetically stable pentagonal BP₅ monolayer, ab initio MD simulations were performed. A large supercell (5×5) was employed by heating the structure to 300 K, 450 K, and 1000 K. For each case, the simulation duration was 10 ps with a step of 1 fs. At the end of each simulation, the final structure was carefully examined. As observed in Fig. 3, the penta-BP₅ could withstand temperatures as high as 1000 K. At high temperatures, the structure became slightly distorted. However, these distortions were not sufficient to destroy the B–P and P–P bonds, and the original structure could be restored by global optimisation. Therefore, the phonon calculations and MD simulations indicated that monolayer penta-BP₅ possesses high dynamical stability.

Mechanical properties. Excellent strain strength is indispensable for an ideal nanomaterial, and the stress–strain curve is an extremely important physical quantity that characterises the mechanical properties of...
nomenon. The valence band maximum (VBM) of the single-layer penta-BP$_5$ lies at a point along the G–X line, comparable to well-known 2D materials such as graphene$^{30}$ and MoS$_2$.$^{31}$ Next, we investigated the case of uniaxial strain. As the x and y axes are equivalent, we introduced the uniaxial strain along the y-axis, and the x-directional lattice was allowed to relax freely$^{32}$. Figure 4b shows the dependence of stress on uniaxial strain. We can see that maximum stress of the structure is 2.93 N/m at uniaxial strain of 18%, indicating that the structure has a high uniaxial tensile capacity.

**Electronic properties.** We calculated the band structures and orbital projected density of states (PDOS) using the PBE method. The results indicate that the penta-BP$_5$ bulk (Fig. 5a) is an indirect semiconductor with a band gap of 1.22 eV. Examination of the band structure of the monolayer (Fig. 5c left) reveals an interesting phenomenon. The valence band maximum (VBM) of the single-layer penta-BP$_5$ lies at a point along the G–X line, and the conduction band minimum (CBM) lies at a point along the M–G line. The energy difference between the second-lowest point of the conduction band (along the G–X line) and the CBM is very small (0.03 eV), such that the monolayer structure can be considered a quasi-direct semiconductor with a band gap of 1.84 eV. A similar phenomenon is also observed in the Si–Ge super lattice system$^{33}$. The PDOS of the monolayer is shown in Fig. 5c (right). Hybridization is observed in the 3s and 3p orbitals of the P atom and 2s and 2p orbitals of the B atom over the entire energy range, indicating that there is a covalent interaction between the P–P and B–P bonds.

Hybrid density functional (HSE06) calculations provide a better description of band gaps in semiconductors; as observed in Fig. 5b and d, the band gaps of the bulk and monolayer penta-BP$_5$ are transformed to 1.94 and 2.68 eV, respectively. Therefore, 2D penta-BP$_5$ is expected to be a promising candidate for photodetectors harvesting photons with wavelengths of less than 620 nm, such as blue and UV LEDs and photodetectors. Note that previous pentagonal layered structures have possessed either large or indirect band gaps, which limits their application in optical devices. Here, we report a novel penta-2D material with a suitable quasi-direct band gap for the first time.

Strain technology is extensively applied to tune the band gap and electronic structures of semiconducting materials via lattice mismatch on the substrate, thermal expansion, or mechanical loading$^{34}$. First, we considered the biaxial strain effect. We observed a transition from semiconductor to metal for penta-BP$_5$ when the compressive strain was larger than $-12\%$, and the metal properties were retained up to $-20\%$. In the range of non-zero band gap, penta-BP$_5$ is an indirect band gap semiconductor. The detailed results are presented in the supplementary materials (see Supplementary Fig. S1). In contrast to biaxial strain, uniaxial strain can further change the structure of a 2D material, resulting in new properties$^{35}$. Therefore, we calculated the uniaxial strain to tune the electronic structure of the penta-BP$_5$ monolayer. In Fig. 6, the band gap was calculated as a function of uniaxial strain ($-20\%$ to $-20\%$) using the PBE exchange functional. The strain range can be divided into three parts: Indirect-I, Direct-II, and Indirect-III. In the strain range from $-4\%$ to $20\%$ (Indirect-I), the monolayer is an indirect semiconductor. With continued expansion, the band gap exhibits an increasing tendency and reaches a maximum (1.95 eV) at $4\%$ before decreasing nearly linearly to $20\%$. Interestingly, the monolayer undergoes a band-gap transition from quasi-direct to direct when a small compression ($-4\%$) is applied. Moreover, this direct band gap can be maintained up to $-12\%$ (Direct-II). Due to the low phonon energy of electron excitation, this strain-tunable direct band gap semiconductor has obvious advantages in the application of optical devices. Up on further increasing the compression, the indirect band gap reappears and is retained until $-20\%$ (Indirect-III).

Next, we mainly discuss the physical mechanism of the band gap transition from quasi-direct to direct. The band structures with different compressive strains are shown in Fig. 7. For convenience, we labelled the CBM point and second-lowest point in energy at 0% strain as A and B, respectively. When not subject to external forces, the structure exhibited the properties of a quasi-direct band gap semiconductor, that is, the energy of point A is slightly smaller than that of point B. Closer inspection reveals that with increasing compressive strain, the

![Image](image-url)
Figure 5. Band structures of penta-BP$_5$ bulk using (a) PBE and (b) HSE06 methods. (c) Band structures (left) and orbital PDOS (right) of the penta-BP$_5$ monolayer determined using PBE methods. (d) Band structures of penta-BP$_5$ monolayer determined using HSE06 methods. The Fermi energy was set to zero.

Figure 6. Variation of band gap with in-plane uniaxial strain for the penta-BP$_5$ monolayer using the PBE method. The inset shows the direction of strains.
energies of both point A and B decrease. However, the energy of point A decreases more rapidly and is consider-
ably lower than that of point B at a compressive strain of $-2\%$. In addition, the VBM and point A of the structure
have similar K-space coordinates. These phenomena indicate that the structure begins to transform into a direct
band gap semiconductor when compressive strains are applied. However, the energy of the VBM at state X slightly
changes. Therefore, the band gap is mainly determined by shifts of the CBM. Overall, the entire conduction band
has a tendency to approach the Fermi surface, which leads to the decrease of the band gap of the monolayer mate-
rial under the compressive stress. At $-4\%$, the penta-BP$_5$ completely transforms into a direct band gap semiconductor
with a band gap of 1.59 and 1.99 eV based on the PBE and HSE06 methods, respectively. This appropriate
direct band gap makes this monolayer a promising candidate for solar cell materials.

In addition, we also studied other compounds of the BP system, including 2D structures with B to P ratios of
1:1, 1:2, 1:3, 1:4, 1:6, and 1:7. The results indicated that the 1:1 compound has a graphene-like structure, which is
consistent with previous studies$^{25-27}$. For the other compounds, we observed a series of stable 2D layered struc-
tures in each compound. The 2D materials for the 1:2 and 1:4 compounds exhibited metal properties, whereas
those for the 1:3, 1:6, and 1:7 compounds exhibited semiconductor characteristics. All the 2D monolayer structures
were dynamically stable, as confirmed by phonon dispersions. These 2D semiconducting materials were
observed to be indirect band gap semiconductors with a band gap range of 0.8–2.0 eV (HSE06 calculations),
indicating that these monolayer structures can be used as optical devices. The optimised structures and corre-
sponding phonon spectral and band structures are presented in Supplementary Figs S2–S6.

Conclusions
In summary, we performed a systematic search for stable 2D materials in the B–P system using the $ab$ $initio$
PSO methodology, including compounds with B to P ratios of 1:1, 1:2, 1:3, 1:4, 1:5, 1:6 and 1:7. A stable 2D structure
was observed for each compound. Surprisingly, we observed a rare penta-graphene-like structure, penta-BP$_5$,
which is a semiconductor with a quasi-direct band gap of 2.68 eV (HSE06 method). The absence of an imagi-
nary mode in the phonon spectrum and a high melting point indicated that the penta-BP$_5$ monolayer exhibits
good dynamical stability. Stress–strain calculations demonstrated that the penta-BP$_5$ monolayer exhibits excellent
mechanical stability with breaking biaxial and uniaxial strains above 20% and 18%, respectively. More impor-
tantly, analysis of the strain effect revealed that small uniaxial strain can trigger a quasi-direct to direct band gap
transition in the penta-BP$_5$ monolayer, whereas moderate biaxial strain can cause the penta-BP$_5$ to transition
from a semiconductor to a metal. This wide and tuneable band gap of monolayer penta-BP$_5$ makes this structure
more advantageous in high-frequency-response optoelectronic materials than the currently popular 2D systems,
such as TMDCs and black phosphorus. In addition, the 2D materials in BP$_3$, BP$_4$, and BP$_5$ have semiconducting
properties with an indirect band gap of 0.8–2.0 eV, whereas the 2D materials in BP$_3$ and BP$_5$ have metallic proper-
ties. These unique structural and electronic properties of 2D B–P sheets make them promising for many potential
applications in future nanodevices.

Methods
The search for an energetically stable 2D B–P monolayer was performed by considering various stoichiometries
of BP$_x$ ($x = 1, 2, 3, 4, 5, 6$ and 7) using simulation cells containing up to four formula units. Structure searches
for all stoichiometries were performed using the PSO methodology as implemented in the CALYPSO cod.$^{35,36}$
Total energy calculations, geometrical optimisations, and electronic properties were computed using the Vienna
Ab Initio Simulation Package (VASP) program$^{37}$. Exchange and correlation of the electrons were treated by the

Figure 7. Strain-manipulated direct-to-indirect band gap in a 2D penta-BP$_5$ monolayer with compressive
strains of 0%, $-2\%$, $-4\%$ and $-6\%$. The Fermi energy was set to zero.
generalised gradient approximation (GGA) with the PBE. The B and P potentials have 2s22p1 and 3s3p3 as valence states, respectively, and the cutoff energy of the plane waves was 500 eV. In all the 2D structures, a vacuum distance of 20 Å was used to separate the periodic images in the perpendicular direction. All the structures were fully relaxed, and the energy was convolved to 1 meV per atom. To obtain highly accurate electronic structures, a hybrid HSE06 method was used. The first-principles MD simulations lasted 10 ps with a time step of 1 fs with the canonical ensemble (NVT). Phonon spectra were calculated using the Phonopy code.

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Author Contributions
B.B.L. designed the research. S.J.L. performed the calculations. S.J.L., B.L., X.H.S., J.Y.L., S.F.N., M.G.Y., Q.J.L., R.L., T.C., and B.B.L. analysed the data. S.J.L. and B.B.L. wrote the paper. All the authors discussed the results and commented on the manuscript.

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