First-principles DFT insights into the structural, elastic, and optoelectronic properties of $\alpha$ and $\beta$-ZnP$_2$: implications for photovoltaic applications

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First-principles DFT insights into the structural, elastic, and optoelectronic properties of α and β-ZnP₂: implications for photovoltaic applications

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Abstract

Binary II–V semiconductors are highly optically active materials, possess high intrinsic mechanical and chemical durability, and have electronic properties ideal for optoelectronic applications. Among them, zinc diphosphide (ZnP₂) is a promising earth-abundant absorber material for solar energy conversion. We have investigated the structural, mechanical, and optoelectronic properties of both the tetragonal (α) and monoclinic (β) phases of ZnP₂ using standard, Hubbard-corrected and screened hybrid density functional theory methods. Through the analysis of bond character, band gap nature, and absorption spectra, we show that there exist two polymorphs of the β phase (denoted as β₁ and β₂) with distinct differences in the photovoltaic potential. While β₁ exhibits the characteristics of metallic compounds, β₂ is a semiconductor with predicted thin-film photovoltaic absorbing efficiency of almost 10%. The α phase is anticipated to be an indirect gap material with a calculated efficiency limited to only 1%. We have also analysed and gained insights into the electron localization function, projected density of states and projected crystal orbital Hamilton populations for the analogue bonds between the α and β-ZnP₂. In light of these calculations, a number of previous discrepancies have been solved and a solid ground for future employment of zinc diphosphides in photovoltaics has been established.

Keywords: electronic structure, semiconductors, photovoltaics, density functional theory

Supplementary material for this article is available online
(Some figures may appear in colour only in the online journal)
Semiconducting II–V compounds are formed from comparatively inexpensive and non-toxic elements (e.g. Zn, N, P), which makes them attractive for large-scale PV applications. Amongst those, Zn₃P₂ and ZnP₂ are promising solar spectrum absorbers due to their favourable band gap energy values. Furthermore, they are mechanically and chemically stable, highly optically active (birefringence and rotation of the polarisation plane), possess long minority-carrier diffusion lengths, and have strong light absorption in the visible region [2–6]. Zinc diphosphide (ZnP₂), which crystallizes in two distinct modifications: tetragonal α-ZnP₂ (red) and monoclinic β-ZnP₂ (black), received little attention despite both phases having narrow band gaps and multivalley conduction and valence bands which result in high thermopower and, combined with the non-cubic symmetry, strong anisotropic optical properties with large birefringence [7]. Moreover, the power production potential of β-ZnP₂, based on the world resource estimates of its elemental constituents, is predicted to be 3.5 × 10² and 2.0 × 10⁵ times larger than InP and GaAs, respectively, for cells operating at 10% efficiency [8]. These features make zinc diphosphides prospective candidates for optoelectronic applications and next-generation solar energy conversion materials.

The development of ZnP₂ materials as efficient and cost-effective solar absorbers requires an atomic-level understanding of their fundamental properties (e.g. band gap energies, near-edge band structures, dielectric function, optical absorption coefficients). However, the basic mechanical and structural characteristics, as well as electronic and optical features of zinc diphosphides are poorly understood and inconclusive.

The crystal structure of α-ZnP₂ was first reported by Stackelberg and Paulus in 1935 [9] as a tetragonal cell containing eight formula units, with each Zn atom surrounded by four P atoms and each P atom surrounded by two Zn and two P atoms. This structure was experimentally confirmed in the 1960s and 1980s by a few groups [7, 10, 11], and again in 2003 by Zanin et al [12]. Elastic constants have been found to depend linearly on temperature in the range from 78 to 400 K, with occasional interruption areas of 10–20 K width [13]. From earlier experimental studies of optoelectronic properties of ZnP₂ materials, it is believed that the top of the valence band of α-ZnP₂ is located at the Γ point and the energy gap is indirect but dipole forbidden [14]. Pressure dependence experiments for vibrational Raman modes, as well as for the band gap of α-ZnP₂, showed very small changes (~0.02 eV) up to 100 kBar [15]. The temperature dependence of the band gap was also reported [15] with a decrease of 0.25 eV in the range from 4.2 to 400 K. Overall, experimental values for the indirect band gap of α-ZnP₂ fall within the range of 2.1–2.3 eV [16, 17]. Theoretical investigations, from pseudopotential methods to more accurate density functional theory calculations, although in fairly good agreement with experiments in the description of structural and elastic properties, failed to accurately reproduce the fundamental electronic band gap (underestimating it by ~0.5 eV) [18–20]. Additionally, the nature of the transition has been wrongly modelled as direct.

In contrast, since it was first synthesized, β-ZnP₂ caused controversy regarding the order of Zn and P atoms. The new crystal form discovered by Hegyi et al [7], monoclinic and black in colour, was suggested to have a fraction of Zn and P atoms bonded to one Zn and three P atoms. However, experimental studies conducted by Fleet [21] suspected the existence of Zn–Zn bonds which they attributed to the insufficiency of high quality intensity data. Further studies have discarded the initial structure and reported each P atom to have two bonds to Zn and two to P atoms, similarly to the α-phase [12, 22]. Changes in the elastic properties that follow the temperature alternations are analogous to those in α-ZnP₂ [13]. From the experimental analysis of the electronic properties, the lowest energy gap of the β-ZnP₂ is identified to be direct [23], with conduction band composed mainly from Zn and the valence band mainly from P ion states [4]. Strong temperature dependence was observed for all optical functions of β-ZnP₂, partially induced by the crystal’s high anisotropy [17]. Experimental band gap values for β-ZnP₂ were reported in the range 1.33–1.60 eV [4, 17]. As in the case of α-ZnP₂, theoretical energies of band gap are smaller than measured values by almost 0.8 eV, but with a reasonable reproduction of structural and mechanical properties [20].

Due to the relative dearth of theoretical works and inconsistency in existing ones, it is clear that a comprehensive study of both α-ZnP₂ and β-ZnP₂ is needed to reveal their possible PV potential. Therefore, density functional theory calculations with standard GGA, Hubbard-corrected GGA+U and screened hybrid DFT methods have been performed to investigate structural and optical properties of α-ZnP₂ and β-ZnP₂. In the case of β-ZnP₂, we adopted two different initial structures within our study and tested them for significant differences. The first structure is based on the data from Hegyi et al [7], hereby named β₁-ZnP₂, while the second follows the atomic arrangement observed by Fleet and Mowles [22], and named β₂-ZnP₂ (for initial coordinates see tables S9–S11 in the supplementary information available online at stacks.iop.org/JPhysCM/31/265501/mmedia). Combining first principles calculated values with an appropriate selection metric, we assessed the suitability of zinc diphosphides as possible photovoltaic absorber materials and obtained maximum values of 10%.

2. Computational details

Electronic structure calculations based on the density functional theory (DFT) [25, 26] were performed with the projector augmented wave (PAW) [27, 28] pseudo-potential method as implemented in the Vienna ab initio simulation package (VASP) [29]. Three different approaches have been considered to describe the exchange-correlation (XC) functional, including the standard Perdew–Burke–Ernzerhof (PBE) general gradient approximation (GGA) [30]; the Hubbard-corrected DFT approach, where an additional potential is added to the GGA total energy to describe the localised (strongly correlated) d-electrons of zinc (GGA+U) [31] method introduced by Dudarev et al [32], with a value of $U_{eff} = 5$ eV; and the screened hybrid XC functional (HSE06),...
where a portion of 25% of the exact non-local Hartree–Fock (HF) exchange is mixed into the PBE total energy [33–35].

The wave function’s kinetic energy cutoff was set to 500 eV (for both zinc diphosphides) together with a Monkhorst–Pack [36] k-mesh of $7 \times 7 \times 3$ ($\alpha$-ZnP$_2$) and $5 \times 5 \times 5$ ($\beta_1$-ZnP$_2$ and $\beta_2$-ZnP$_2$) to sample the first Brillouin zone. For structural optimisations the change in the total energy and interatomic forces between two steps was required to be less than $10^{-6}$ eV and 0.01 eV $^{-1}$, respectively. The optimisation included Van der Waals interactions via the DFT-D3 method developed by Grimme et al [37]. Band structure and density of state (DOS) calculations were performed at the optimized structure along high-symmetry directions obtained from the Bilbao Crystallographic Server, using the crystal symmetry data as referenced in table 1 [38–40]. The band structures were plotted using the Wannier90 code [41].

The LOBSTER package, which permits the electronic structure reconstruction through the projection of PAW-based wave functions onto atomic-like basis sets was employed for the chemical-bonding analysis [42, 43]. To obtain the actual projections, local basis functions as given by Bunge were used [44].

In order to study the mechanical properties of the zinc diphosphides, calculations of the elastic constants have been performed within the strain–stress formalism embedded in VASP [45]. The elastic tensor was determined by performing six finite distortions of the lattice and deriving the elastic constants from the strain–stress relationship. Elastic constants, including both the contribution from distortions with rigid ions and the contribution from ionic relaxations, have been calculated. The phonon spectra were calculated using the small-displacement method via the Phonopy code [46]. The plane wave cutoff was increased systematically until convergence up to 2 GPa has been achieved. While exploring the anisotropy properties, we calculated the bulk and the shear modulus using the Voigt–Reuss–Hill approximation [47],

$$K = (K_V \cdot K_R)^{1/2},$$

$$G = (G_V \cdot G_R)^{1/2},$$

with

$$K_V = \frac{1}{9} \left[ C_{11} + C_{22} + C_{33} + 2 (C_{12} + C_{13} + C_{23}) \right],$$

$$K_R = \left[ S_{11} + S_{22} + S_{33} + 2 (S_{12} + S_{13} + S_{23}) \right]^{-1},$$

$$G_V = \frac{1}{15} \left[ C_{11} + C_{22} + C_{13} - (C_{12} + C_{13} + C_{23}) \right. + 3 \left( C_{44} + C_{55} + C_{66} \right),$$

$$G_R = 15 \left[ 4 (S_{11} + S_{22} + S_{33}) - 4 (S_{12} + S_{13} + S_{23}) + 3 (S_{44} + S_{55} + S_{66}) \right]^{-1}.$$  

Here, $C_{ij}$ are the elastic constants of zinc diphosphides, $S_{ij}$ elements of the compliance matrix, $K_V$ and $K_R$ denote the upper (Voigt) and lower (Reuss) bound of the effective bulk modulus, $K$, of the polycrystalline compound, while $G_V$ and $G_R$ represent those for the effective shear modulus, $G$. In addition, the effective Young’s modulus, $E$, and Poisson’s ratio ($\nu$) can be estimated from the calculated bulk and shear modulus according to the following equations,

$$E = \frac{9KG}{3K+G},$$

$$\nu = \frac{3K-2G}{2(3K+G)}.$$  

To gain insight into the optical properties of the zinc diphosphides, the frequency-dependent dielectric function $\varepsilon(E) = \varepsilon_1(E) + i\varepsilon_2(E)$ at energy $E$ has been computed in the independent-particle (IP) picture. Local field effects have been neglected and the XC functional has been treated in the

<table>
<thead>
<tr>
<th>Table 1. Crystallographic properties of zinc diphosphides.</th>
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<tr>
<td>Crystal structure</td>
</tr>
<tr>
<td>Space group</td>
</tr>
<tr>
<td>$P4_12_12$</td>
</tr>
<tr>
<td>$P4_12_12$</td>
</tr>
<tr>
<td>Class</td>
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<tr>
<td>Pearson symbol</td>
</tr>
</tbody>
</table>

Figure 1. Crystal structures of $\alpha$-ZnP$_2$ (top), $\beta_1$-ZnP$_2$ (middle), and $\beta_2$-ZnP$_2$ (bottom) together with their corresponding bonding atoms.
GGA and HSE06 approximation. From the dielectric function, we have calculated the absorption coefficient \( \alpha \)

\[
\alpha (E) = \frac{4 \pi E}{hc} k (E),
\]

where \( h \) is Planck’s constant, \( c \) the speed of light, and \( k(E) \) the extinction coefficient,

\[
k (E) = \left( \frac{|\epsilon (E)| - \epsilon_1 (E)}{2} \right)^{1/2}.
\]

Also, to assess the solar-absorbing potential of zinc diphosphides, the ‘spectroscopic limited maximum efficiency’ (SLME) has been computed. This concept, proposed by Yu and Zunger [48], offers a calculable selection metric (SLME) has been computed. This concept, proposed by Yu and Zunger [48], offers a calculable selection metric. It expands the traditional Shockley–Queisser (SQ) limit to capture underpinning mechanisms that show relevance to real-time applications.

Spin–orbit coupling (almost negligible experimental value [49]) and excitonic effects have not been included in this work.

3. Results and discussion

3.1. Structural properties

The optimised crystal structures of \( \alpha- \), \( \beta_{1-} \), and \( \beta_2 \)-ZnP\(_2\) are shown in figure 1, whereas the calculated lattice parameters in comparison with experimental and earlier theoretical results are summarised in table 2.

In general, we found good agreement between our calculated unit cell parameters and those reported in literature. A small but not insignificant difference between lattice constants of \( \beta_1 \)-ZnP\(_2\) and \( \beta_2 \)-ZnP\(_2\) was noticed, with \( \beta_2 \)-ZnP\(_2\) being somewhat closer to the experimental data. None of the studied compounds showed magnetic ground state properties. Obtained values of bonds and angles for all three structures are listed in tables S1–S6 (these can be found in the supplementary information file).

Within \( \alpha \)-ZnP\(_2\) (figure 1—top), there are three crystallographically independent atoms (Zn, P\(_1\), and P\(_2\)) with each zinc atom being bonded to four P atoms (two of each, P\(_1\) and P\(_2\)), whereas each phosphorus atom has two bonds with Zn and two with P atoms of the other crystallographic type. The overall arrangement gives four phosphorus chains per unit cell parallel to either \( a \) or \( b \) lattice vector, with possible chain formations in the \( c \) direction being disrupted by zinc atoms. When projected in the \( a \) or \( b \) direction (\( bc \) and \( ac \) planes), chains resemble tetragonally shaped wells. Every zinc interacts with three chains (two running in \( a \) direction and one in \( b \) or vice versa) through four Zn–P bonds building a network of pentagons alternating with existing P chain wells. The P–P bonds of all chains are 2.17 and 2.21 Å, and match distances in pure black phosphorus [50]. The average Zn–P bond length is 2.37 Å, which is shorter than the distance calculated from Pauling’s tetrahedral radii, 2.41 Å [51]. The biggest deviation from the ideal tetrahedral angle (109.5°) comes from a Zn bonded to
two P atoms (124.3°) and is a direct consequence of the 0.05 Å difference in the bond length. Furthermore, bond lengths shorter than covalent ones indicate fractional ionic character and increase in the bond strength and could therefore lead to an opening of the band gap.

Compared to the α-ZnP2 phase, two modifications of the monoclinic β-ZnP2 phase can be distinguished, which are denoted in this study as β1-ZnP2 and β2-ZnP2 based on the presence or absence of zinc–zinc bonds in the structure.

For β1-ZnP2 (figure 1 middle), unlike α-ZnP2, six crystallographically independent atoms can be identified in the unit cell: Zn1, Zn2, P1, P2, P3, and P4. Zn1 is bonded to four P atoms as expected (2 × P1, P2, P3), with P2, P3, and P4 atoms having two bonds with zinc and two bonds with phosphorus atoms. Both Zn2 and P1 are, however, surrounded by one Zn2 and three independent phosphorus atoms (P1, P2, P3). The final structure has only one P chain per unit cell which changes path along all three directions. P–P distances alternate from 2.19 Å to 2.26 Å, and even to 2.32 Å, causing Zn–P bonds and angles to vary over a wider range than in the tetragonal structure. Zn–Zn bonds are in the plane perpendicular to the c axis and tilted by 56.673° from the b axis which agrees well with experimental studies (±57°) [7]. The distance between bonded zinc atoms is very short compared to the covalent radius of the zinc atom, but it is not the first time this has been noticed [53]. According to the structural parameters, it could also be said that the Zn2 atom resembles Zn in α-ZnP2, followed by P1 and P4 atoms behaving like P1 and P3 in the α-modification. Moreover, the presence of the metallic bonds may result in the delocalization of electrons and a decrease of the band gap.

The β2-ZnP2 phase (figure 1 bottom) does not possess zinc atoms bonded to each other, but it still has six crystallographically independent atoms (Zn1, Zn2, P1, P2, P3, and P4). The different types of Zn atoms arise from their positions—half of the atoms sit inside phosphorus chains, while the other half is in between the two chains. Therefore, all zinc atoms are bonding four phosphorus atoms, and all phosphorus atoms have two bonds with zinc and two bonds with P atoms of a different type, similar to α-ZnP2. There are two phosphorus chains per unit cell and their projection is in the shape of an open irregular pentagon. Most angles are closer to the perfect tetrahedral angle compared to β1-ZnP2, leading to better stability. Comparison of the angles of β2-ZnP2 with α-ZnP2 reveals analogues in the two structures: the tetragon surrounding Zn1 closely resembles the tetragon of Zn from α-ZnP2, while P1 and P4 tetragons have their twins in P1 and P2, respectively. Despite these similarities, three additional crystallographic species in the β2-ZnP2 phase disrupt the formation of the same structural patterns as in α-ZnP2. Longer Zn–P bonds indicate less ionic character and lower band gap values.

In both the tetragonal and monoclinic phases, all atoms are tetrahedrally coordinated with Zn–P and P–P bonds within the same tetrahedron differing by ~2%–3%. These differences account for deviations in ideal tetrahedral angles and lead to tetrahedra being somehow distorted when forming a continuous structure. Variations in bond lengths and angle distortions between the two phases provide an insight in the ionic share of bonding and can serve as a guide for the expected electronic structures.

3.2. Elastic properties and lattice dynamics

To obtain information about the mechanical stabilities of the zinc diphosphides, the elastic stiffness constants were calculated. For tetragonal and monoclinic structures, there are six
and thirteen independent elastic constants, respectively. The calculated elastic constants, together with available experimental and theoretical data are given in tables S7 and S8 (supplementary information). As shown in these tables, both zinc diphosphide phases satisfy the mechanical stability criteria by Cowley [54]. Elastic constants obtained from DFT are usually based on single-crystal zinc diphosphides, while the Voigt–Reuss–Hill approximation (VHR) can induce the elastic properties of polycrystalline compounds. Equations for the calculated bulk modulus ($B$), shear modulus ($G$), and Young’s modulus ($E$) in the VHR approximation can be found in the literature [55]. $B$ measures the resistance of material to volume change, $G$ the resistance to shape change, while $E$ often gives a measure of the stiffness of the solid.

Calculated values of $B$, $G$, and $E$ for zinc diphosphides, shown in table 3, vary only slightly from each other, indicating that the three compounds have almost identical resistance to external mechanical forces. The Poisson’s ratio allows us to test the ductility/brittleness of material. A material is characterised as ductile if $G/B < 0.5$, otherwise it is classed as brittle. According to this criterion, $\alpha$-ZnP$_2$, $\beta_1$-ZnP$_2$, and $\beta_2$-ZnP$_2$ are all ductile materials.

Most materials undergo phase transitions driven by lattice instability, both those induced by pressure and those induced by temperature. This instability can be an elastic one (leading to a change of the unit cell shape) or a phonon instability (a so-called soft phonon, whose energy goes to zero) [56]. For a material to be stable, it needs to satisfy a set of conditions [57]. One of the stability requirements of a crystal lattice is its invariability towards any small displacements of atoms. This condition equals to the statement of all phonon frequencies possessing positive values. The identified soft phonon mode appears whilst an external parameter is changed, leading to a crystal instability, resulting with a new crystal structure.

Calculated full phonon dispersion curves along high-symmetry lines in the Brillouin zone are depicted in figure 2. All three studied zinc diphosphide compounds are dynamically stable at atmospheric pressure, since there is no occurrence of imaginary frequencies. The result for $\beta_1$-ZnP$_2$ is somewhat surprising considering the fact that it was evaluated as misinterpreted structural data, and it indicates that $\beta_1$-ZnP$_2$ might be present in real monoclinic crystals as an impurity and future experimental and theoretical studies should not rule it out a priori.

### Table 4. Bader charge analysis of zinc diphosphides.

<table>
<thead>
<tr>
<th></th>
<th>Zn$_1$</th>
<th>Zn$_2$</th>
<th>P$_1$</th>
<th>P$_2$</th>
<th>P$_3$</th>
<th>P$_4$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\alpha$-ZnP$_2$</td>
<td>0.7071</td>
<td></td>
<td>-0.3531</td>
<td>-0.3543</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\beta_1$-ZnP$_2$</td>
<td>0.7139</td>
<td>0.3669</td>
<td>-0.0976</td>
<td>-0.2956</td>
<td>-0.3075</td>
<td>-0.3758</td>
</tr>
<tr>
<td>$\beta_2$-ZnP$_2$</td>
<td>0.7006</td>
<td>0.7014</td>
<td>-0.3716</td>
<td>-0.3318</td>
<td>-0.3484</td>
<td>-0.3502</td>
</tr>
</tbody>
</table>

*Figure 4.* Electron localization function (ELF) for considered three zinc diphosphides calculated using hybrid DFT.
To distinguish further between the stability of the monoclinic $\beta_1$-ZnP$_2$ and $\beta_2$-ZnP$_2$ structures, the equilibrium properties were calculated by fitting the computed total energy versus volume curve with the Birch–Murnaghan equation of state (EOS), which shows the $\beta_2$-ZnP$_2$ structure to be thermodynamically more stable (figure 3).

3.3. Charge analysis

Following the general trend of shorter bonds having a higher share of ionic character, $\beta_1$-ZnP$_2$ with average Zn–P bond length of ~2.35 Å would be expected to have the most pronounced ionic nature. However, this effect could be compensated for long P–P and metallic Zn–Zn bonds, which dominate the whole structure. All Zn–P bonds in $\alpha$-ZnP$_2$ are shorter than pure covalent ones between these species, indicating partial ionic nature. For $\beta_2$-ZnP$_2$ this is true only for 50% of the Zn–P bonds with the remaining bonds either being the length of the perfect covalent bond or longer, indicating less ionic character compared to $\alpha$-ZnP$_2$. To confirm those indications, charge analysis calculations have been performed and the results are shown in table 4. The calculated charge density difference for $\alpha$-ZnP$_2$ shows a shift of bonding electrons towards P atoms as a result of the ionic character, with corresponding Bader charges of $-0.35$ e$^-$ for P and $0.71$ e$^-$ for Zn. The metallic zinc–zinc bonds in $\beta_2$-ZnP$_2$ cause the delocalisation of electrons. Consequently, electrons of P$_1$ atoms tend to migrate in delocalised clouds, leaving one fourth of phosphorus atoms with charge of $0.37$ e$^-$ and bonded zinc atoms with reduced positive charge of $0.37$ e$^+$. Interestingly, in the case of $\beta_2$-ZnP$_2$ only minor differences in charge are obtained compared to the $\alpha$-phase ($0.70$ e$^-$ for Zn and $-0.33$, $-0.34$, $-0.35$, and $-0.37$ e$^-$ for P atoms).

In figure 4, the electron localisation function (ELF) as a direct space depiction of the electron distribution has been plotted for all three zinc diphosphate phases. The ELF value at a considered point indicates higher localisation of electrons in that area compared to a uniform electron gas of the same density. It is evident that the electrons are localised around phosphorus atoms with P atoms in P–P interactions in the $\alpha$-phase compared to the Zn–P interactions in the $\beta$-ZnP$_2$ phases, which can lead...
to increased band gap values. Cyan areas between zinc atoms denote delocalised electrons of the metallic bond and exist only for $\beta_1$-ZnP$_2$.

Since the total charge for single atoms includes electrons from four bonds, not much can be concluded about the nature and strength of each bond. Direct bond analysis can, however, reveal important information and has therefore been conducted through the crystal orbital Hamiltonian population (COHP) technique. COHP data allow comparison between bonds of analogous atoms in each structure and do not rely on any other parameter. Projected COHP (pCOHP) for matching Zn–P, Zn–Zn, and P–P bonds are shown in figure 5. Although bonding peaks for Zn–P analogues appear at different energy ranges, the similarity in the shapes of the pCOHP curves reveals...
the resemblance of the Zn–P bonds in the three zinc diphosphides. The highest peak at \( \sim -8 \text{ eV} \) indicates the strongest bond amongst the three phases and belongs to \( \alpha \)-ZnP\(_2\). In the case of Zn–Zn bonds, \( \alpha \)-ZnP\(_2\) and \( \beta_2 \)-ZnP\(_2\) do not show any bonding states, while the Zn–Zn bond appears to be more than double the strength of Zn–P bonds in \( \beta_1 \)-ZnP\(_2\). All P–P bonds are similar with differences arising mostly from discrepancies in length.

### 3.4. Electronic properties

The electronic band structures, total densities of state (DOS), and projected densities of state (pDOS) for \( \alpha \)-ZnP\(_2\), \( \beta_1 \)-ZnP\(_2\), and \( \beta_2 \)-ZnP\(_2\) were calculated using standard GGA, Hubbard-corrected GGA + U, and screened hybrid DFT approaches and are summarised in table 5.

Overall, our calculated results are consistent with experimental data and earlier DFT predictions. Although the standard GGA functional captures accurately the indirect nature, the obtained band gap value of \( \alpha \)-ZnP\(_2\) is \( \sim 0.70 \text{ eV} \) smaller than the experimentally determined one. Introducing an additional on-site potential (+U\(_{\text{eff}}\)) to GGA increases the band gap for most common semiconductors. With \( \alpha \)-ZnP\(_2\) this is not the case, as a counterintuitive shrinking of the band gap value in comparison with the pure GGA one was observed (for \(-0.05 \text{ eV}\)). A close examination of the calculated densities of states clarified this behaviour. The applied potential localizes the \( d \) orbitals of zinc atoms thereby lowering their energy level by \(-1 \text{ eV}\), as has been depicted in figure S4. Because the top of the valence band consists mostly of phosphorus \( p \) orbitals, inducing (or increasing) the +U\(_{\text{eff}}\) value does not influence the value of the band gap.

Adding a fraction of the short-range screened exchange parameter (HSE06 functional, \( \alpha = 0.25 \)) to the GGA-PBE functional resulted in opening of the band gap, where \( \alpha \)-ZnP\(_2\) now has an indirect band gap of 2.13 eV, which is in a very good agreement with experimental data. The calculated band structure along high-symmetry directions together with the accompanying DOS and pDOS of \( \alpha \)-ZnP\(_2\) obtained by the hybrid functionals are shown in figure 6. The bottom of the conduction band is located at the \( \Gamma \) point, while the top of the valence band can be found along the \( M - \Gamma \) path. Analysis of the orbital pDOS reveals that the majority of carriers at the top of the valence band originate from P-\( p \) orbitals with a small contribution from Zn-s and Zn-\( d \) states, whereas the bottom of the conduction band mainly arises from P-s states.

The band structures and DOS results of \( \beta_1 \)-ZnP\(_2\) and \( \beta_2 \)-ZnP\(_2\) extend the proposed mechanisms obtained from the corresponding charge analysis (shown in figure 6). Results derived from the GGA functional identified \( \beta_1 \)-ZnP\(_2\) as a pure metal, without any bands separated by a gap. Hybrid functionals, on the other side, indicate a small band gap value which was tentatively assigned as an artefact of the calculation itself, rather than an intrinsic property. DOS calculations have shown how the top of the valence band arises from equally contributing Zn-\( p \), Zn-s and P-\( p \) orbitals, which contrasts with \( \alpha \)-ZnP\(_2\) where Zn states play a minor role. The bottom of the conduction band originates from P-\( p \) and P-s states, similar to the tetragonal phase. As proposed, high Zn-\( p \) and Zn-s states originating from the highly metallic character of Zn–Zn bonds prevent the material from showing semiconducting properties, which is shown in figure 7.

This metallic character is in contrast with \( \beta_2 \)-ZnP\(_2\), which already showed a direct band gap from GGA calculations, located at the highly symmetric \( \Gamma \) point, although with a value underestimating experimental ones for \(-0.80 \text{ eV}\). Employing of hybrid functionals gives a direct first transition between valence and conduction band with a corrected value of 1.46 eV, which is in very good agreement with available experimental data. Similar to \( \alpha \)-ZnP\(_2\), the top of the valence has been linked with majority states originating from P-\( p \) orbitals, leaving Zn-\( d \) orbitals with only a minor influence.

Overall, there is a strong overlap between the zinc and the phosphorus components of the DOS structures over the whole range of energies, which confirms the formation of covalent bonds between the two species and a low level of ionicity.

Moreover, it is noted how previous theoretical studies have failed to include all relevant high-symmetry points in the first Brillouin zone. The multivalley nature of the band structure of zinc diphosphides can easily lead to band gap values which do not correspond to the global lowest transitions.

### 3.5. Optical properties and SLME

The optical properties of semiconducting materials are inherently linked to their electronic and vibrational properties. An understanding of how macroscopic properties can be linked to microscopic parameters is important in many applications, especially photovoltaics. At a microscopic level in bulk materials, the complex dielectric function is closely connected with the band structure. The dielectric function generally describes the reaction of a system to an external force. As long as the reaction is linear, the linear response functions (describing the reaction) are properties of the system itself, rather than being force dependent. This concept then can be
applied to the whole electromagnetic spectrum whilst studying complex materials [61].

The calculated real (dispersive, $\varepsilon_1$) and imaginary (absorptive, $\varepsilon_2$) parts of the dielectric function for $\alpha$-ZnP$_2$ and $\beta_2$-ZnP$_2$ are shown in figure 8. The monoclinic phase $\beta_1$-ZnP$_2$ has been omitted from the optical analysis due to its pronounced metallic character, which makes the compound less attractive as a potential PV absorber. The calculated absorption coefficient, an ingredient needed for the PCE analysis, together with the material’s reflectivity has been calculated and shown in figure 9.

The value of the static dielectric function $\varepsilon_1(0)$ is equal to 7.993 for $\alpha$-ZnP$_2$ and 8.017 for $\beta_2$-ZnP$_2$, which corresponds well with previous experimental results yielding values in the range 9–14 for temperatures between 78–100 K [59]. Within the imaginary part of the dielectric function, in principle, one can associate certain peaks with transitions in the band structure.

For $\alpha$-ZnP$_2$, the absorption onset is noted around an energy of 2 eV, which corresponds to the calculated fundamental band transition between the top of the valence and bottom of the conduction band. However, the aforementioned transition is indirect in nature and forbidden by symmetry. This corresponds to a vanishing electronic dipole transition matrix element resulting in a weaker response of the system to incoming electromagnetic radiation. The first significant peak that can be observed lies around 3 eV and arises from transitions deeper within the band structure. Most of the transitions up to that point are still forbidden by symmetry and contribute only weakly to the response functions. On the other hand, $\beta_2$-ZnP$_2$ has a direct allowed transition between the bands separated by an electronic gap. This transition takes place around 1.5 eV, as one would expect from the calculated band gap value. By looking at the probabilities of absorbing photons of a certain wavelength, which is depicted in figure 9, one can clearly observe the earlier absorption onset for $\beta_2$-ZnP$_2$ compared to $\alpha$-ZnP$_2$.

After calculation of the band structures and dielectric function of the zinc diphosphides, we have obtained all the required information to assess their photovoltaic potential. The calculated SLME as a function of the film thickness for zinc diphosphides was calculated and shown in figure 10. For a reasonable reference thickness of 0.5 µm we obtained 0.28% and 9.39% efficiencies for $\alpha$-ZnP$_2$ and $\beta_2$-ZnP$_2$, respectively. The origin of this somewhat surprising difference comes from the distinct band nature of the two compounds. For $\alpha$-ZnP$_2$, there is a non-zero difference between the direct allowed and fundamental band transition ($\Delta = E_{gA} - E_{g} > 0$) which indicates that non-radiative processes are significant within this compound.
Although the SLME metric was found to overestimate the reverse saturation current, in our case it is of the order of $10^{-3}$, which seems to be a reasonable value compared to other similar classes of materials [62]. If we would consider radiative recombination as the only process taking place, we would resemble the classic SQ limit and obtain an efficiency value of 5.86\% for $\alpha$-ZnP$_2$ (considering a 0.5 $\mu$m thick absorber). However, the fundamental transition for $\beta_2$-ZnP$_2$ is already dipole allowed and we can expect radiative recombination as the main recombination process governing absorption.

4. Conclusion

A comprehensive computational study of the structures and properties zinc diphosphides has been undertaken. Calculations employing the HSE06 XC functional have been proven accurate in describing structural, mechanical, electronic, and optical properties of all three targeted zinc diphosphide compounds. Unlike the stable tetragonal phase, the two monoclinic structures show different stabilities and bonding character. Specific Zn–Zn bonds within $\beta_1$-ZnP$_2$ cause metallic behaviour and a lower stability than $\beta_2$-ZnP$_2$. Moreover, electronic structure calculations that reproduce the experimentally observed features have been conducted. They show diverse band transitions and electric dipole moments which lead to substantially different optical properties. In addition, we assessed the thin-film photovoltaic potential of zinc diphosphides, showing $\beta_2$-ZnP$_2$ as a promising absorbing material. Future investigations will expand the work presented here to include the effect of doping, or the creation of heterojunctions on tuning the electronic and optical properties of zinc diphosphides to achieve efficient charge carrier separation and improved photovoltaic efficiencies.

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References

[12] Zinin I E, Aleinikova K B and Antipin M Y 2003 Analysis of chemical bonding in the $\alpha$ and $\beta$ modifications of zinc diphosphide from x-ray diffraction data Crystallogr. Rep. 48 190–204

Figure 10. Calculated spectroscopic limited maximum efficiency (SLME) for $\alpha$-ZnP$_2$ and $\beta_2$-ZnP$_2$ plotted as a function of the film thickness.

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[53] Vold C L 1961 The crystal structure of NbZn$_2$, Acta Crystallogr. 14 1289–90


[57] Lukačević I 2011 High pressure lattice dynamics, dielectric and thermodynamic properties of SrO Physica B 406 3410–6

[58] Stavros I, Syrbu N N and Zalama V V 2014 Optical properties and band structure of ZnP$_2$–D$_x$$_4$ J. Lumin. 149 19–27

