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Towards an Accurate Estimate of the Exfoliation Energy of Black Phosphorus: A Periodic Quantum Chemical Approach

Giuseppe Sansone1,∗ Lorenzo Maschio1, Denis Usvyat2,∗ Martin Schütz2, and Antti Karttunen3∗

1Dipartimento di Chimica and NIS (Nanostructured Interfaces and Surfaces) Centre, Università di Torino, via Giuria 5, I-10125 Torino, Italy

2 Institute for Physical and Theoretical Chemistry, University of Regensburg, Universitätsstraße 31, D-93040 Regensburg (Germany)

3Department of Chemistry, Aalto University, Kemistintie 1, 02150 Espoo, Finland

E-mail: giuseppe.sansone@unito.it; denis.usvyat@chemie.uni-regensburg.de; antti.j.karttunen@iki.fi

Abstract

The Black phosphorous (black-P) crystal is formed of covalently bound layers of phosphorene stacked together by weak van der Waals interactions. An experimental measurement of the exfoliation energy of black-P is not available presently, making theoretical studies the most important source of information for the optimization of phosphorene production. Here, we provide an accurate estimate of the exfoliation energy of black-P on the basis of multilevel quantum chemical calculations, which include the periodic local Möller-Plesset perturbation theory of second order, augmented by higher-order corrections, which are evaluated with finite clusters mimicking the crystal.
Very similar results are also obtained by density functional theory with the D3-version of Grimme’s empirical dispersion correction.

Our estimate of the exfoliation energy for black-P of -151 meV/atom is substantially larger than that of graphite, suggesting the need for different strategies to generate isolated layers for these two systems.

Phosphorus is well known for its ability to form a variety of allotropes in the solid state. In this respect it closely resembles carbon, whose nanostructures (0D-fullerenes, 1D-nanotubes, 2D-graphene) have led to the impressive development of nanoscience in recent years. Solid phases of phosphorus, although known since long, are studied nowadays with a renewed interest,¹ as precursors of nanostructures analogous to carbon-based ones. Among the graphene²³-inspired 2D materials, such as silicene,⁴ germanene⁴ and stanene,⁵ phosphorene has been very recently fabricated⁶⁷ and proposed as a candidate for future electronics, optoelectronics, photovoltaics and spintronics applications.⁸⁹ A systematic development of new modifications and applications of phosphorene requires a complete understanding of structure and energetics of its parent crystal, i.e., black phosphorus (black-P). The latter is the most stable allotrope of phosphorus,¹⁰¹¹ and is built from an infinite stacking of phosphorene layers, cf. Figure 1.

Notably, a detailed understanding of the interactions in black-P can show new ways to fabricate high-quality phosphorene samples and novel phosphorene-based technologies. From a computational point of view, black-P is a very challenging system because of the simultaneous presence of covalent (intralayer) and weak dispersive (interlayer) forces. It is well-known that Kohn-Sham density functional theory (DFT)¹²¹³ utilizing commonly used approximations of the exchange correlation functional is unable to properly describe van der Waals interactions.¹⁴ Nonempirical van der Waals functionals,¹⁵¹⁷ dispersion-corrected atom-centered potentials (DCACPs)¹⁸, and Grimme’s empirical dispersion corrections¹⁹²¹ offer a practical solution to that problem. Yet, they typically rely on some form of parametrization and may therefore fail unexpectedly. Computational studies of solid phosphorus using such DFT
Figure 1: The black phosphorus crystal structure is built from phosphorene layers. The “chair” interconnection of the six-membered rings gives rise to the characteristic puckered shape of the layers. Lowercase letters (a,b,c) refer to the lattice parameters. Along the direction of the b lattice vector, the interlayer distance (d_{int}) is depicted. Uppercase letters (A,B,C,D) have been used to define the so-called hinge angle (\angle CBD) and the adjacent P–P bond length (d_{BC}), the dihedral angle (\angle ABC), and the connecting bond length (d_{AB}).

approaches and Quantum Monte Carlo methods are found in the recent literature.\(^8,^{22–26}\) Nevertheless, when looking for a reliable estimate of a physical quantity, accurate \textit{ab initio} wavefunction methods are indispensable tools since they can be employed to assess and rank the quality of different DFT approaches for a given system. Post-HF approaches, as employed in this work, have already provided important insight into the formation of other allotropes of phosphorus, such as that of red of phosphorus, where systematic MP2 studies on phosphorus clusters have been performed already over 20 years ago.\(^27\)

Here we apply periodic local second-order Møller-Plesset perturbation theory (p-LMP2)\(^{28–30}\) to study the interlayer interactions between the phosphorene layers in black-P, which is corrected for basis set and method errors by using finite cluster calculations. These results are compared to those obtained from different flavours of dispersion-corrected DFT. Post-Hartree-Fock (HF) calculations on crystalline solids within the periodic theoretical framework are nowadays still not yet routine.\(^{31–33}\) For the present study the recent developments in
the p-LMP2 method,\textsuperscript{34} such as the implementation of orbital-specific virtuals (OSV),\textsuperscript{35} were very instrumental, since they substantially simplify calculation of smooth potential energy surfaces (\textit{vide infra}).

Following a computational protocol, recently proposed for molecular crystals,\textsuperscript{36} and adsorption on surfaces\textsuperscript{37,38} we have evaluated finite cluster corrections accounting for the method and basis-set errors as well as the neglect of core-electron correlations. The corrections are computed by using the interaction energies of the P$_4$H$_6$ – P$_{16}$H$_{10}$ dimer (see supplementary information), cut out from the crystal structure within the given interlayer distance, with the dangling bonds substituted by P-H ones. The correcting energy increments are evaluated as the differences in interaction energies on top of the OSV-LMP2 (or OSV-spin-component-scaled-LMP2, see below) results. The high-end quantum chemical treatment reaches the coupled cluster with singles, doubles and perturbative triples CCSD(T) level (often considered the “gold standard” of quantum chemistry) in a reasonably large basis set, \textit{i.e.}, of augmented triple zeta quality. Due to the relatively large size of the dimer, local\textsuperscript{39,40}, rather than canonical CCSD(T) calculations are performed. To this end we utilize the LCCSD(T0)|LCCD[S]-R$^{-6}$ method.\textsuperscript{41,42} This local approximation to the CCSD(T) method provides a very high accuracy for intermolecular systems.\textsuperscript{42}

MP2 is notorious for overestimating dispersion for highly polarizable systems. As will be clear from the results presented below, this is also the case for black-P (which is a small band gap semiconductor), if an extended and diffuse basis set is used. For such systems, the spin-component-scaled(SCS)-MP2\textsuperscript{43} technique is known to be much more stable and closer to CCSD(T). Therefore we evaluate the basis set and CCSD(T) corrections with respect to the periodic OSV-SCS-LMP2 treatment. For brevity, we denote in the following the sum of all the three corrections, \textit{i.e.}, method, basis set, and core correlation, by the one acronym $\Delta$CCSD(T).

All calculations have been performed by using the CRYSTAL\textsuperscript{44} program for HF and DFT, and CRYSCOR\textsuperscript{30} for p-LMP2, and MOLPRO\textsuperscript{45} for the LCCSD(T) calculations and other
cluster corrections. All these codes employ a basis set of atom-centered Gaussian-type orbitals. In periodic calculations we used basis sets derived from the molecular Karlsruhe basis sets \(^{46}\) (see Supporting Information – SI – for details). We applied basis sets of different quality: split-valence plus polarization (SVP) and triple-zeta-valence with three different sets of polarization functions (TZVP with 1d; TZVPP with (2d1f)|2d1f| and TZVPP2 with (3d1f)|2d1f|). Finally, in the periodic LMP2 and SCS-LMP2 calculations for determining the exfoliation energy we augmented the TZVPP2 basis with diffuse d- and f-orbitals from Dunning’s aug-cc-PVTZ basis set, \(^{47}\), which are essential for an accurate description of dispersion.\(^{38,48}\) Since such diffuse orbitals are harmful for periodic HF, they were added only \textit{a posteriori} for the correlated calculations by means of the dual basis set technique.\(^{29}\)

The finite cluster corrections were computed using basis sets from the Dunning family, \textit{i.e.}, aug-cc-pVTZ for the CCSD(T) - SCS-LMP2 energy difference; (ii) aug-cc-pVTZ and aug-cc-pVQZ for the basis set extrapolation at the SCS-MP2 level; (iii) aug-cc-pwCVTZ\(^{49}\) for the correlated core correction.

As a first step, we optimised the cell parameters of black-P at the p-LMP2 level. Since p-LMP2 analytical nuclear gradients are presently not available in the CRYSCOR code, we sampled the potential energy surface through single point calculations on a 3D grid around the experimental structure. Five symmetry irreducible geometrical parameters need to be optimised for black-P: three lattice parameters (\(a, b, c\)) and two fractional coordinates of the atoms \(y/b\) and \(z/c\). We tackled these cell and atomic coordinates separately. 2D potential energy surfaces were then obtained from fitting a quadratic function at each \(b\) value to the corresponding subset of data points. Figure 2 shows an example of such a fitting. For more details on the fit see SI. The resulting potential energy surface is indeed smooth. Such smoothness is not automatically achieved in the local correlation approach, based on projected atomic orbitals,\(^{28}\) due to a possible domain mismatch along the potential surface scan. However, the recent implementation of the OSV-based p-LMP2\(^{34}\) to a large extent eliminates this problem.
Figure 2: Potential energy surface of black-P fitted to explicitly computed energies at the p-LMP2/TZVPP level of theory. Left Panel: Energy as a function of $a$ and $c$ cell parameters (with fixed $b = 10.38 \text{Å}$). Right Panel: Energy as a function of the internal coordinates of the phosphorus atoms in the fractional units $y/b$ and $z/c$, with fixed $x/a = 0$.

Note that the lattice parameter $b$ corresponds to the stacking direction, and both interlayer distance and layer thickness are related to it. Note that the change in the latter, which also involves contraction or elongation of the P-P bonds, corresponds to a “hard” coordinate (with large curvature of the potential energy surface along this direction). On the other hand, the interlayer distance is a typical “weak” coordinate. Therefore, the lattice parameter $b$ strongly couples these hard and weak coordinates. The so “optimized” $b$-parameter value hence is governed by the hard coordinate.

In order to decouple the inter- from the intra-layer interaction, we computed a one-dimensional potential energy profile in which all coordinates were frozen at the previously found minimum, and only the interlayer distance $d_{int}$ was varied. Furthermore, here we focused directly on the exfoliation energy, which allowed us not only to eliminate the basis set superposition error for this delicate quantity, but also to incorporate the finite cluster $\Delta \text{CCSD(T)}$ corrections into the potential energy curve. The counterpoise corrected exfoliation energy is given by the expression:

$$E_{\text{exf}} = E_{\text{bulk}} - E_{\text{lay}}^{\text{gh}} + \Delta E_{\text{relax}},$$  

(1)
where $E_{\text{bulk}}$ and $E_{\text{lay}}^{\text{gh}}$ are the total energy of the bulk black-P and the total energy of a single layer of black-P in the bulk geometry with ghost functions on the two nearest-neighboring layers, respectively (both in energies are per atom). The relaxation energy for the single layer

\[ \Delta E_{\text{relax}} = E_{\text{lay}} - E_{\text{lay}}^{\text{opt}}, \tag{2} \]

involves the total energies (also per atom) of the isolated layer in the bulk ($E_{\text{lay}}$) and the optimized ($E_{\text{lay}}^{\text{opt}}$) geometries. In the absence of periodic LMP2 nuclear gradients, the optimization of the layer, as well as the evaluation of $\Delta E_{\text{relax}}$ was done at the PBE0-D3 level.

First, in Table 1 we report the optimized (according to the described scheme) p-LMP2 lattice parameters for the TZVPP and TZVPP2 basis sets and compare them to the fully optimized (cell parameters and fractional coordinates) values from two DFT functionals PBE\textsuperscript{52} and PBE0\textsuperscript{53} with and without dispersions corrections D2\textsuperscript{19} and D3\textsuperscript{21}. From these results several observations can be made:

(i) Van der Waals forces are crucial for a correct description of black-P. A wide range of values for the $b$ parameter is obtained by the different functionals. The PBE and PBE0 functionals with no dispersion corrections tend to substantially overestimate the cell volume.

(ii) The better performance of the plain PBE or PBE0 functionals in conjunction with smaller basis sets (compared to saturated basis sets) is due to basis set superposition error (BSSE) effects that act as spurious attractive interaction between the layers.

(iii) The agreement with experiment of the dispersion corrected DFT and LMP2 is very good as far as $a$ and $c$ are concerned. For the $b$ parameter only DFT-D2 gives a result close to the experimental value. We note, that in previous works it was shown that the D3 correction can generally be expected to be superior to D2,\textsuperscript{54} even though weak interactions are overestimated.\textsuperscript{55,56} Here we see that both DFT-D3 and LMP2 similarly provide a noticeably smaller value for $b$ than DFT-D2 as well as the experiment. At the same time, experimental values corresponds to room temperature and therefore it remains unclear if it can be regarded as a proper reference for the electronic-energy-only optimization of the $b$ parameter.
Table 1: Summary of the black-P structural data obtained from the p-LMP2 and DFT structure optimisations. The relaxed cell parameters \((a, b, c)\) and inter-layer distance \((d_{\text{int}})\) have been compared with two experimental studies.

<table>
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<th>Method</th>
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<th>(a) (Å)</th>
<th>(b) (Å)</th>
<th>(c) (Å)</th>
<th>(V) (Å(^3/\text{atom}))</th>
<th>(d_{\text{int}}) (Å)</th>
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All computed data are for 0 K, thermal expansion has not been considered.
Experimental data obtained at room temperature.
In order to investigate this discrepancy in more detail we focus on the exfoliation energy as a function of $d_{\text{int}}$. $\Delta E_{\text{exf}}$, computed at different levels, is presented in Fig. 3. Importantly, here we have access to a very accurate reference, obtained by adding $\Delta \text{CCSD}(T)$ to the periodic SCS-LMP2/aug(d,f)-TZVPP2 result.

![Figure 3](https://example.com/figure3.png)

**Figure 3**: Plot of the exfoliation energy as a function of the interlayer distance in bulk black-P, as evaluated with different methods. Curves have been fitted with a 1D Morse potential. The curves are qualitatively comparable to graphite studied using Quantum Monte Carlo (Spanu *et al.*, 57), but the absolute value of the exfoliation energy is clearly larger for black-P.

First we see from Fig. 3 that the electronic-energy minimum in the reference curve is rather close to that of DFT-D3 and LMP2 with TZVPP2 basis, hence is shifted to smaller $d_{\text{int}}$ relative to the experimental and DFT-D2 values. At the same time we note an extreme shallowness of the potential curve for a wide range of $d_{\text{int}}$ values and thus of the lattice parameter $b$. This indicates that the thermal expansion of the lattice parameter $b$ can be appreciable in black-P. The experimental linear thermal expansion coefficient in the $b$-direction is $39 \times 10^{-6}$ 1/K at room temperature (10). It is about 1.5 times larger than that of graphite ($25 \times 10^{-6}$ 1/K at room temperature) (58) and implies a small but not negligible change in the $b$-parameter with temperature. The thermal expansion thus can be a reason
for the discrepancy between the best theoretical and the experimental results. Considering this, the DFT-D2 minimum is likely to be incorrect, despite the good agreement with the experimental structural parameters obtained at the room temperature. In this respect, a theoretical modeling of the thermal expansion of black-P appears is very interesting and will be explored in future work.

As noted above the LMP2 with extended and diffuse basis substantially overestimates binding. Smaller basis sets, especially TZVPP, provide a reasonably good description due to a cancellation of method error vs. basis set error. The SCS-LMP2 method, on the contrary, provides a quite accurate interaction energy for black-P only with a sufficiently large basis set.

The exfoliation energy of a layered material such as black-P is equal to the interlayer binding energy (also known as interlayer cohesive energy) and an understanding of the energetics of the interlayer bonding in the layered compounds is fundamental for optimizing their exfoliation processes. Notably, for black-P, a controllable exfoliation process could be exploited to produce few-layer phosphorene systems with a tunable band gap. In other words, few-layer phosphorene is a semiconductor with a finite gap, which is strongly affected by the number of layers, making it highly attractive for applications in, e.g., optoelectronics. However, apart from the archetypal graphite/graphene case, experimental data on exfoliation energies are very rarely available for weakly bonded layered inorganic compounds. In particular, no experimental estimate on the exfoliation energy of black-P has been reported, to our knowledge. For this reason, an accurate estimate of such quantity is of fundamental importance for a better understanding of the physico-chemical properties of the black-P/phosphorene system at the nanoscale.

The results for the exfoliation energy for Black-P are compiled in the Table 2. PBE-D2 and PBE0-D2 predict it to be around -90 to -100 meV per atom. Similar values for the DFT-D2 exfoliation energy have recently been obtained in Refs. However, both
Table 2: The exfoliation energy ($E_{\text{exf}}$) of black-P calculated using different approaches and corrections. $E_{\text{exf}}$ has been calculated from Eq. (1).

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<tr>
<th>Method</th>
<th>Basis set</th>
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<th>$d_{\text{int}}$ (Å)</th>
<th>$E_{\text{exf}}$ (meV/atom)</th>
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<td>TZVPP</td>
<td>10.19</td>
<td>2.96</td>
<td>-155</td>
</tr>
<tr>
<td></td>
<td>TZVPP2</td>
<td>10.17</td>
<td>2.97</td>
<td>-148</td>
</tr>
</tbody>
</table>

\textsuperscript{a} Includes finite-cluster corrections for the method, basis set errors and the core-valence correlation.

p-SCS-LMP2+CCSD(T) and DFT-D3 provide a much stronger interaction in the range of about -140 to -160 meV/atom. Since this estimate is predicted by a sophisticated high-level quantum chemical treatment and is furthermore supported by DFT with the most recent parameterization of the dispersion correction, we consider it as the most accurate benchmark so far for the electronic contribution to the exfoliation energy of black-P.

For graphite, the exfoliation energy has been estimated to be in the range of -30 to -50 meV per atom.\textsuperscript{61–63} Hence, the predicted exfoliation energy of black-P is considerably larger than that of graphite. Such a substantial difference in the inter-layer energetics between black-P and graphite suggests that the optimal exfoliation strategies for these two systems might be quite different.

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Supporting Information Available

This material is available free of charge via the Internet at http://pubs.acs.org/.

References


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Graphical TOC Entry

Black phosphorus, phosphorene, periodic quantum chemical methods, van der Waals interactions in solids