¹ Phosphorene–Fullerene nanostructures:

A first-principles study

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9Abstract. Hybrid materials formed by carbon fullerenes and layered materials have emerged due 10to their advantages for several technological applications, and phosphorene arises as a promising 11two-dimensional semiconductor for C₆₀ adsorption. However, the properties of phosphorene-12fullerene hybrids remain mainly unexplored. In this work, we employed density functional 13theory to obtain structures, adsorption energies, electronic/optical properties, binding (AIM, 14NBO), and energy decomposition analyses (ALMO-EDA) of nanostructures formed by 15phosphorene and fullerenes (C₂₄ to C₇₀). We find fullerenes form covalent and non-covalent 17even in solution. Two classes of covalent complexes arise by cycloaddition-like reactions: the 18 first class, where short-range effects (charge-transfer and polarization) determines the stability; 19and the second one, where short-range effects decay to avoid steric repulsion, and balanced long-20range forces (electrostatics and dispersion) favors the stability. Otherwise, high-size fullerenes 21(C₅₀ to C₇₀) only form non-covalent complexes due to strong repulsion at shorter intermolecular 22distances and lack of dissociation barriers. In terms of electronic properties, fullerenes act as 23mild *p*-dopants for phosphorene, increasing its polar character and ability to acquire induced 24dipole moments (polarizability). Also, small energy-bandgap fullerenes (<0.8 eV) largely 25 increase the phosphorene metallic character. We also note fullerenes retain their donor/acceptor 26 properties upon adsorption, acting as active sites for orbital-controlled interactions and 27 maximizing the phosphorene light absorbance at the UV-Vis region. Finally, we strongly believe 28our study will inspire future experimental/theoretical studies focused on phosphorene-fullerene anode materials, sensing, phosphorene bandgap engineering, 29uses for storage, and 30optoelectronics.

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31**Keywords:** Phosphorene; DFT calculations; nanotechnology; composites; nanostructures; 32surface science.

331. Introduction

34 Among low-dimensional carbon allotropes (such as graphene and carbon nanotubes), the 35buckminsterfullerene molecule (C_{60}) is a class of 0-dimensional (0-D) organic compounds of 36spherical molecular shape and high surface area, displaying remarkable physical and chemical 37properties due to surface/interfacial effects and quantum confinement¹. In particular, C₆₀ displays 38a remarkable electron acceptor character with a semiconductor bandgap. Also, C₆₀ is merely 39composed of sp^2 -hybridized carbon atoms, which confer it an electron-deficient polyalkene 40nature, and thus, it is chemically reactive¹. Likewise, small fullerenes have been synthesized and 41characterized (e.g., from high-resolution transmission electron microscopy), which have 42associated an increased reactivity with attaching to substrates because of containing paired or 43adjacent pentagons²⁻³, and their electronic properties are significantly influenced by the shape 44and size ⁴. The unique properties of fullerenes turn into useful building blocks for constructing 45supramolecular assemblies and micro/nanofunctional materials¹. Consequently, non-covalent and 46covalent nanostructures have been fulfilled by depositing fullerene onto emerging low 47dimensional substrates such as graphene, perovskites, graphite-like carbon nitride, transition-48metal disulfides, and hexagonal boron nitride^{1, 5}. These hybrid nanostructures show potential 49applications in different technologies such as catalysis, nanoelectronics, optoelectronics, storage, 50batteries, solar cells, and spintronics, among others^{1, 5}.

Phosphorene emerged as a new generation of nanomaterials with an anisotropic 2-52dimensional (2-D) structure in a puckered honeycomb shape with sp^3 -orbital hybridization⁶. 53Relevant properties of phosphorene are high electrical conductivity (2·10⁵ cm² V⁻¹ s⁻¹), high

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54carrier mobility ($\approx 1000 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$), high thermal conductivity (4840–5300 W m⁻¹ K⁻¹), and high 55tensile strength up to 130 GPa⁶⁻⁸. Phosphorene also displays a remarkable ability to bind organic/ 56inorganic molecules through covalent or non-covalent interactions, mainly favored by 57electrostatic attraction⁹⁻¹²; thus, phosphorene emerges as an ideal template for the formation of 58well-ordered structures for assembly of fullerenes and related phosphorene/organic 59nanostructures. Furthermore, surface coordination of phosphorene has been shown to enhance its 60stability under air/water dispersion, which is a major challenge in practical applications¹³⁻¹⁵. 61Despite widespread works on hybrids of fullerenes with other 2D nanomaterials, studies on 62Phosphorene-Fullerene hybrids have infrequently been reported and mainly focused on C₆₀ 63adsorption. Experimentalists and theoreticians have synthesized/proposed Phosphorene-C₆₀ 64nanostructures with relevant properties for technological applications such as high specific 65capacity battery electrode materials¹⁶, solar energy conversion¹⁷⁻¹⁸, new molecular doped 66crystalline superlattices for semiconductor industry¹⁹, and heterojunction photodetectors²⁰. In this 67way, properties of phosphorene and fullerenes are compensated synergistically; for example, it is 68found in photovoltaic research that fullerenes act as excellent electron acceptors in polymer solar 69cells, while phosphorene improves the energy alignment in the devices, which lead to improved 70power conversion efficiency by favoring the charge transfer and exciton dissociation¹⁸.

Additionally, the packing of fullerenes onto substrates must be influenced by the shape 72and size, which would play a key role in the properties and stability of Phosphorene-Fullerene 73hybrids; then the physical/chemical phenomena that dominate the interaction strength is essential 74to be clarified. Mechanochemical reactions in a high energy mechanical milling process have 75been used as a strategy to form phosphorus-carbon (P–C) bonds between phosphorene and 76carbon materials, including C_{60} , graphite, and graphite oxide¹⁶; in this way, the content of the

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77P–C bond in the phosphorene-C₆₀ hybrids is only 0.8%, denoting C₆₀ is not preferably bonded to 78phosphorene *via* covalent interactions until breaking the *sp*² C=C/C–C bonds to form defects¹⁶. 79In addition, low-temperature scanning tunneling microscopy, X-ray, ultraviolet photoelectron, 80and scanning tunneling spectroscopy measurements also show that C₆₀ molecules are mainly 81physisorbed at room temperature on the honeycomb lattice of blue and black phosphorene 82synthesized by epitaxial growth, where an interfacial charge transfer is evidenced upon 83interaction with C₆₀²¹⁻²². The Phosphorene-C₆₀ non-covalent interaction has also been confirmed 84by the disappearance of the C₆₀ absorption band at 340 nm in phosphorene Langmuir–Blodgett 85films after a simple toluene wash²⁰. In this regard, density functional theory (DFT) computations 86supported a simple physical interaction between C₆₀ and phosphorene characterized by strong 87electron density rearrangements (charge transfer) and adsorption energies of ~1 eV²²⁻²³.

The background mentioned above indicates phosphorene could be implemented as a 89remarkable substrate for the assembly of novel hybrid nanostructures with carbon fullerenes. 90Nevertheless, it is still not well understood yet the influence of fullerenes size on Phosphorene-91Fullerene hybrids' properties, such as the stability and preferred binding mechanism 92(covalent/non-covalent), electronic/optical properties after bonding, and the contribution of 93intermolecular forces determining the adsorption stability. To answer these questions, we 94employed a dispersion-corrected DFT scheme to obtain adsorption energies and conformations, 95electronic/optical properties, electron density-based analyses, and energy decomposition analyses 96of Phosphorene-Fullerene nanostructures, which provide deep insights into the stability, 97structure, adsorption mechanism, and useful potential properties for technological applications. 98We consider either the covalent or non-covalent binding of fullerenes in different sizes and 99symmetries as representative classes, i.e., C_{24} (D_{6h}), C_{26} (D_{2h}), C_{40} (D_{2}), C_{40} (D_{2}), C_{40}

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100(D_2), C_{50} (D_3), C_{60} (l_h), and C_{70} (D_{5h}); in this way, it is provided a wide family of adsorbates 101resulting in different assemblies for characterization.

1022. Computational Details

103 We used the PBE functional with the all-electron def2-SVP basis sets for all the DFT 104calculations in the ORCA4.1²⁴⁻²⁶. The PBE functional has been used to describe the interactions 105of phosphorene with a wide range of adsorbates. The DFT-D3(BJ) procedure included dispersion 106corrections into the PBE functional for energies and gradients²⁷. Molecular structures were 107optimized without geometrical constraints and verified through frequency analyses, where 108positive vibrations were associated with all vibrational modes. Convergence tolerance values of $1091 \cdot 10^{-8}$ and $1 \cdot 10^{-6}$ Ha were used for SCF and geometry optimizations; the geometries converged 110 with tolerance values in gradients and coordinate displacements of $3 \cdot 10^{-4}$ Ha/Bohr and $4 \cdot 10^{-3}$ 111Bohr, respectively. Continuum solvent effects were included by the universal continuum 112solvation model (SMD) based on the quantum mechanical charge density of a solute molecule 113interacting with a continuum description of the solvent²⁸. Excited states were obtained with the 114simplified time-dependent density functional theory (sTD-DFT) approach combined with the 115meta-hybrid TPSSh functional (10% Hartree-Fock exchange)²⁹⁻³⁰; configuration state functions 116were included up to an energy threshold of 6.2 eV. Phosphorene nanoflakes (P₁₂₆H₃₀) were used 117 for adsorption studies and with a surface area of at least \sim 1585 Å² considering its electron 118density, which is relatively larger to obtain well-converged adsorption energies concerning the 119 fullerenes surface area (up to ~456 Å²). Adsorption energies (E_{ads}) were computed as:

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$$E_{ads} = E_{Phos} + E_{Fullerene} - E_{Phos-Fullerene} + \Delta ZPE$$
(1)

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121where E_{Phos} , $E_{Fullerene}$, and $E_{Phos-Fullerene}$ are the total energies of the free phosphorene, free fullerenes, 122and the complex, respectively; ΔZPE stands for the zero-point energy correction. Thus, the more 123positive the E_{ads} values, the more stable the complex is. The counterpoise correction was used to 124avoid BSSE in the E_{ads} values³¹. Adsorption energies were further decomposed by the second-125generation energy decomposition analysis based on absolutely localized molecular orbitals 126(ALMO-EDA) of the Q-Chem5.2 program at the PBE-D3/def2-TZVP level³². Hence, the 127adsorption energy for one AB complex expresses as³³⁻³⁴:

$$-E_{ads} = \Delta E_{CT} + \Delta E_{POL} + \Delta E_{ELEC} + \Delta E_{DISP} + \Delta E_{PAULI} + \Delta E_{PREP}$$
(2)

129where ΔE_{CT} , ΔE_{POL} , ΔE_{ELEC} , and ΔE_{DISP} stand for the energy lowering due to charge-transfer (inter 130and intramolecular charge flow between fragments), polarization (induced electrostatics), 131Coulombic attractions (classical intermolecular electrostatics), and dispersion forces (van der 132Waals interactions), respectively. ΔE_{DISP} is obtained with the dispersion-free revPBE functional. 133 ΔE_{PAULI} is the energy destabilization due to Pauli repulsion when two fragments are close 134enough, i.e., electrons cannot have the same position (Pauli principle). ΔE_{PREP} is the preparation 135energy penalty due to the geometric distortion of fragments to reach the complex geometry.

Intermolecular interactions were also revealed by the Atoms-in-Molecules (AIM) 137method³⁵, where the electron density (ρ_i) of the bond critical points (BCPs) connecting fragments 138through intermolecular bond paths serve as a measure of the interaction strength. In this way, 139covalent bonds, closed-shell interactions, and weak electrostatic interactions are characterized by 140 ρ_i values of $\rho_i \ge 0.10$, $\rho_i \approx 0.10$ -0.04, and $\rho_i \le 0.01 \ e/Bohr^3$, respectively ³⁵. Furthermore, the 141signature of weak interactions were examined by the Independent Gradient Model (IGM)³⁶, 142which introduce the δg^{inter} descriptor that uniquely defines intermolecular interaction regions:

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143 $\delta g^{\text{inter}} = |\nabla \rho^{\text{IGM,inter}}| - |\nabla \rho|$, where $\nabla \rho$ stands for the electron density gradient and $\nabla \rho^{\text{IGM,inter}}$ is an 144upper limit to $\nabla \rho$ as defined by Lefebvre and co-workers³⁶. AIM, IGM (based on the Hirshfeld 145partition), CM5 charges, and wavefunction analyses were performed in Multiwfn3.7³⁷.

1463. Results and Discussions

1473.1. Structure and stability.

We place fullerenes initially ~5 Å away from the phosphorene surface; at least ten 1490rientations for each molecule were considered according to its symmetry. The **Phos-Fullerene** 150systems form two groups of stable complexes depending on the fullerene size (Fig. 1a): i) 151covalent and ii) non-covalent complexes. C_{24} to C_{44} fullerenes form covalent complexes with 152phosphorene *via* chemisorption, showing intermolecular distances of $d_{inter} \approx 2.0-2.2$ Å; while C_{50} 153to C_{70} fullerenes form non-covalent complexes *via* physisorption, showing intermolecular 154distances of $d_{inter} \approx 2.6-2.8$ Å (Fig. 1b).



156**Fig. 1.** Properties of **Phos-Fullerene** complexes: a) Ground state structures. b) Average 157intermolecular distances (d_{inter} , in Å). c) Adsorption energies of ground states (E_{ads} , in eV). d) 158Potential energy surface for the formation of covalent complexes at shorter intermolecular 159distances; the energy of the physisorbed state is set to zero.

In the case of the covalent complexes, the adsorption stability decreases as the size of 161fullerenes increases. C_{24} , C_{26} , and C_{34} form highly stable complexes, reaching positive adsorption 162energies of ~2.1, 1.4, and 1.3 eV, respectively (favorable adsorption, Fig. 1c). Interestingly, 163**Phos-C**₂₄ and **Phos-C**₂₆ complexes form a [4+4]-cycloaddition bonding, which compares to 164prototype graphene–fullerene hybrids called graphene nanobuds, where covalently bonded 165fullerenes or fusing fragmented C_{60} are designed onto a graphene monolayer *via* cycloaddition 166reactions³⁸⁻³⁹. Otherwise, C_{36} , C_{40} , and C_{44} reach relative medium stability upon adsorption, with 167positive adsorption energies of up to 0.9 eV (favorable adsorption). Structurally, C_{34} , C_{36} , C_{40} , and 168C₄₄ fuse to phosphorene through cycloaddition of up [2+2] order, and adsorption energy 169decreases up to 67% compared to **Phos-C**₂₄ and **Phos-C**₂₆.

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170 On the other side, the C₅₀, C₆₀, and C₇₀ fullerenes are physisorbed onto phosphorene with 171positive adsorption energies of up to 1.0 eV, denoting stable adsorption that compares to the 172stability reached by C₃₆ and C₄₀ via chemisorption (Fig. 1c). Note also adsorbed fullerenes could 173be further stabilized by neighboring molecules onto phosphorene through strong molecule-174molecule interactions, arising two-dimensional self-assembly in the physisorption regime as 175reported for C₆₀ adsorption¹⁹. Compared to our results, periodic vdW-KBM calculations and 176 molecular dynamics simulations have reported non-covalent **Phos-C**₆₀ complexes with E_{ads} 177 values of ~ 0.7 and ~ -1.0 eV, respectively^{19, 23}; then, our results agree with previous reports. 178Additionally, UV-VIS-NIR absorbance, temperature scanning tunneling microscopy, and 179scanning tunneling spectroscopy measurements indicate that C60 molecules physically adsorb on 180phosphorene, where desorption occurs with annealing 400 K²⁰⁻²¹. As can be seen, $E_{ads} = \sim 0.9$ eV 181 for **Phos-C**₆₀ complexes agrees with the experimental favorable non-covalent adsorption of 182 fullerenes on phosphorene-based materials. Furthermore, the C₆₀ adsorption stability increases at 183least 30% compared to related 2D materials such as graphene, where $E_{ads} \approx 0.7$ eV is reported for 184Graphene-C₆₀ non-covalent complexes⁴⁰⁻⁴⁴. Therefore, phosphorene serves as excellent support 185 for fullerenes to form new hybrid nanostructures, ensuring a stable interaction.

We also compute the potential energy surface (PES) of non-covalent complexes to search 187for possible states where chemisorption occurs (Fig. 1d). PES begins at intermolecular distances 188of $d_{inter} \approx 2.8$ Å, where the physisorption state is set to zero. The non-covalent complexes turn into 189covalent complexes at $d_{inter} \approx 2.0$ Å, reaching [2+2]-cycloaddition configurations. Nevertheless, 190covalently bonded states do not reach saddle points in the PES, even without energy barriers. 191The latter indicates that PES is repulsive for $d_{inter} < 2.8$ Å, denoting chemisorption is unstable for 192**Phos-C**₅₀, **Phos-C**₆₀, and **Phos-C**₇₀ complexes. In other words, covalently bonded metastable

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193states will have a short lifetime and poor stability due to the lack of dissociation barrier to reach 194the non-covalent states. This behavior is different from graphene and carbon nanotubes, where 195C₆₀ fullerenes reach highly stable chemisorbed states at room temperature due to the relatively 196high dissociation barrier to reach the physisorbed states ($\Delta E^{\neq}>0.7 \text{ eV}$)^{39, 45}. The absence of an 197energy barrier to reach chemisorbed states is a consequence of phosphorene nonplanarity due to 198its shape of structural ridges; in contrast, the chemical Graphene-C₆₀ attachment requires a high 199energy barrier due to the lack of local puckering in the graphene structure^{39, 41}.



Fig. 2. Adsorption energies of the **Phos-Fullerene** complexes in solvents (*E*_{ads}, in eV).

Regarding stability in solvent media, almost all **Phos-Fullerene** complexes) show high 201stability in different solvents, independent of the solvent polarity (Fig. 2). In this way, solvation 202energies cause a slight penalty in the adsorption energies (\sim 30%). The exception is the **Phos-C**₄₄ 203and **Phos-C**₇₀ complexes, whose stability is decreased up to 96% in high/medium polar solvents 204due to differences in solute-solvent polarity with respect to the free fragments. Consequently, 205solvents with low dielectric constants would be adequate for **Phos-C**₄₄ and **Phos-C**₇₀ dilution, 206e.g., toluene and n-hexane. Another key point to emphasize, phosphorene can be sensitive and

2070xidized underwater and oxygen conditions⁴⁶. However, phosphorene oxidation does not imply 208that fullerene adsorption will be hindered, and new synthetic techniques are developed to 209improve the stability of phosphorene layered materials for several applications⁴⁷⁻⁴⁸. Despite the 210latter, recent reports highlight the physisorption of C_{60} during an assembly at the air-water 211interface contribute to protecting phosphorene thin films from oxidation and inhibiting the 212overlapping stacking or agglomeration of phosphorene nanosheets in solvents²⁰.

2133.2. Electronic properties.

Relevant electronic properties of the **Phos-Fullerene** complexes are displayed in Table 1. 215In terms of charge distribution, all systems show electron transfer in the Phos \rightarrow Fullerene 216direction. The negative charge on fullerenes after interaction (Q_{full}) is of the order of \sim -0.1 to 217-0.3|*e*| for covalently bonded fullerenes (C_{24} - C_{44}), while physisorbed fullerenes gain up to \sim 0.1|*e*| 218(C_{50} - C_{70}). Consequently, carbon fullerenes act as mild *p*-dopants for phosphorene, introducing up 219to \sim 0.3 holes/molecule on the substrate. Moreover, carbon, boron, boron nitride, and silicon 220carbide fullerenes have also behaved as *p*-dopants upon adsorption on related 2D materials such 221as pristine graphene and metal-doped graphene^{40, 42, 49-50}.

The electron transfer in the Phos \rightarrow Fullerene direction for all the complexes is consistent 223with the chemical potential (μ) of the fragments (Table 1): μ characterizes the response of the 224system energy with respect to changes in the electron number, thus related to the 225electronegativity χ through $\mu = -\chi^{51}$. Hence, electrons move from the system with high chemical 226potential (phosphorene, μ =-4.4 eV) to the one with low chemical potential (fullerenes, μ =-4.7 to 227-5.3 eV) for establishing new electronic equilibrium. Further, we found the low energy of the 228HOMO level of fullerenes avoids the electrons to flow from the HOMO of fullerenes to the

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236density difference [$\Delta \rho$ (r)], where yellow and red regions denote accumulation and depletion of 237electron density, respectively (Fig. 3a-c). For the covalent complexes (e.g., **Phos-C**₂₄ and 238**Phos-C**₄₀, Fig. 3a-b), there is a strong electron accumulation at the bonded carbon atoms of 239fullerenes, which is also accumulated in their surrounding C-C bonds. While electron depletion 240mainly comes from the interacting 3*p* lone-pair orbitals of P atoms in the substrate. These results 241suggest the holes remain localized in specific sites of the phosphorene after electron transfer and 242in the absence of external bias potential. The non-covalent complexes (e.g., **Phos-C**₆₀, Fig. 3c)



Fig. 3. a-c) Electron density difference $[\Delta \rho(\mathbf{r})]$ of representative **Phos-Fullerene** complexes. ³⁴Yellow and red densities stand for accumulation or depletion of electron density after the ³⁵Interaction, respectively. d) Dipole moment (μ_D) vs magnitude of fullerene charge ($|Q_{Full}|$) of all ³⁶the **Phos-Fullerene** complexes.

243displays an analogous $\Delta\rho(\mathbf{r})$ pattern but with a low magnitude due to the weak electron-transfer. 244The electron density rearrangements also induce dipole moments in the range of ~0.5-4.3 Debye 245(μ_D , Table 1); note all free systems are non-polarized in their free states ($\mu_D \approx 0$). The induced 246dipole moments show a directly proportional correlation to the charge transfer magnitude (Fig. 2473d). Also, dipole polarizability α of the complexes increases compared to free phosphorene 248(α >6.47·10⁻²² esu vs. 6.17·10⁻²² esu, Table 1), denoting an improving ability to acquire an electric 249dipole moment in proportion to external perturbations such as an electric field. Likewise, the 250polarizability of complexes increases as the polarizability of free fullerenes increases, which is 251directly proportional to the electron density volume of fullerenes.

Considering that electron transfer could be responsible for changes in the substrate energy 253levels because of the charge doping, we analyze the bandgap of the systems (Table 1). The 254HOMO-LUMO energy difference (Δ_{HL}) of intrinsic phosphorene is ~1.3 eV, denoting its 255semiconductor character as reported from previous first-principles calculations (~1.4-1.5 eV)⁵²⁻ 256⁵³. Conversely, carbon fullerenes show Δ_{HL} values in the range of 0.4 to 1.7 eV, which increase as 257their size increases (see parenthesis in Table 1). Upon interaction, Δ_{HL} values of **Phos-Fullerene** 258nanostructures decrease up to ~1.0 eV (78%) with respect to intrinsic phosphorene, this is 259increasing the metallic character. The covalent adsorption of fullerenes causes a larger decrease 260in the bandgap (56-78%). Nevertheless, we can establish a more general relation:

Small energy-bandgap fullerenes (Δ_{HL}<0.8 eV, typified by C₂₄ to C₅₀) largely increase the
phosphorene metallic character.

Medium energy bandgap fullerenes (Δ_{HL}>0.8 eV, typified by C₅₀ to C₇₀, and possibly the
bigger ones) slightly increase the phosphorene metallic character.

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265 In the final analysis, deposition of small fullerenes could be an efficient strategy for 266modulation of phosphorene bandgap *via* molecular doping, i.e., increasing its metallic character. 267The bandgap modulation mechanism is mainly due to LUMO stabilization and HOMO 268destabilization for **Phos-C**₂₄ to **Phos-C**₄₀ complexes because of the high charge transfer (Fig. 4). 269In the case of complexes Phos-C₄₄ to Phos-C₇₀, the bandgap decreasing emerges due to 270stabilization of the LUMO level of phosphorene. Considering related 2D materials such as 271graphene, the fullerene adsorption induces bandgap opening of $\sim 0.3-0.4 \text{ eV}^{54-55}$, which is due to 272the stress/strain externally exerted by fullerenes on the substrate structure⁵⁶. Therefore, we can 273conclude bandgap modulation in phosphorene could result from charge transfer and/or stress-274strain effects exerted upon fullerene attachment. About the reliability of these results, periodic 275DFT calculations have predicted bandgap values of ~ 1.0 eV for **Phos-C**₆₀ complexes²³, which 276 agrees with this work (Δ_{HL} =1.1 eV for **Phos-C**₆₀). Moreover, we compute bandgap values of $277 \sim 1.7$ eV for C₆₀ and C₇₀, which agree with those reported via photothermal deflection spectra and 278temperature-dependent microwave conductivity measurements⁵⁷⁻⁵⁸: 1.6-1.8 eV (C₆₀) and 1.6 eV 279(C₇₀). The latter ensures the reliability of the employed theoretical methodology.



Fig. 4. HOMO and LUMO energies of Phos-Fullerene complexes. HOMO and LUMO densities are displayed below and above, respectively; isosurface value of 0.003 a.u.

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Regarding the topology of frontier molecular orbitals in the complexes (Fig. 4), covalent 281complexes show the HOMO located in both phosphorene and fullerene moieties due to the 282orbital interactions. However, the fullerene contribution to the HOMO decreases as the chemical 283bonding character decreases until it almost disappears in non-covalent complexes. Conversely, 284LUMO is almost entirely located in fullerene for all the complexes. Thus, fullerenes act as active 285sites for orbital-controlled interactions, mainly retaining their donor/acceptor properties when 286chemisorbed on phosphorene. While fullerene mainly will act as acceptor sites when physisorbed 287on phosphorene.

2883.3. Interaction mechanism.

We provide a quantitative and readily physical interpretation of the adsorption 290mechanism by examining the specific role of the stabilizing effects into the adsorption energies 291and employing the EDA method. The attention is focused on the stabilizing contributions of Eq. 292(2) (ΔE_{CT} , ΔE_{POL} , ΔE_{ELEC} , ΔE_{DISP}) because destabilizing terms were up to 95% only due to Pauli 293repulsion (ΔE_{PAULI}). For a standardized interpretation among systems, relative single percentage 294contributions of EDA terms are discussed (% ΔE_i , Fig. 5a).



Fig. 5. Energy decomposition analyses: **a)** Relative single percentage contributions (% ΔE_i , in %) of stabilizing terms. **b)** Comparison between combined percentual contributions of short-range ($\Delta E_{CT}+\Delta E_{POL}$) and long-range ($\Delta E_{ELEC}+\Delta E_{DISP}$) terms. **c)** Adsorption energies (E_{ads} , in eV)

First, the stability of covalent complexes (**Phos-C**₂₄ to **Phos-C**₄₄) emerges from the 296charge-transfer term (ΔE_{CT}) because of the strong orbital interactions with a contribution of ~38-29748% to the stabilizing energy (Fig. 5a), agreeing with the high magnitude of electron transfer in 298the covalent complexes (Table 1). The second highest contribution to the stability emerges from 299polarization effects as expected in short-range orbital interactions (ΔE_{POL} , up to 32%). The latter 300denotes strong intramolecular charge rearrangements due to the on-fragment relaxation of each 301species to the presence of the nuclei and electrons of all other fragments. Then, covalent bonding 302is demonstrated from the physical viewpoint. It is worth mentioning that an obvious correlation 303between the number of transferred electrons and ΔE_{CT} magnitude cannot be established because 304 ΔE_{CT} term also accounts for energy lowering associate with intramolecular electron flow, 305allowing to delocalize electrons over the complex. The latter explains why the ΔE_{CT} contribution

306is high in **Phos-C**₂₄ (43%) in spite that the whole intermolecular charge transfer is low ($\sim 0.08|e|$, 307Table 1), which could be caused by strong intrafragment electron density rearrangements and 308bond polarization in the [4+4]-cycloaddition as noted from the high ΔE_{POL} contribution (32%).



Fig. 6. Electron density at the bond critical points of intermolecular interactions (ρ_1 , in *e*/Bohr³) for representative Phos-Fullerene complexes.

The AIM analysis allows assessing the binding mechanism from the values of electron 310density (ρ_i) at intermolecular bond critical points [BCPs, points in space at which the first 311derivatives of the electron density vanish $\nabla \rho(\mathbf{r})=0$, Fig. 6]. Accordingly, the covalent complexes 312reveal short-range intermolecular C-P bonding with ρ_i values of $\rho_i \approx 0.10-0.14 \ e/Bohr^3$, which 313associate to polarized covalent bonding by sharing electrons (red numbers in Fig. 6a-b). Natural 314bond orbital analyses (NBO) reveal at least 60% of the C-P bond density is polarized to fullerene 315carbon atoms due to their strong acceptor character. Therefore, the polarized covalent bonding in 316covalent complexes agrees with the high ΔE_{CT} and ΔE_{POL} contributions. In the molecular orbital 317picture, the bonding occurs when interacting carbon atoms of fullerenes hybridize from $sp^2 \rightarrow sp^3$, 49

318forming σ C-P polarized covalent bonds with the low-occupied lone-pair 3*p* orbitals of 319phosphorus atoms in the substrate. Additionally, BCPs appear at long-range C-P interactions 320($\rho_i \leq 0.02 \ e/Bohr^3$), which are associated with weak electrostatic interactions (blue numbers in 321Fig. 6a-b). In this regard, EDA shows the electrostatic term (ΔE_{ELEC}) reaches the third largest 322contribution to the stability (17-31%). The electrostatic contribution emerges from the electron-323sufficient carbon atoms of fullerenes, which acquire a negative charge that electrostatically 324attracts the electron-deficient phosphorous atoms of phosphorene as noted from electron density 325rearrangements [see the $\Delta \rho$ (r) surface, Fig. 3a-c].

It is necessary to note ΔE_{CT} and ΔE_{POL} contributions mainly decay as the intermolecular 327distances increase due to the connection to orbital overlap, becoming ΔE_{CT} and ΔE_{POL} mostly 328short-range terms. In that case, the combined contribution of short-range terms defines two 329classes of covalent complexes (Fig. 5b):

330 i. Members of the first class (typified by **Phos-C**₂₄, **Phos-C**₂₆, **Phos-C**₃₄) have short-range 331 terms influencing the magnitude of adsorption energies, which highly contribute 332 compared to long-range terms (electrostatics and dispersion): $\Delta E_{CT} + \Delta E_{POL} > \Delta E_{ELEC} + \Delta E_{DISP}$. 333 ii. The second class of covalent complexes is represented by **Phos-C**₃₆, **Phos-C**₄₀, and 334 **Phos-C**₄₄, where a balanced contribution of long-range and short-range driving forces 335 determine the magnitude of adsorption energies: $\Delta E_{CT} + \Delta E_{POL} \approx \Delta E_{ELEC} + \Delta E_{DISP}$.

In the framework of steric effects, we can establish that a relatively larger volume of 337fullerenes (C_{36} , C_{40} , C_{44}) would cause a larger steric repulsion at shorter intermolecular distances 338in the second class of complexes. In this manner, intermolecular distances slightly increase 339($d_{inter} \approx 2.1$ -2.2 Å) to compensate for the volume exclusion effects when fragments are brought

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340into close interaction. As a result, the strong stabilization gained by short-range contributions 341decay and weak long-range E_{ELEC} and ΔE_{DISP} effects compensate the steric destabilization; 342consequently, lower adsorption energies are reached compared to the first class of covalent 343complexes (Fig. 1b).

Focusing now on non-covalent complexes (Phos-C₅₀, Phos-C₆₀, and Phos-C₇₀), the 344 345stabilizing part of the adsorption energy mainly arises from permanent Coulombic electrostatic 346and dispersion driving forces ($\Delta E_{ELEC} + \Delta E_{DISP}$, up to 90%, Fig. 5a). In this regard, the 347minimization of dispersion and electrostatic interactions also plays a key role in forming organic 348molecular crystals, phthalocyanine self-assemblies, and DNA nucleobase patterns onto 349phosphorene-based substrates, standing for \sim 85% of the stability^{19, 59-61}. It can be pointed out 350although phosphorene is not a π -electron system, it behaves with a similar attractive ability than 351graphene or carbon nanotubes to bind aromatic molecules, where fullerenes adsorb by π - π 352stacking³⁹⁻⁴⁵. In this regard, the 3D δq^{inter} isosurface representation of weak intermolecular 353 interactions is visually displayed for **Phos-C**₆₀ complex as a representative case and compared 354against **Graphene-C**₆₀ (Fig. 7a). The weak interaction pattern shows a similar shape in both 355**Phos-C**₆₀ and **Graphene-C**₆₀, so dispersion forces (green regions) play a key role in both 356systems. This ability arises because of the larger amount of 3*p* lone pair electrons in the 357phosphorene upper lattice, which can correlate with the π -electrons of aromatic molecules and 358cause transitory induced dipoles responsible for dispersion forces⁶¹.

359 Additionally, δg^{inter} pattern shows that polar interactions (blue regions) have a higher 360intensity on phosphorene compared to graphene, denoting the major role of electrostatic driving

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362complexes displays two type of electrostatic interactions (Fig. 6c-d): i) weak electrostatics 363interactions at longer distances, typified by $\rho_1 \leq 0.02 \ e/Bohr^3$ (blue numbers); ii) strong 364electrostatic interactions, typified by $\rho_1 \approx 0.02-0.03 \ e/Bohr^3$ (green numbers). The NBO analysis 365also reveals strong electrostatics interactions involve a weak orbital overlapping due to electron 366donation from phosphorene 3*p* lone pairs towards low-occupied π^*C-C orbitals of fullerenes, 367while fullerenes show back-donation from single-occupied 2*p* orbitals towards unoccupied 368 σ^*P-P bonds of the substrate. When compared against the **Graphene-C**₆₀ complex, AIM 369analyses also show electrostatics interactions are favored in the **Phos-C**₆₀ case with lower 370electron density values for the former ($\rho_1 \leq 0.01 \ e/Bohr^3$, Fig. 7b). These results clarify how 371electrostatics driving forces favor the adsorption onto phosphorene with respect to related 2D 372materials. For instance, the **Phos-C**₆₀ system shows ΔE_{ELEC} and ΔE_{DISP} contributions of 45 and 37339%, respectively (major contribution of electrostatics, Fig. 5a); while the **Graphene-C**₆₀ system 374is reported with ΔE_{ELEC} and ΔE_{DISP} contributions of 33 and 67%, respectively (major contribution 375of dispersion)⁴⁰.

Otherwise, ΔE_{CT} and ΔE_{POL} terms show low stabilizing contributions in the non-covalent 377complexes because they are mainly short-range terms. Specifically, ΔE_{CT} shows a slightly larger 378contribution than ΔE_{POL} , with a contribution of up to 18% (**Phos-C**₅₀). ΔE_{CT} contribution in these 379cases relates to the donor-acceptor orbital interactions as noted above: donation $3803p(\text{Phos}) \rightarrow \pi^*\text{C-C}(\text{Fullerene})$, and back-donation $2p(\text{Fullerene}) \rightarrow \sigma^*\text{P-P}(\text{Phos})$. In this way, $381\Delta E_{\text{CT}}$ contributions decrease as the number of transferred electrons decreases.

382 Additionally, the choice of either covalent or non-covalent configurations can also be 383elucidated based on steric effects. Fig. 5c displays the adsorption energies and considering

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384chemisorption and physisorption adsorption configurations when possible. First, complexes 385Phos-C₂₄ prefer bind to Phos-C₄₀ to by covalent bonding, i.e., $386E_{ads}$ (chemisorption)> E_{ads} (physisorption); consequently, although covalent complexes can reach a 387stable physisorption state, the chemisorption will be thermodynamically preferred. Also, Fig. 4c 388displays the Pauli repulsion energy (ΔE_{PAULI}) in the most stable adsorption configurations. 389Accordingly, covalent complexes show high ΔE_{PAULI} values, but the charge-transfer and 390polarization effects overcompensate the Pauli repulsion ($\Delta E_{\rm CT} + \Delta E_{\rm POL} > \Delta E_{\rm PAULI}$), leading to stable 391adsorption. Note that as the fullerene size increases, the intermolecular distance increases to 392compensate for the repulsion energy. The energy penalty emerges as kinetic energy pressure 393exerted by electrons when atoms are brought into close interaction³⁴. As a result, the ΔE_{PAULI} term 394decreases as the adsorption energy decreases until the Phos-C₄₄ complex. In other words, we 395 found at this point the fullerene size at which chemisorption is no longer allowed, which is set 396between C₄₄ and C₅₀ (Fig. 5c). This limit arises because the intermolecular separations must 397 increase to decrease the repulsion energy; consequently, physisorption is a stable state at longer 398intermolecular distances for bigger fullerenes (C₅₀, C₆₀, and C₇₀). Therefore, ΔE_{PAULI} is the 399primary driving force that compensates all the stabilizing effects to establish the energetically 400optimal intermolecular separation between phosphorene and fullerenes.

Finally, destabilizing terms mainly arise from Pauli repulsion; however, the covalent 402complexes involve geometrical preparation energies ΔE_{PREP} due to the required orbital 403hybridization $sp^2 \rightarrow sp^3$ of fullerenes and rearrangements of phosphorene sublattices, which equals 404to contributions of 10-16% to the destabilizing energy. The root-mean-square deviation RMSD 405of the fullerene geometries in the complex ranges from 0.04 to 0.94 Å with respect to the free 406molecules, which increase with the order of cycloaddition reaction. Although there is an energy

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407penalty due to weak structural changes to reach the covalent bonding, the low RMSD values 408show that the electronic/structural rearrangements are low-scale and are not prohibited.

4093.4 Outlook of Phos-Fullerene nanostructures.

This work provides a first pivotal stepping-stone to exploring the immense possibilities of 411the proposed nanostructures in future theoretical/experimental studies. All analyses considered, 412in addition to comparisons with related hybrid structures, show some interesting potential uses 413for **Phos-Fullerene** nanostructures. For instance, covalent **Phos-Fullerene** complexes display a 414graphene nanobud-like structure and strong electron transfer in the Phos→Fullerene direction. In 415this regard, graphene nanobuds have shown potential uses in energy and gas storage devices with 416high stability at room temperature, new spintronics, position sensing, and graphene band 417structure engineering^{38-39, 54, 62}. Furthermore, Li adsorption was enhanced for graphene–C₆₀ 418nanobuds by the high electron affinity of C₆₀ and the charge transfer from graphene to C₆₀^{55, 63}. 419Accordingly, the strong charge transfer in the Phos→Fullerene direction turns covalent 420complexes into remarkable candidates to be studied as novel storage and anode materials in 421Li/Na-ion battery applications.

Additionally, orbital analyses show that fullerenes act as active sites for orbital-controlled 423interactions, mainly retaining their donor/acceptor properties when chemisorbed on phosphorene. 424While fullerene mainly will act as acceptor sites when physisorbed on phosphorene. Also, 425fullerenes increase the metallic character of phosphorene. The latter suggest fullerenes could act 426as signal amplifiers for specific redox applications considering that phosphorene-based materials 427have shown interesting electrochemical performance as electrode materials. Deposited 428mesoporous and organic molecules have caused effective signal amplification on phosphorene-

429based sensors to determine prostate-specific antigen, hemoglobin, clenbuterol, and 430polychlorinated biphenyls, reaching an excellent specificity, chemical stability, low detection 431limits, and reproducibility⁶⁴⁻⁶⁶. The electrochemical amplification mechanism emerges from the 432good charge transfer in the hybrid material, an enhanced electrical conductivity of the electrode, 433and the excellent electrocatalytic activity of the deposited molecules upon adsorption⁶⁵. 434Therefore, our results open avenues for the study of **Phos-Fullerene** hybrids in electrochemical 435biosensing.

On the other hand, charge-doping and bandgap changes reveal phosphorene conductance 437can be tailored by deposition of fullerenes. Since the bandgap value is directly proportional to the 438conductance ($\sigma \propto \Delta_{HL}/kT$, where *k* is the Boltzmann constant and *T* the temperature), our results 439define a framework to explore the charge-transport and optoelectronic response of **Phos**-440**Fullerene** nanostructures. As an illustration, the increased charge transport of graphene due to 441C₆₀ deposition⁵⁴⁻⁵⁵, and the strong electron-accepting characteristics of C₆₀ causing a hole-doped 442nature in graphene, lead to hybrid structures allowing photogenerated carriers for high-443performance photodetector devices⁵⁰. It is also noticed that a larger electron transfer in the 444Phos→Fullerene direction is responsible for lower bandgap values, resulting in a higher 445conductance. In this way, functionalized fullerenes with strong acceptor functional groups could 446be implemented to maximize the electron transfer and charge transport.

Furthermore, small bandgaps would enable UV-Vis radiation absorption by 448Phos-Fullerene nanostructures as required for optoelectronic applications. In fullerene-based 449polymer solar cells, phosphorene has enabled a strong absorption between 300-800 nm and better 450energy alignment in the device layers, which is significantly beneficial to the charge transfer, 451exciton dissociation, and reduce charge recombination, consequently enhancing the power

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452conversion efficiency¹⁸. In addition, C₆₀ allowed absorption in the visible range for mm-scale 453heterojunction phosphorene-based photodetectors, where synergistic effects enable high 454photoresponse from the visible to near-infrared spectrum²⁰. Accordingly, synergistic properties 455of **Phos-Fullerene** complexes could inspire experimental/theoretical studies on its uses for future 456generation solar energy harvesters or as active layers within photodetector devices. To support 457these points, we verified the UV-Vis spectrum of the proposed nanostructures in n-hexane as a 458solvent commonly implemented to study the fullerene absorbance (Fig. 8). Intrinsic phosphorene 459shows a wide absorbance from the infrared, which increases in intensity below 500 nm as 460recorded from UV–vis spectrophotometry⁶⁷. In comparison, most free fullerenes absorb high 461molar absorption coefficients at the UV region below λ_{abs} <400 nm (Fig. 5a). **Phos-Fullerene** 462nanostructures almost resemble the absorption spectra of free phosphorene (Fig. 5b), but a 463remarkably improved absorption at higher frequencies is reached ($\lambda_{abs} \approx 400$ -500 nm). Then, 464fullerenes maximize the phosphorene absorbance at the UV-Vis region, denoting the emerging 465synergistic effects for optoelectronics and photodetectors.

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467**Fig. 8.** Absorption spectra of **a**) free phosphorene and fullerenes; **b**) Phos-Fullerene complexes.

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4684. Conclusions

469 We have computationally elucidated the structure, stability, and interaction mechanisms 470of phosphorene-fullerene nanostructures, which form covalent and non-covalent complexes 471depending on the fullerene size and outstanding stability compared to related 2D materials, even 472in solution. Two classes of covalent complexes arise by cycloaddition-like reactions of up to 473[4+4] order: the first class (C_{24} - C_{34}), where short-range stabilizing effects dominate; and the 474second one (C₃₆-C₄₄), where short-range effects decay to avoid steric repulsion, and balanced 475long-range forces favor the stability. High-size fullerenes (C₅₀-C₇₀) form non-covalent complexes 476due to strong repulsion at shorter intermolecular distances and lack of dissociation barriers. 477Furthermore, fullerenes act as mild *p*-dopants for phosphorene, increasing its polar character and 478ability to acquire induced dipole moments (polarizability). Also, small energy-bandgap 479fullerenes (<0.8 eV) largely increase the phosphorene metallic character. We also note fullerenes 480retain their donor/acceptor properties upon adsorption, acting as active sites for orbital-controlled 481interactions and maximizing the phosphorene absorbance at the UV-Vis region. Future work 482should focus on relevant applications of the proposed nanostructures for storage, anode materials 483in Li/Na-ion batteries, electrochemical sensing, bandgap engineering, and/or optoelectronics.

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