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Substrate charge transfer drives the adsorption site of metal-phthalocyanines and porphyrins on coinage metal surfaces†

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The Frontier electronic structure of tetraphenylporphyrinato (TPP²⁻) and phthalocyaninato (Pc²⁻) square planar transition metal complexes (MTPP and MPc; M = V, Cr, Mn, Fe, Co, Ni, Cu, and Zn) has been revisited through DFT calculations. The different ground state symmetry and spin multiplicity between MPc and MTPP of the same M is shown to originate from the different Pc²⁻ and TPP²⁻ ligand field, stronger in the former ligand than in the latter. The corresponding spatial localization and symmetry of the unoccupied molecular orbitals postulate unescapable geometric constraints to their overlap with the electron cloud of a crystalline metal surface. From comparison with literature experimental evidence, we show that the adsorption geometry (atomic site and azimuthal orientation) of MTPPs and MPcs on the low index crystal planes of coinage metals (CM = Au, Ag, Cu) may be predicted when two conditions are satisfied: (i) evidence of a surface → adsorbate charge transfer, (ii) absence of significant distortion of the macrocycle upon adsorption. In this regard, the overall susceptibility to charge transfer is determined by the strength of the molecular ligand field (*i.e.*, charge transfer to MPc is more favoured than to MTPP) and inversely linked to the electronegativity of the surface atoms (being Au the most inert CM substrate thanks to its highest electronegativity).

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† Electronic supplementary information (ESI) available: Frontier electronic structure of MTPP and MPc (M = V, Cr, Mn, Fe, Co, Ni, Cu, and Zn) isolated complexes. Optimized coordinates of Py, i-Ind, H₂P, H₂Pz, H₂TPP, H₂Pc, P²⁻, Pz²⁻, TPP²⁻, and Pc²⁻ in Tables S1–S10; optimized coordinates of MTPP and MPc in Tables S13, S16–S24, S27, S28, S30, S31, S33, and S34; optimized coordinates of CoPc⁺, NiPc⁺, CuPc⁺, and ZnPc⁺ in Tables S25, S29, S32, and S35; H₂TPP, H₂Pc, MTPP, and MPc TSEs in Table S11; MTPP and MPc optimized/experimental M–N^{Py} bond lengths in Table S12; MTPP and MPc GS terms and low-lying excited state terms sharing the same GS spin multiplicity in Table S14; MTPP and MPc TSEAs in Table S15; theoretical/experimental ν values of the IR-active 22e_u vibrational mode of MPc and MPc⁺ (M = Co, Ni, Cu, and Zn) in Table S25; selected internuclear distances (Å) in: D_{2h} H₂P and H₂Pz (Fig. S1); D_{2h} H₂TPP and H₂Pc (Fig. S2); D_{4h} P²⁻ and Pz²⁻ (Fig. S3); D_{4h} TPP²⁻ and Pc²⁻ (Fig. S4); energy position of H₂P, H₂TPP, H₂Pc, and H₂Pz GS FMOs (Fig. S5); 3D CPs of the Py and i-Ind HOMOs (Fig. S6); 3D CP of the i-Ind 1a₂ π MO (Fig. S7); 3D CPs of H₂P FMOs generated by the 1a₂ Py HOMO (Fig. S8); 3D CPs of low-lying H₂P π MOs (Fig. S9); parenthood between ^{pmc} π and ^{pmc} π^* Frontier MOs upon D_{2h} → D_{4h} switching (Fig. S10); 3D CPs of low-lying TPP²⁻ and Pc²⁻ π^* MOs (Fig. S11); superposition of the CoPc (²A_{1g}) and CoPc⁺ (²A_{1u}), NiPc (¹A_{1g}) and NiPc⁺ (²A_{1u}), CuPc (²B_{1g}) and CuPc⁺ (²B_{1u}), ZnPc (¹A_{1g}) and ZnPc⁺ (²A_{1u}) optimized structures in Fig. S12; schematic representation of the ⁵CM *ns* AOs SALCs in Fig. S13–S17. See DOI: <https://doi.org/10.1039/d5cp01576f>

1. Introduction

The control of the interfacial structure between organic/metal-organic molecular layers and the underlying substrate is crucial to reproducing the functional properties of the organic/metalorganic component in surface-supported devices. Even tiny alterations of the interface structure may severely modify the behavior of the overlayer both at the supramolecular and the single-molecule level, thus generating significant changes independently of the investigated property (mechanical,¹ electronic,^{2–4} optical,^{5,6} magnetic^{7,8}) or the application field (sensing, catalysis, light-to-energy conversion, molecular electronics, nonlinear optics, spintronics). Among surface-supported layers, those made up of transition-metal complexes of porphyrin-related molecules (see the upper panel of Fig. 1)‡ are particularly relevant because of the pivotal role played by porphyrins not only in fundamental biological processes such as oxygen transport and storage, the photosynthesis, and the electron transport during cellular respiration

‡ Porphine (H₂P) is an aromatic (4*n* + 2; *n* = 4 for the shortest cyclic path), heterocyclic, organic compound consisting of four pyrrole fragments held together by as many methylene bridges, which makes it the simplest tetrapyrrole (C₂₀H₁₄N₄). Because of the H₂P low solubility, the interest for the pristine molecule is mainly theoretical. Substituted porphines correspond to porphyrins; *meso*-tetraphenylporphyrin (H₂TPP) is the most common.⁹ The acronym herein used for M complexes of H₂TPP is MTPP.



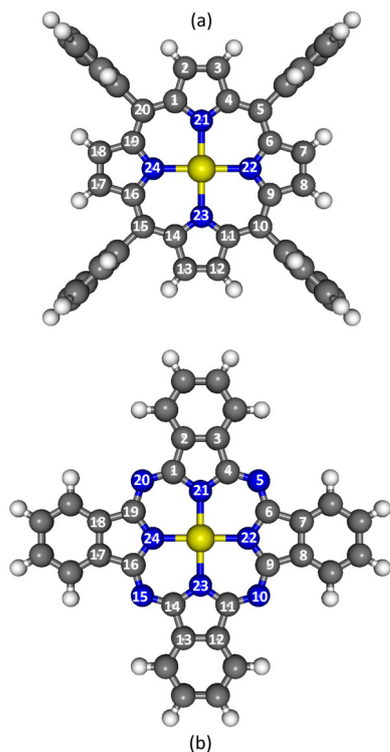


Fig. 1 Idealized representation of D_{4h} MTPP and MPC isolated molecules. White, gray, blue, and yellow spheres are representative of H, C, N, and M atoms, respectively. In the adopted framework, M lies in the origin O of the coordinate axes, M–N^{Py} bonds are aligned with OX and OY axes, and the σ_h plane corresponds to the XY plane. The atom numbering adopted for the planar, aromatic macrocycle core (pmc) is that recommended by IUPAC.

and photosynthesis,^{10,11} but also in technological fields ranging from electronics¹² to solar cells,¹³ and sensors,¹⁴ thus justifying the cross-disciplinary interest towards them and motivation to develop new porphyrin-like species, whose electronic and optical properties may be tuned through molecular engineering.^{15–23}

Unlike porphyrins, neither the H₂Pc free-base nor MPC complexes (see the lower panel of Fig. 1) are present in Nature. Nonetheless, they have been attracting a growing interdisciplinary interest^{20,24–30} since their synthesis by serendipity at the end of the twenties of the last century^{31–39} because their technological potential spans a huge range of applications including catalysts,²⁶ dyestuffs for textiles and inks, intrinsic semiconductors, chemical sensors, organic light-emitting diodes, organic photovoltaic cells, thin-film transistors, materials for nonlinear optics, spintronics, and laser recording as well.^{40–43}

From a structural point of view, isolated MTPPs are usually characterized by a planar, aromatic macrocycle core (hereafter, pmc) consisting of four Py-like rings held together by four methine groups in *m* positions (see the upper panel of Fig. 1 where the adopted atom numbering is that recommended by IUPAC and *m* sites correspond to the 5, 10, 15, and 20 positions). Additionally, 1, 4, 6, 9, 11, 14, 16, and

19 (2, 3, 7, 8, 12, 13, 17, and 18) positions are commonly referred to as α -positions (β -positions). Similarly to MTPPs, isolated MPCs consist of four *i*-Ind (a benzene ring fused with Py) fragments held together by four aza-nitrogen atoms occupying the pmc *m* positions (see the lower panel of Fig. 1 where the adopted atom numbering is again the one recommended by IUPAC).

Upon moving from the isolated to the adsorbed species, the coordinative arrangements of the on-surface stabilized arrays of M ions may be exploited to boost free molecules' peculiar properties,^{44,45} to stabilize M in unusual spin and oxidation states^{46–53} or, in wider terms, to control and tune their chemical reactivity.^{54–58} As such, a crucial issue is the pmc adsorption site, usually determined by combining the outcomes of STM measurements with the results of expensive and time-consuming quantum mechanical calculations, commonly based on DFT.^{20,23} However, DFT-based numerical experiments have often been restricted to the generation of phenomenological descriptions rather than exploring in detail the anchoring configurations of the adsorbate–substrate pair.^{20,23}

Li *et al.*²⁵ tackled the single-molecule chemistry of MPCs on the (111) surface of group 11 elements (CM) by combining first-principles simulations with STM, which, besides information about the local adsorption geometry and the electronic properties of chemisorbed MPCs, was exploited to carry out the so-called single-molecule surgery to control the Kondo effect⁵⁹ at the MPC/CM(111) interface. Nevertheless, the DFT numerical experiments carried out by Li *et al.*²⁵ did not provide any atomistic modeling of the MPC–CM(111) anchoring configuration to explain, for instance, why the chemisorption site of the same MPC may be different on the (111) surface of diverse CM or why the chemisorption site of diverse MPCs may be different on the (111) surface of the same group 11 element. In this regard, we recently proposed convincing modeling of the NiTPP anchoring to the Cu(100) surface, as well as an atomistic view of the Cu(100) → NiTPP charge transfer taking place at the interface.⁶⁰ Starting from the experimentally reported Ni^{II} (d⁸) → Ni^I (d⁹) reduction at the interface,⁶¹ the NiTPP chemisorption site and the molecular orientation on Cu(100) can be unequivocally assigned as confirmed by the structural outcomes of PED measurements.⁶⁰ The charge transfer from specific substrate atoms to specific adsorbate atoms follows the spatial superposition of the symmetry-allowed substrate and molecular orbitals, according to the matching of the intramolecular atomic structure with the substrate lattice spacing. More specifically, symmetry and geometry arguments made it practically unessential to carry out costly numerical experiments to assess the chemisorption configuration. Our analysis relied on identifying the NiTPP lowest-lying unoccupied MOs and the SALCs of the substrate topmost atoms (⁶³Cu) 4s AOs involved in the charge transfer process. The latter point implies that each ⁶³Cu participates in the adsorbate/substrate interaction, with its single electron occupying its 4s AO, while the electrons of the completely occupied 3d shell are simple bystanders.



The use of MTPP and MPc symmetry, orbitals, and spectra⁶² coupled with a descriptor able to provide information about the different nobleness of group 11 elements turned out to be Hobson's choice to set up a protocol able to provide a semi-quantitative modeling of the $^5\text{CM} \rightarrow \text{MTPP}/\text{MPc}$ charge transfer when MTPP/MPc complexes are chemisorbed on CM (100), (110), and (111) surfaces. In this regard, it is notable that Kepp thoroughly investigated the chemical causes of M nobleness a few years ago.⁶³ More specifically, he considered all the group 10 and 11 elements as well as the heaviest element of group 12 (Hg), and he tested 12 different descriptors, including, by the way, the d-band center energy of the solid, the first IE and the EA, the Pauling electronegativity χ ,⁶⁴ the bulk polycrystalline M work function, the relativistic s-shell contraction, the oxophilicity of M, and the cohesive free energy (free energy of atomization) of the bulk M state, to conclude that the M's χ explains best the M nobleness.⁶³ The noblest metal is then Au, whose high χ (2.54 Pauling's units, the highest among metals)^{65,66} is determined by concurrent effects such as the high effective nuclear charge and the contraction of the half-filled 6s AO.⁶⁷ As such, Pyykkö^{67–69} pointed out that the ratio of relativistic and nonrelativistic 6s shell radii in the atomic GS of the elements with the atomic number Z ranging from 55 to 100 has a marked minimum for $Z = 79$, and there is often talk of the “gold maximum” of relativistic effects in group 11, whose elements are all characterized by the electronic configuration $d^{10}s^1$.⁶⁵

Even though the high $^{\text{Au}}\chi$ ^{65,66} provides a rationale for understanding the inertness of gold surfaces independently of Miller indices and chemisorption sites' local symmetry, genuine unpublished results and literature data will be compared with the provisions of the proposed approach in the forthcoming discussion. In more detail: (i) results on the Frontier electronic structure of isolated MTPP and MPc ($M = \text{V}, \text{Cr}, \text{Mn}, \text{Fe}, \text{Co}, \text{Ni}, \text{Cu}, \text{Zn}$), critically revisited and compared with literature experimental and theoretical evidence in the ESI,[†] are summarized in Paragraph 3.1; (ii) clean, bulk-terminated CM (100), (110), and (111) surfaces have been considered in Paragraph 3.2 by focusing on the local symmetry properties of their most common chemisorption sites; (iii) the proposed approach is applied in Paragraph 3.3 to selected MTPP/CM and MPc/CM interfaces (sub-Paragraphs 3.3.1–3.3.5) representing as many case studies.

2. Computational details

The GS electronic configurations of MPcs ($Z^{\text{M}} = 25\text{--}30$) have been thoroughly investigated by multireference electronic structure methods in the past.⁷⁰ Nonetheless, the GS structural and electronic properties of free MTPP and MPc complexes ($Z^{\text{M}} = 23\text{--}30$) have been re-investigated herein at the DFT level by exploiting the ADF package.⁷¹ ADF numerical experiments have been carried out within the assumption of an idealized D_{4h} symmetry⁷² (see Fig. 1) and by running nonrelativistic, spin-polarized calculations with generalized gradient corrections

self-consistently included through the BP86 formula.^{73,74} A triple- ζ with a polarization function Slater-type basis set has been adopted for all the atoms; moreover, the $(1s-2p)^{\text{M}}, 1s^{\text{N}}$, and $1s^{\text{C}}$ cores have been kept frozen throughout the calculations. \S ¶ IEs and EAs of valence MOs have been estimated through spin-polarized TS calculations.⁷⁵ The adopted set-up allowed the comparison of ADF results with homogeneous theoretical outcomes pertinent to CoTPP,^{51,52,56,76} NiTPP,^{52,53,60} CuTPP,^{77,78} MPc ($M = \text{V},^{79-81} \text{Cr},^{79} \text{Mn},^{79} \text{Fe},^{79}$ and Cu^{81-84}). In a few selected cases (*vide infra*), further numerical experiments have been carried out by employing the same ADF package, the same basis sets, and a hybrid functional (B3LYP),⁸⁵ incorporating a portion of exact exchange (20%) from Hartree–Fock theory.

3. Results and discussion

3.1 Isolated MTPP and MPc

The square planar arrangement of M^{II} ions in both MTPP and MPc lifts the five-fold degeneracy of the M 3d AOs to generate, within the assumption of a local D_{4h} symmetry⁷² and the framework adopted in Fig. 1, five spin-up orbitals (SO^{\uparrow}) of symmetry a_{1g} (z^2), b_{1g} (x^2-y^2), b_{2g} (xy), e_g (xz, yz) and as many SO^{\downarrow} of the same symmetry. These SOs may be grouped in parallel (\parallel) and perpendicular (\perp) $\pi_{t_{2g}}$ -like ($\pi_{\parallel}b_{2g} + \pi_{\perp}e_g$) SOs and σ_{e_g} -like SOs ($\sigma_{a_{1g}} + \sigma_{b_{1g}}$) by exploiting the parenthood between D_{4h} and O_h complexes.⁷² It is noteworthy that the M σ_{e_g} -like $\sigma_{b_{1g}}$ SO, antibonding in nature with respect to the M–N^{Py} σ interaction, is certainly occupied in the $3d^5$ HS MnTPP,⁸⁶ $3d^9$ CuTPP^{78,87,88} and CuPc,^{81-84,89-91} and $3d^{10}$ ZnTPP and ZnPc complexes;⁹⁰⁻⁹² moreover, only the closed-shell ZnTPP and ZnPc have no vacancy in the Zn 3d-based SOs. Additionally, the Pc^{2-} ligand field is experimentally and theoretically found slightly stronger than the TPP^{2-} one^{78,86,93} (see Section ESI.1 of the ESI[†]). This last evidence scarcely affects MTPP and MPc complexes whose $\pi_{t_{2g}}^{\uparrow}/\pi_{t_{2g}}^{\downarrow}$ -like SOs are filled, and the $^2A_{1g}$ ($\text{Co}^{\text{II}}, 3d^7$), $^1A_{1g}$ ($\text{Ni}^{\text{II}}, 3d^8$), $^2B_{1g}$ ($\text{Cu}^{\text{II}}, 3d^9$), and $^1A_{1g}$ ($\text{Zn}^{\text{II}}, 3d^{10}$) GSs uniquely determined; \parallel however, it could be relevant in lighter complexes, whose GS electronic terms are still debated^{79,94} and could be different upon moving from MTPP to MPc.^{78,86,93}

As mentioned in the Introduction, the frontier electronic structure of MTPP and MPc isolated complexes has been critically revisited and thoroughly compared with literature data in the ESI[†] (Section ESI.2). The relevant results of such a review will be briefly discussed by referring to Tables 1, 2, and Fig. 2.

Starting from the occupied FMOs and, more specifically, from the $^{\text{pmc}}\pi$ ones; *i.e.*, the $\text{H}_2\text{TPP } 13b_{1u}$ and $10a_u$ FMOs, the

\S ADF outcomes about isolated MTPP and MPc are compared with the results of the literature in the ESI.[†]

¶ Throughout the paper, the MO numbering corresponds to all-electron calculations independently of adopting the frozen core approximation.

\parallel Both $\text{Co}^{\text{II}} (3d^7)$ and $\text{Ni}^{\text{II}} (3d^8)$ complexes are characterized by a low-spin (LS) state with $S = \frac{1}{2}$ and $S = 0$, respectively (see Section ESI.2, ESI[†]).



Table 1 H₂TPP (H₂Pc) TSIEs (eV) of the 10a_u and 13b_{1u} (7b_{1u} and 4a_u) MOs.[†] MTPP and MPc TSIEs of the M-based $\sigma_{a_{1g}}$, $\sigma_{b_{1g}}$, $\pi_{b_{2g}}$, π_{e_g} , and pmc-based $\pi_{a_{1u}}$ and $\pi_{a_{2u}}$ SOs. Values of the lowest TSIE are in bold[‡]

	$\sigma_{a_{1g}}$	$\sigma_{b_{1g}}$	$\pi_{b_{2g}}$	π_{e_g}	a_{1u}^{\downarrow}	a_{2u}^{\downarrow}
H ₂ TPP	—	—	—	—	6.77 (10a _u)	6.48 (13b _{1u})
H ₂ Pc	—	—	—	—	6.57 (4a _u)	7.74 (7b _{1u})
VTTP	6.29 [†]	—	6.37 [†]	5.69[†]	6.72	6.50
VPc	6.74 [†]	—	6.59 [†]	6.17[†]	6.49	7.98
CrTPP	7.28 [†]	—	7.69 [†]	6.63 [†]	6.69	6.54
CrPc	7.73 [†]	—	7.44 [†]	7.03 [†]	6.50	7.99
MnTPP	9.61 [†]	5.88[†]	9.49 [†]	10.53 [†]	6.55	6.36
MnPc	8.40 [†]	—	9.33 [†]	5.99[†]	6.57	8.00
FeTPP	6.18[†]	—	6.45 [†]	7.08 [†]	6.70	6.57
FePc	6.65 [†]	—	6.77 [†]	8.63 [†]	6.51	8.01
CoTPP	8.94 [†]	—	7.60 [†]	6.85 [†]	6.69	6.60
CoPc	9.14 [†]	—	7.62 [†]	7.32 [†]	6.54	8.01
NiTPP	7.65 [†]	—	8.63 [†]	7.03 [†]	6.70	6.62
NiPc	8.36 [†]	—	8.03 [†]	7.47 [†]	6.55	8.03
CuTPP	9.72 [†]	6.95 [†]	9.39 [†]	9.64 [†]	6.72	6.55
CuPc	9.17 [†]	7.28 [†]	8.08 [†]	7.77 [†]	6.56	8.01
ZnTPP	—	—	—	—	6.72	6.54
ZnPc	—	—	—	—	6.55	7.98

[†] HS MnTPP TSIE calculations have been run by adopting the B3LYP functional.

H₂Pc 7b_{1u} and 4a_u FMOs, the MTPP 12a_{2u} and 2a_{1u} FMOs, and the MPc 6a_{2u} and 2a_{1u} FMOs, the following statements hold: (i) TSIEs of the 12a_{2u} (13b_{1u} in H₂TPP)** FMO are very similar along the whole TPP series (the IE range they cover is less than 0.2 eV wide; see Table 1 and Fig. 2, where the MnTPP 12a_{2u} FMO TSIE is not considered because evaluated by running B3LYP-based TS calculations (see Section ESI.2, ESI[†])); (ii) TSIEs of the 6a_{2u} (7b_{1u} in H₂Pc)** FMO are very similar along the whole Pc series and vary between 7.74 and 8.01 eV (see Table 1 and Fig. 2); (iii) nodal properties of the 2a_{1u} MO (the 10a_u FMO in H₂TPP and the 4a_u FMO in H₂Pc)** make corresponding TSIEs very similar to each other; the $|\Delta\text{TSIE}|$ between the 12a_{2u} and 2a_{1u} (10a_u and 13b_{1u} in H₂TPP)** FMOs is tiny (~ 0.2 eV) in MTPP; in addition, the ΔTSIE between the 6a_{2u} and the 2a_{1u} (7b_{1u} and 4a_u in H₂Pc)** FMOs is rather constant (~ 1.5 eV) in MPc. Moving to the M 3d-based occupied FMOs, photoemission processes involving an M 3d-based MO and lying at the lowest IE are limited to VTTP, FeTPP (TSIEs of MnTPP FMOs have been computed by running B3LYP-based TS calculations; see Section ESI.2, ESI[†]), VPc, and MnPc.

Data on unoccupied MTPP and MPc FMOs are, for this contribution, much more interesting to be jointly considered. Let us start from the $\text{pmc}\pi^*$ FMOs; *i.e.*, the quasi degenerate 12b_{2g}/12b_{3g} and the 11a_u FMOs in H₂TPP, the quasi degenerate 6b_{2g}/6b_{3g} and the 5a_u FMOs in H₂Pc, and the e_g and b_{1u} FMOs in MTPP and MPc (see Table 2 and Fig. 2). Similarly to the free H₂TPP and H₂Pc species, the $\text{pmc}e_g$ and $\text{pmc}b_{1u}$ MPc TSEAs lie deeper in energy than the MTPP ones; moreover, the MPc $\text{pmc}b_{1u}$ TSEAs (see Table 2 and Fig. 2) and, separately, the MTPP $\text{pmc}b_{1u}$ TSEAs are scarcely affected by the presence of different

** Upon the D_{2h} \rightarrow D_{4h} switching, the following correlations hold between irreducible representations (ir): (B_{2g} + B_{3g}) \rightarrow E_g; A_u(C₂) \rightarrow A_{1u}; A_u(C₂) \rightarrow B_{1u}; B_{1u}(C₂) \rightarrow A_{2u} (see Appendix 3 of ref. 72).

M along the two series because of the symmetry forbidden participation of the M AOs to the b_{1u} MOs. Moving to the M 3d-based unoccupied FMOs, two things are particularly relevant: (i) both in MTPP and MPc, the largest EA corresponds to an M 3d-based unoccupied FMO only in V ($\pi_{e_g}^{\uparrow}$) and Co ($\sigma_{a_{1g}}^{\downarrow}$) complexes (see Table 2 and Fig. 2); (ii) among the MTPP complexes, CoTPP is the only one with both σ_{e_g} -like FMOs (the $\sigma_{a_{1g}}^{\downarrow}$ and $\sigma_{b_{1g}}^{\downarrow}$ SOs) able to be involved in a charge transfer processes (see Table 2 and Fig. 2).

3.2 Bulk-terminated CM surfaces

Copper, silver, and gold share the same space group 225 (*Fm*3*m*) and the cubic-close-packed (ccp) crystal structure; corresponding cell parameters are $a = b = c$ ($a^{\text{Cu}} = 3.61496$ Å; $a^{\text{Ag}} = 4.0853$ Å; $a^{\text{Au}} = 4.0782$ Å) and $\alpha = \beta = \gamma = 90^\circ$.⁹⁵ A schematic representation of the two outermost layers of the bulk terminated CM(100) and CM(110) surfaces is displayed in Fig. 3,^{††} which also includes a top view of the three CM(111) unreconstructed topmost layers (high-symmetry chemisorption sites are also pinpointed in the figure).

The inspection of Fig. 3 reveals that local symmetry properties of these sites are quite different. More specifically, both CM(100) T and H sites are characterized by a local fourfold symmetry (*C*₄), reduced to a local *C*₂ symmetry when the B site is considered; (ii) no CM(110) chemisorption site (T, H, SB, and LB) exceeds a local *C*₂ symmetry; (iii) CM(111) T, H, and B sites have a local *C*₆, *C*₃, and *C*₂ symmetry, respectively. Incidentally, details of possible surface reconstructions, such as those affecting all the Au low-index planes, are neglected herein.^{‡‡} More generally, an interacting adsorbate (*i.e.*, subject to charge transfer) will locally disrupt the collective (periodic) surface properties, and the frontier orbitals of the ^SCM atoms underneath the adsorbate will recover their atomic-like symmetry.

Coming back to the topmost layer of the bulk terminated CM surfaces, the single *ns* AO ($n = 4, 5,$ and 6 for Cu, Ag, and Au, respectively) localized on a ^SCM T site (^TCM), independently of the local symmetry, will be a basis for the totally symmetric *ir* (a, in the Schoenflies notation).⁷² The situation is a bit more complicated when SALCs of the *ns* AOs centered on chemisorption sites' neighbours are considered: (i) the four SALCs of the *ns* AOs centered on the ^TCM(100) nnn and ^HCM(100) nn (^SCM_{100}^nnn and ^SCM_{100}^nn, respectively) are bases for the *irs* a, b, and e (see Fig. S13 and S14 of the ESI[†]) of the *C*₄ point group,⁷²}}

^{††} The internuclear distance between ^SCMⁿⁿ is the same, independently of the surface Miller indexes: 2.556, 2.889, and 2.884 Å in Cu, Ag, and Au, respectively.⁹⁵

^{‡‡} The complex herringbone reconstruction of Au(111) corresponds to a compressed atomic layer with an average lattice spacing of 2.76 Å along the [110] direction,⁹⁶ instead of the corresponding bulk plane value of 2.88 Å, despite the hexagonal symmetry is locally preserved. The Au(100) surface also bears a large cell reconstruction, resulting in a surface layer with an atomic density 25% larger than in the (100) bulk plane. The corresponding quasi-hexagonal surface layer displays an average interatomic spacing (~ 2.78 Å) very close to that of the (111)-herringbone reconstruction.^{97,98} Finally, the Au(110) surface displays the renowned (1 \times 2) missing row reconstruction, which exhibits local (111) microfacets. The missing row reconstruction may be lifted by the adsorption of large heteroaromatic molecules like CuPc, which locally deconstructs the surface beneath the molecular scaffold.⁹⁹



Table 2 MTPP (MPc) TSEAs (eV) of low-lying unoccupied SOs. Values of the highest TSEA are in bold^{ab}

	$\sigma_{a_{1g}}$	$\sigma_{b_{1g}}$	$\pi_{b_{2g}}$	π_{e_g}	$p_{mc}e_g$	$p_{mc}d_{1u}$
VTPP	0.66 [↓]	0.31 [↑]	0.01 [↓]	2.23 [↑]	1.81 [↓]	0.43 [↑]
VPc	1.18 [↓]	-0.95 [↑]	0.53 [↓]	2.94 [↑]	2.50 [↓]	0.99 [↑]
CrTPP	0.39 [↓]	0.32 [↑]	0.13 [↓]	-0.37 [↓]	1.76 [↓]	0.38 [↑]
CrPc	1.13 [↓]	-0.59 [↑]	0.87 [↓]	0.96 [↓]	2.47 [↓]	0.97 [↑]
MnTPP	-0.20 [↓]	-3.59 [↓]	-1.17 [↓]	-1.57 [↓]	1.33 [↓]	-0.09 [↑]
MnPc	2.16 [↓]	1.14 [↑]	2.25 [↓]	1.69 [↓]	2.84 [↓]	0.93 [↑]
FeTPP	—	0.41 [↑]	—	0.71 [↓]	1.85 [↓]	0.35 [↑]
FePc	—	0.30 [↑]	—	1.44 [↓]	2.61 [↓]	0.95 [↑]
CoTPP	2.13 [↓]	1.06 [↑]	—	—	1.52 [↑]	0.26 [↑]
CoPc	3.00 [↓]	1.45 [↑]	—	—	2.17 [↑]	0.90 [↑]
NiTPP	—	1.47 [↑]	—	—	1.53 [↑]	0.27 [↑]
NiPc	—	1.88 [↑]	—	—	2.19 [↑]	0.91 [↑]
CuTPP	—	1.35 [↓]	—	—	1.61 [↑]	0.33 [↑]
CuPc	—	1.78 [↓]	—	—	2.25 [↑]	0.94 [↑]
ZnTPP	—	—	—	—	1.64 [↑]	0.38 [↑]
ZnPc	—	—	—	—	2.26 [↑]	0.96 [↑]

^a HS MnTPP TSEA calculations have been run by adopting the B3LYP functional. ^b TSEA positive values correspond to the IE of the MTPP⁻ (MPc⁻) species; negative values indicate an unfavorable (costly) electron capture.

while those of the *ns* AOs centered on ${}^S_{B}CM_{100}^{nn}$ transform as the *irs* a and b (see Fig. S15 of the ESI[†]) of the C_2 point group,⁷² and simply correspond to the in-phase and out-of-phase linear combinations of the two *ns* AOs; (ii) SALCs of the *ns* AOs centered on ${}^S_{B}CM_{110}^{nn}$ and ${}^S_{H}CM_{110}^{nn}$ transform as the *irs* a and b (see above); (iii) SALCs of the *ns* AOs centered on ${}^S_{T}CM_{111}^{nn}$ are bases for the *irs* a, b, and e (see Fig. S16 of the ESI[†]); SALCs of the *ns* AOs centered on ${}^S_{H}CM_{111}^{nn}$ transform as the *irs* a and e (see Fig. S17 of the ESI[†]); SALCs of the *ns* AOs centered on ${}^S_{B}CM_{111}^{nn}$ are of symmetry a and b (see Fig. S15 of the ESI[†]).§§

3.3 MTPP and MPc chemisorbed on CM surfaces

The core of the present paper consists of providing a convincing molecular picture of the MTPP and MPc grafting to CM surfaces and then a local point of view of the substrate → adsorbate charge transfer, if present, by combining elementary symmetry and geometry arguments with readily accessible information about the unoccupied frontier electronic structure of the free adsorbates. Far from attempting a systematic analysis of the adsorption of all the MTPP and MPc herein considered on the different chemisorption sites present on CM(100), CM(110), and CM(111) surfaces, we will limit ourselves to focus on selected case studies for which (i) the adsorbate → substrate charge transfer is established and (ii) the structural perturbations undergone by the pmc upon adsorption are tiny.

3.3.1 VPc on Ag(111) and Au(111) surfaces. Despite lacking any literature data for the isolated VTPP, Eguchi *et al.* succeeded in characterizing VPc on Ag(111) by NEXAFS and

XMCD.¹⁰⁰ The authors emphasized a wispy Ag(111) → VPc charge transfer and a flat orientation on the substrate.¹⁰⁰

A few years ago, Mabrouk and Majewski¹⁰¹ theoretically investigated the stability and the electronic and magnetic properties of VPc grafted to Au(111) by exploiting VASP.¹⁰² The authors considered all the possible high-symmetry chemisorption sites reported in the bottom panel of Fig. 3 as well as different molecular orientations to conclude that the VPc chemisorption on Au(111) is weakly dependent on the adsorption site (the ΔE between the most (H_{fcc}) and least (T) stable sites amounts to 120 meV) and the substrate → adsorbate charge transfer is weak and quantifiable in 0.62 electrons.¹⁰¹ Even without supercell periodic calculations and STM measurements, let us see how VPc ADF results combined with symmetry and geometrical considerations may provide useful insights into the grafting of VPc to Ag(111).

VPc TSEAs of the V 3d-based SOs are all but one positive (see Table 2 and Fig. 2); moreover, both the $\pi_{e_g}^{\uparrow}$ and $p_{mc}e_g^{\downarrow}$ TSEAs are very high (2.94 and 2.50 eV, respectively). In addition, the experimentally revealed role played by the VPc σ_{e_g} -like $\sigma_{a_{1g}}$ SO in the substrate → adsorbate charge transfer¹⁰⁰ necessarily implies the interaction of the V 3d₂ AO with 5s ^SAg SALCs transforming as the *ir* a in the chemisorption site local symmetry. As such, it can be useful to remember that the $\sigma_{a_{1g}}^{\downarrow}$ TSEA (1.18 eV; see Table 2 and Fig. 2) is significantly smaller than the $\pi_{e_g}^{\uparrow}$ and $p_{mc}e_g^{\downarrow}$ TSEAs (see above). Schematic representations of the 5s ^SAg SALCs transforming as the *ir* a for the T, B, and H $\uparrow\uparrow$ sites are superimposed to the optimized $p_{mc}VPc$ placed at 2.8 Å_{||} above the bulk terminated Ag(111) in Fig. 4 (${}^S_{T}Ag_{111}^{nn}-V$, ${}^S_{B}Ag_{111}^{nn}-V$, and ${}^S_{H}Ag_{111}^{nn}-V$ internuclear distances are 2.80, 3.15, and 3.26 Å, respectively).

Elementary symmetry, geometrical, and overlap considerations allow the following statements: (i) the participation of the VPc $\pi_{e_g}^{\uparrow}$ SOs to the ${}^S_{T}Ag_{111}^{nn} \rightarrow VPc$ charge transfer is symmetry forbidden at the T site (see the 3D CPs displayed in Fig. 5); (ii) the contribution of the VPc $\pi_{e_g}^{\uparrow}$ SOs to the ${}^S_{T}Ag_{111}^{nn} \rightarrow VPc$ charge transfer should be very weak as the ${}^S_{T}Ag_{111}^{nn}-V$ internuclear distance is 4.02 Å; (iii) similarly weak should be the contribution of both the VPc $\pi_{e_g}^{\uparrow}$ and $p_{mc}e_g^{\downarrow}$ SOs (see Fig. 5) to the ${}^S_{H}Ag_{111}^{nn} \rightarrow VPc$ charge transfer (the local C_3 symmetry of the H site is incompatible with the adsorbate local C_4 symmetry); (iv) both the VPc 3d-based σ_{e_g} -like $\sigma_{a_{1g}}^{\downarrow}$ (see Fig. 4) and $\pi_{b_{2g}}$ -like $\pi_{e_g}^{\uparrow}$ (see Fig. 5) may effectively participate to the ${}^S_{B}Ag_{111}^{nn} \rightarrow VPc$ charge transfer; (v) the VPc $p_{mc}e_g^{\downarrow}$ (see Fig. 5) is well tailored to participate to the ${}^S_{B}Ag_{111}^{nn}/{}^S_{B}Ag_{111}^{nn} \rightarrow VPc$ charge transfer (with VPc sitting at the B site, placed at 2.8 Å above the bulk-terminated Ag(111), and oriented as in Fig. 5, the ${}^S_{B}Ag_{111}^{nn}-N^{Py}$ and ${}^S_{B}Ag_{111}^{nn}-N^{m}$ internuclear distances are almost identical (2.85 and 2.84 Å, respectively)). The VPc chemisorption B site, with V-N^{Py} bonds oriented along the [011] and [211] directions,

¶¶ Besides CM(111) T and B sites, two types of H sites characterize the surface: the H_{hcp} and the H_{fcc} in Fig. 3. If the attention is limited to the outermost CM(111) layer, H_{hcp} and H_{fcc} have the same environment.

||| Blades *et al.* have thoroughly investigated the electronic and molecular properties of VCu_x^+ , VAg_x^+ , and VAu_x^+ ($x = 3, 14$) clusters. The V-Ag bond distance varies between 2.76 Å (VAg_3^+) and 2.95 Å (VAg_{14}^+) with a mean value of 2.8 Å.¹⁰³

§§ The schematic representations of the CM(111) ${}^S_{T}CM^{nn}$ and ${}^S_{H}CM^{nn}$ *ns* AOs SALCs sketched in Fig. S16 and S17 (ESI[†]) do not consider the *ns* AOs normalization coefficients; the radii of the *ns* AOs localized on different ${}^S_{CM}^{nn}$ are then the same. The interested reader may refer to Chapter 7 of ref. 72.



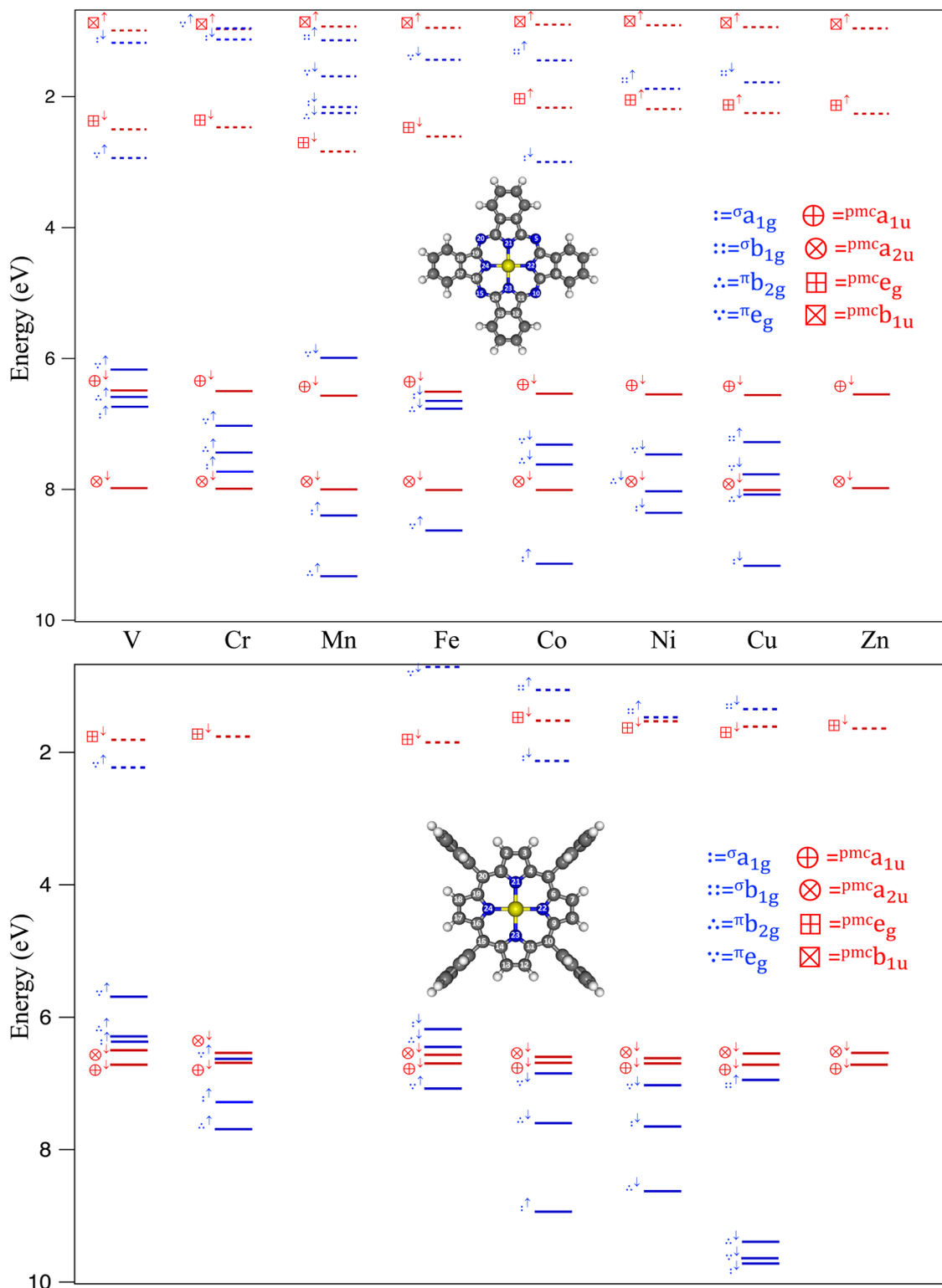


Fig. 2 TSEIs (solid lines) and TSEAs (dotted lines) of MPC and MTPP M 3d-based (blue lines) and $\text{pmc}\pi$ -based (red lines) spin orbitals. MnTPP TSEIs and TSEAs are not included in the figure, having been estimated by adopting the B3LYP^{73,74,85} functional rather than the BP86 one^{73,74} (see the main text).

seems then inescapable. This conclusion is only apparently in contrast with Mabrouk and Majewski¹⁰¹ results because one must consider on one side the lower nobleness of Ag compared

to Au (A_{Ag}^{χ} and A_{Au}^{χ} are 1.94 and 2.54 Pauling's units^{65,66}) and, on the other side, the weak dependence on the adsorption site of the VPC chemisorption on Au(111). Results so far presented are



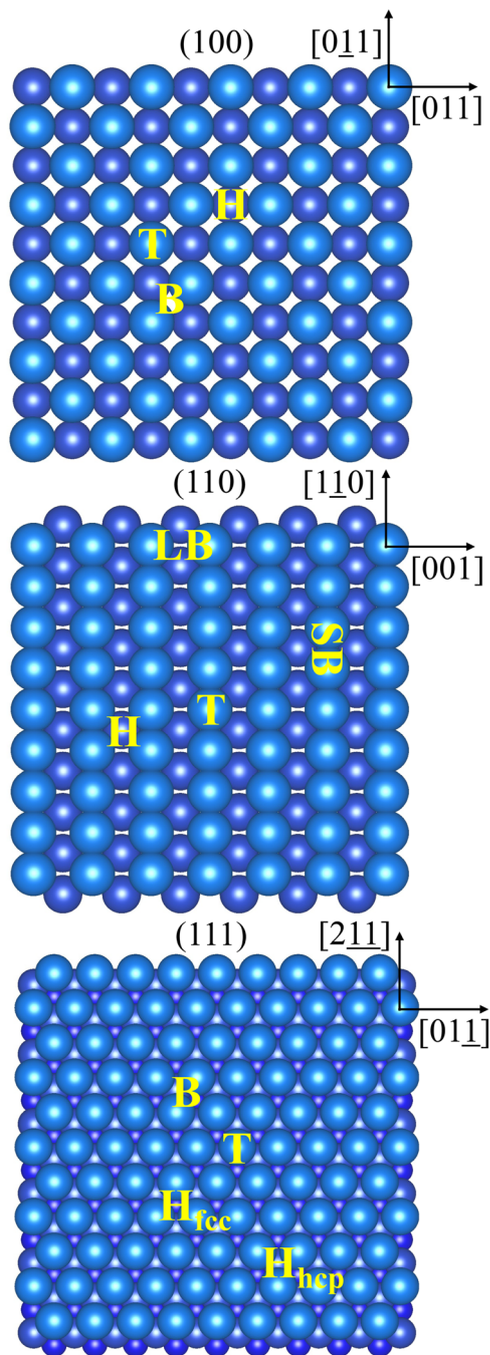


Fig. 3 Schematic representation of the high symmetry chemisorption sites (in yellow) on the CM(100), CM(110), and CM(111) bulk-terminated surfaces. T, B, and H stand for top, bridge, and hole, respectively. The topmost layer lies in the XY plane; moreover, the shades of blue correspond to different layers, with the most intense color corresponding to the deepest one.

useful to get information about the faint VPC species generated *in situ* on Ag(111),¹⁰⁰ and can be straightforwardly extended to other interfaces.

3.3.2 CoPc on CM surfaces. CoPc/CM(111) interfaces are particularly interesting because of the lack of a consensus about the nature and strength of the adsorbate–substrate

interaction. Li *et al.*,²⁵ based on DFT numerical experiments, proposed a H_{hcp} chemisorption site (see Fig. 3) for CoPc on Ag(111) and Au(111) and the occupation of a B site for CoPc on Cu(111). These differences were ascribed to an adsorbate/substrate interaction significantly different upon moving from Cu(111) to Ag(111) and Au(111). In contrast, the experimental and theoretical outcomes gathered by Baran *et al.*^{104,105} and Schmid *et al.*¹⁰⁶ for sub-ML of CoPc on Ag(111) prompted them to conclude that: (i) the CoPc chemisorption is accompanied by a spin state quenching from $S = \frac{1}{2}$ to $S = 0$; (ii) the Co 2p XPS features of the chemisorbed species shift towards lower binding energy consistently with an Ag(111) \rightarrow CoPc charge transfer; (iii) the CoPc σ_{e_g} -like $\sigma_{\text{a}_1^g}$ empty SO plays a leading role in determining the $\text{Co}^{\text{II}} \rightarrow \text{Co}^{\text{I}}$ reduction upon adsorption; (iv) the most stable chemisorption site corresponds to the T one with the Co^{II} species lying $2.90 \pm 0.05 \text{ \AA}$ above the Ag(111) surface.^{104,105} Interestingly, Zhao *et al.* reported similar evidence for CoPc on Au(111).⁵⁹

As already stressed, the inspection of Table 2 and Fig. 2 indicates that: (i) the TSEAs of M 3d-based SOs exceed those of the $\text{P}^{\text{m}^{\text{c}}}\pi^*$ ones only in the V and Co complexes; (ii) the CoPc σ_{e_g} -like $\sigma_{\text{a}_1^g}$ TSEA has the highest value (3.00 eV) among those reported in the table and displayed in the figure. Moreover, Fig. 4 reveals at glance that ${}^5\text{Ag}^{\text{nn}}$ allows the best overlap between the Co^{II} $3d_{z^2}$ AO and the ${}^5\text{Ag}$ SALCs transforming as the *ir a*. The huge CoPc EA also makes possible the Au(111) \rightarrow CoPc charge transfer (see above) despite the high ${}^{\text{Au}}\chi$ (2.54 Pauling's units^{65,66}), while the different behavior of CoPc⁵⁹ (T site, large charge transfer) and VPC¹⁰¹ (B site, weak charge transfer) on Au(111) may be reasonably traced back to (i) the fulfillment of CoPc $\pi_{\text{t}_{2g}^-}/\text{t}_{2g}^-$ -like SOs and (ii) the different TSEA of the σ_{e_g} -like $\sigma_{\text{a}_1^g}$ in VPC and CoPc (see Table 2 and Fig. 2).

Results about the CoPc/Au(111) interface may be now exploited to shed new light into the CoPc/Au(110)¹⁰⁷ and CoPc/Au(100)^{108,109} ones and for which the CoPc chemisorption site is still unknown. As such, it is experimentally established that the CoPc molecular plane lies parallel to the surface and results firmly anchored to the substrate both in CoPc/Au(110) and CoPc/Au(100);^{107,108} moreover, NEXAFS at the Co $L_{2,3}$ -edges and valence band photoemission suggested a reduction of the spin magnetic moment for CoPc/Au(110).¹⁰⁷ Photoemission studies of CoPc on Au(100) also demonstrate that the strong Au(100) \rightarrow CoPc charge transfer determines the $\text{Co}^{\text{II}} (\text{d}^7) \rightarrow \text{Co}^{\text{I}} (\text{d}^8)$ surface reduction, most likely involving the Co-based σ_{e_g} -like $\sigma_{\text{a}_1^g}$ SO, and then the quenching of the adsorbate magnetic moment ($S = 0$).¹⁰⁹ The giant CoPc first EA level (see Table 2 and Fig. 2) and the significant charge transfer from the Au(110) and Au(100) surfaces into the Co 3d-based SOs prompt us to indicate the T site as the most favorite not only for the CoPc chemisorption on Au(111) but also on Au(110) and Au(100). Concerning the azimuthal orientation of CoPc on Au(110), the inspection of Fig. 3 and 6 reveals that an effective substrate \rightarrow adsorbate charge transfer involving the lowest-lying $\text{P}^{\text{m}^{\text{c}}}\pi^*$ orbitals (the CoPc $\text{P}^{\text{m}^{\text{c}}}\pi^*$ SOs TSEA = 2.17 eV; see Table 2 and Fig. 2) should imply the out-of-phase linear combination



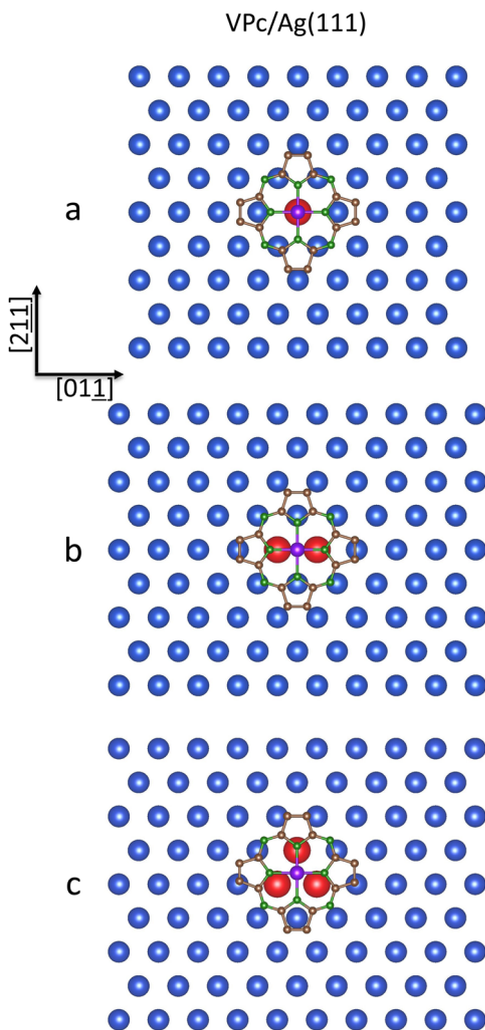


Fig. 4 Schematic representation of the ${}^5\text{Ag}_{111}^{\text{nn}}$ (a), ${}^5\text{Ag}_{111}^{\text{nn}}$ (b), and ${}^5\text{Ag}_{111}^{\text{nn}}$ (c) 5s AOs SALCs of symmetry a on the Ag(111) bulk-terminated surface superimposed to the optimized ${}^{\text{pmc}}\text{VPc}$ placed at 2.8 Å above the substrate and oriented with V–N^{Py} bonds aligned along the [211] and [011] directions. Only Ag(111) topmost layer atoms are displayed for clarity. Large red spheres represent ${}^5\text{Ag}^{\text{nn}}$ 5s AOs.

of the ${}^5\text{Au}_{110}^{\text{nn}}$ 6s AOs (see the upper panel of Fig. 6) and the CoPc ${}^{\text{pmc}}\text{e}_g^\uparrow$ SO having a node localized on the N^{Py} atoms aligned to the [001] direction (see the lower panel of Fig. 6). Incidentally, with the planar CoPc placed at 3 Å above the bulk-terminated Au(110) surface, the Co– ${}^5\text{Au}_{110}^{\text{nn}}$ and N^{Py}– ${}^5\text{Au}_{110}^{\text{nn}}$ internuclear distances are 4.16 and 3.15 Å, respectively.

For the Au(100) surface, the experimental evidence reported by Lindner *et al.*,¹⁰⁹ combined with the CoPc and CM(100) local fourfold symmetry, is also consistent with CoPc occupying a T site and the Co–N^{Py} bonds oriented along the [011] and [011] directions (see Fig. 3 and 7). As such, it is noteworthy that the chemisorption T site and the proposed orientation are best suited to favor the charge transfer from the ${}^5\text{Au}_{100}^{\text{nn}}$ 6s AO into the σ_{e_g} -like $\sigma_{\text{a}_g}^\uparrow$ SO through a direct Co– ${}^5\text{Au}_{100}^{\text{nn}}$ through-space interaction, which determines the experimentally detected Co^{II} (d⁷) → Co^I (d⁸) surface reduction.¹⁰⁹ Incidentally, the CoPc

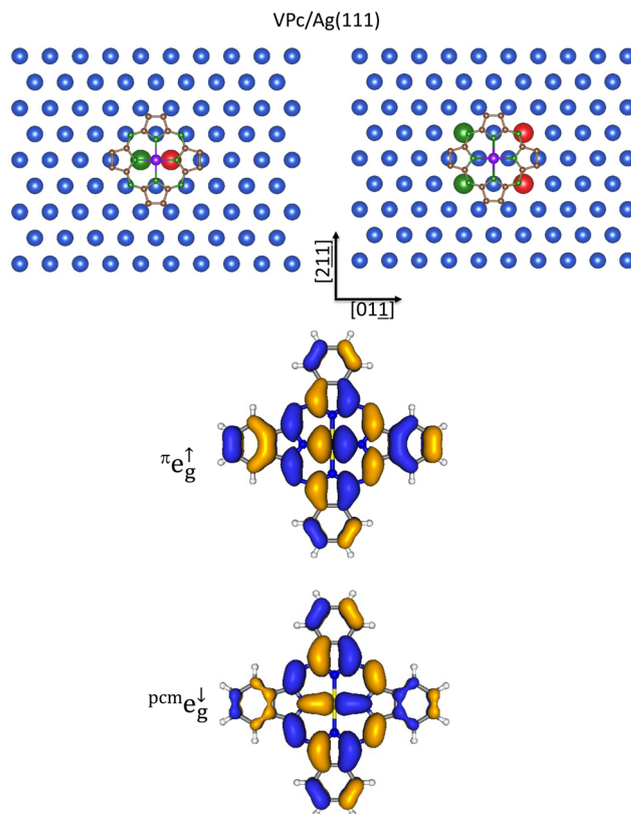


Fig. 5 Schematic representation of the ${}^5\text{Ag}_{111}^{\text{nn}}$ (upper left panel) and ${}^5\text{Ag}_{111}^{\text{nn}}$ (upper right panel) 5s AOs SALC of symmetry b on the Ag(111) bulk-terminated surface superimposed to the optimized ${}^{\text{pmc}}\text{VPc}$ placed at 2.8 Å above the substrate and oriented with V–N^{Py} bonds aligned along the [211] and [011] directions. Only Ag(111) topmost layer atoms are displayed for clarity. Large spheres represent the ${}^5\text{Ag}^{\text{nn}}$ and ${}^5\text{Ag}^{\text{nn}}$ 5s AOs, while different colors account for their different phases. 3D CPs of one component of the V^{II} $\pi_{t_{2g}}$ -like $\pi_{\text{e}_g}^\uparrow$ SOs (middle panel) and one component of the ${}^{\text{pmc}}\text{e}_g^\downarrow$ π^* SOs (lower panel). Displayed isosurfaces correspond to $\pm 0.02 \text{ e}^2 \text{ \AA}^{-3/2}$ values.

chemisorption at the Ag(100) H site, as proposed by Mugarza *et al.*,¹¹⁰ is contradictory with the simultaneously reported charge transfer of one electron into the molecule; as such, it can be discarded.

The overlap between the SALCs of Au 6s AOs localized on the ${}^5\text{Au}_{100}^{\text{nn}}$ and ${}^5\text{Au}_{100}^{\text{nn}}$, transforming as the *ir e* in the local C_4 symmetry, and the CoPc ${}^{\text{pmc}}\text{e}_g^\uparrow$ SO (see Fig. 7), provides a theoretical rationale for the tight anchoring of the adsorbate to the substrate. By the way, within the assumption of the CoPc molecular plane at 3 Å from the surface and the Co–N^{Py} bonds oriented along the [011] and [011] directions (see Fig. 7), the ${}^5\text{Au}_{100}^{\text{nn}}$ –N^{Py} and ${}^5\text{Au}_{100}^{\text{nn}}$ –N^m internuclear distances are 3.15 and 3.08 Å, respectively.

3.3.3 FePc on (100) and (110) CM surfaces. A planar arrangement of FePc was reported on both Au(110)¹⁰⁷ and Au(100) surfaces.¹⁰⁸ Moreover, Betti *et al.*¹⁰⁷ emphasized the quite strong bonding determined by the interaction of the Fe^{II} 3d-based AOs with the Au(110) surface.

The FePc ${}^3\text{A}_{2g}$ IS GS determined by the $[\sigma_{\text{a}_g}^\uparrow \pi_{\text{b}_g}^\downarrow \pi_{\text{e}_g}^\uparrow]$ configuration (see Table S14 of the ESI[†]) would prevent the



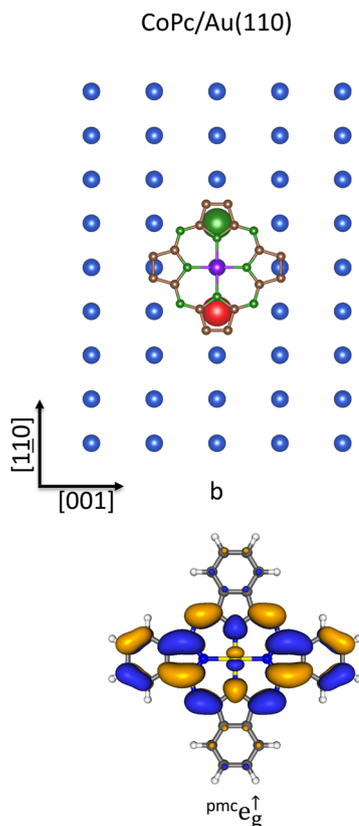


Fig. 6 Schematic representation of the ${}^5\text{Au}_{110}^{\text{nnn}}$ 6s AOs SALC of symmetry b on the Au(110) bulk-terminated surfaces superimposed to the optimized ${}^{\text{pmc}}\text{CoPc}$ placed at 3.0 Å above the substrate and oriented with Co–N^{Py} bonds aligned along the [110] and [001] directions. Only Au(110) topmost layer atoms are displayed for clarity. Large spheres represent ${}^5\text{Au}_{110}^{\text{nnn}}$ 6s AOs, while different colors account for their different phases (upper panel). 3D CP of one component of the $e_g^{\text{pmc}}e_g^{\uparrow}$ SOs (lower panel). Displayed isosurfaces correspond to $\pm 0.02 e^{\frac{1}{2}} \text{Å}^{-3/2}$ values.

CoPc/Au(100)

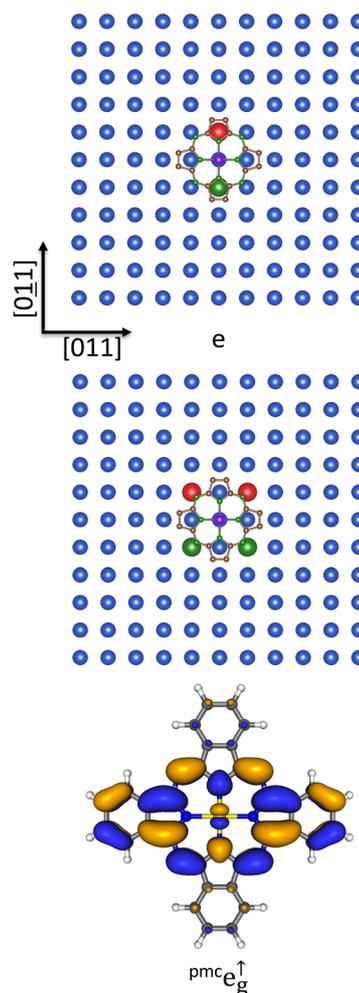


Fig. 7 Schematic representation of one component of the ${}^5\text{Au}_{100}^{\text{nnn}}$ (upper panel), ${}^5\text{Au}_{100}^{\text{nnn}}$ (middle panel) 6s AOs SALC of symmetry e on the Au(100) bulk-terminated surface superimposed to the optimized ${}^{\text{pmc}}\text{CoPc}$ placed at 3.0 Å above the substrate and oriented with Co–N^{Py} bonds aligned to the [011] and [011] directions. Only Au(100) topmost layer atoms are displayed for clarity. Large spheres represent ${}^5\text{Au}_{100}^{\text{nnn}}$ and ${}^5\text{Au}_{100}^{\text{nnn}}$ 6s AOs, while different colors account for their different phases. 3D CP of one component of the $e_g^{\text{pmc}}e_g^{\uparrow}$ SOs (lower panel). Displayed isosurfaces correspond to $\pm 0.02 e^{\frac{1}{2}} \text{Å}^{-3/2}$ values.

chemisorption site T because the antibonding component of the interaction between the completely occupied FePc σ_{e_g} -like $\sigma_{a_{1g}^{\uparrow\downarrow}}$ MO (the Fe^{II} 3d_{z²-based MO) and the ${}^5\text{Au}_{100}^{\text{nnn}}$ 6s AO would result partially occupied. Nevertheless, the minute ΔE between the FePc ${}^3E_g^{(1)}$ excited state generated by the $[\sigma_{a_{1g}^{\uparrow\downarrow}}\pi_{b_{2g}^{\uparrow\downarrow}}\pi_{e_g^{\uparrow\downarrow}}\pi_{e_g^{\uparrow\downarrow}}]$ configuration and the ${}^3A_{2g}$ GS (47 meV; see Table S14 of the ESI†) must be kept in mind because tiny ligand field perturbations, for instance, those induced by chemisorption, could generate different occupation numbers of the Fe^{II} 3d-based SOs. As such, the TSEAs of the ${}^3E_g^{(1)}\pi_{t_{2g}}$ -like ($\pi_{\perp}e_g^{\downarrow}$) 12e_g LUMOs (3.04 eV) and σ_{e_g} -like ($\sigma_{a_{1g}^{\downarrow}}$) 21a_{1g} LUMO+2 (2.70 eV), much higher than those reported in Table 2 for the low-lying empty SOs of the isolated molecule in its ${}^3A_{2g}$ GS, are consistent with a higher FePc electron-withdrawing capability in the ${}^3E_g^{(1)}$ state than in the ${}^3A_{2g}$ one. The $\pi_{\perp}e_g^{\downarrow}$ and $\sigma_{a_{1g}^{\downarrow}}$ SOs bareness and their high TSEAs make them very well suited to actively participate to the substrate → adsorbate charge transfer and then to the anchoring of FePc to both Au(110) and Au(100).}

Besides the abnormal gold nobleness^{65,66} and the ${}^3E_g^{(1)}\pi_{\perp}e_g^{\downarrow}$ SOs TSEA (3.04 eV), higher than the ${}^3E_g^{(1)}\sigma_{a_{1g}^{\downarrow}}$ one (2.70 eV), we may further account for the symmetry and geometry

constraints to determine the FePc chemisorption site on Au(110). Starting with symmetry considerations: (i) the participation of the Fe^{II} $\pi_{t_{2g}}$ -like $\pi_{\perp}e_g^{\downarrow}$ SOs to the substrate → adsorbate charge transfer is symmetry forbidden for FePc chemisorbed at a T site of the Au(110) and Au(100) surfaces but symmetry allowed for FePc chemisorbed at the SB site in the former case and the B one in the latter (see Fig. 8);*** (ii) the participation of the Fe^{II} σ_{e_g} -like $\sigma_{a_{1g}^{\downarrow}}$ SO to the anchoring of FePc to both Au(110) and Au(100) is symmetry allowed for FePc chemisorbed

*** The Au(110) LB chemisorption site has not being herein considered because, with FePc positioned 2.85 Å above the bulk-terminated surface,¹¹¹ the Fe–_{LB}Agⁿⁿ internuclear distance amounts to 3.504 Å.



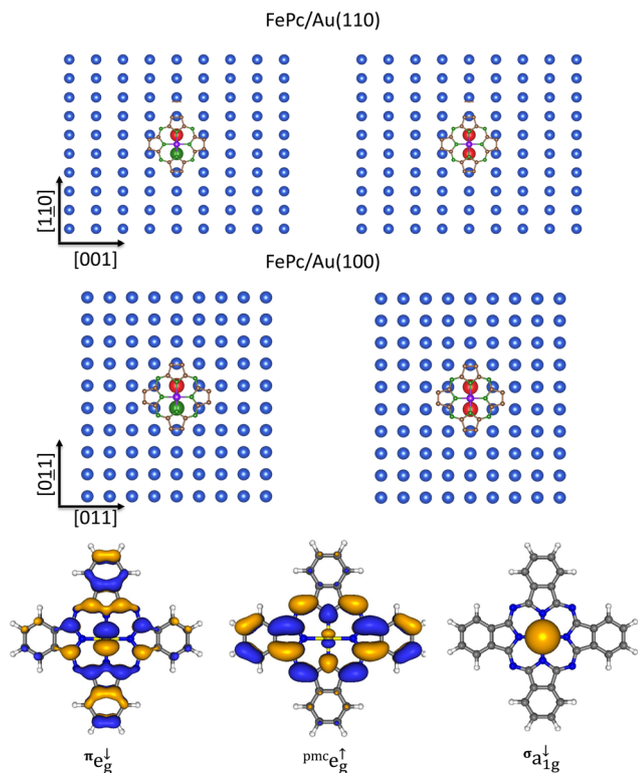


Fig. 8 Schematic representation of the $\text{S}_{\text{SB}}\text{Au}_{110}^{\text{nn}}$ 6s AOs SALCs of symmetry b (upper, left panel) and a (upper, right panel) on the Au(110) bulk-terminated surface superimposed to the optimized pmc^{FePc} placed at 2.85 Å above the substrate and oriented with Fe–N^{Py} bonds aligned to the [110] and [001] directions. Only Au(110) topmost layer atoms are displayed for clarity. Large spheres represent $\text{S}_{\text{SB}}\text{Au}_{110}^{\text{nn}}$ 6s AOs, while different colors account for their different phases. Schematic representation of the $\text{S}_{\text{B}}\text{Au}_{100}^{\text{nn}}$ 6s AOs SALCs of symmetry b (middle, left panel) and a (middle, right panel) on the Au(100) bulk-terminated surface superimposed to the optimized pmc^{FePc} placed at 2.85 Å above the substrate and oriented with Fe–N^{Py} bonds aligned to the [110] and [011] directions. Only Au(100) topmost layer atoms are displayed for clarity. Large spheres represent $\text{S}_{\text{B}}\text{Au}_{100}^{\text{nn}}$ 6s AOs, while different colors account for their different phases. 3D CPs of one component of the partially occupied FePc ${}^3\text{E}_g^{(1)}$ π^* SOs (lower, left panel), unoccupied FePc ${}^3\text{E}_g^{(1)}$ pmc^{FePc} π^* SOs (lower, middle panel), and unoccupied FePc ${}^3\text{E}_g^{(1)}$ $\sigma_{\text{a}1g}$ SO (lower, right panel). Displayed isosurfaces correspond to $\pm 0.02 \text{ e}^2 \text{ \AA}^{-3/2}$ values.

at the SB site in the former case and the B one in the latter (see Fig. 8);*** (iii) the FePc pmc^{FePc} π^* SOs (TSEA = 2.03 eV) may actively participate to the anchoring of FePc to both Au(110) and Au(100) when chemisorbed at the SB site in the former case and the B one in the latter. With specific reference to the third point, it has to be noted that the substrate → adsorbate charge transfer will involve different pmc atoms upon moving from the FePc/Au(110) interface to the FePc/Au(100) one (see Fig. 9).

From a geometry perspective, starting with FePc positioned 2.85 Å¹¹¹ above the bulk terminated Au(110) at the SB site*** (see Fig. 8 and 9), the $\text{S}_{\text{SB}}\text{Au}_{110}^{\text{nn}}\text{–Fe}$, $\text{S}_{\text{SB}}\text{Au}_{110}^{\text{nn}}\text{–N}^{\text{Py}}$, and $\text{S}_{\text{SB}}\text{Au}_{110}^{\text{nn}}\text{–C}^{\beta}$ internuclear distances are 3.19, 2.89, and 2.94 Å, respectively; moreover, with FePc positioned 2.85 Å¹¹¹ above the bulk terminated Au(100) at the B site (see Fig. 8 and 9), the $\text{S}_{\text{B}}\text{Au}_{100}^{\text{nn}}\text{–Fe}$, $\text{S}_{\text{B}}\text{Au}_{100}^{\text{nn}}\text{–N}^{\text{Py}}$, $\text{S}_{\text{B}}\text{Au}_{100}^{\text{nn}}\text{–C}^{\alpha}$, and $\text{S}_{\text{B}}\text{Au}_{100}^{\text{nn}}\text{–N}^m$ internuclear distances are 3.19, 2.89, 2.87, and 3.04 Å, respectively.

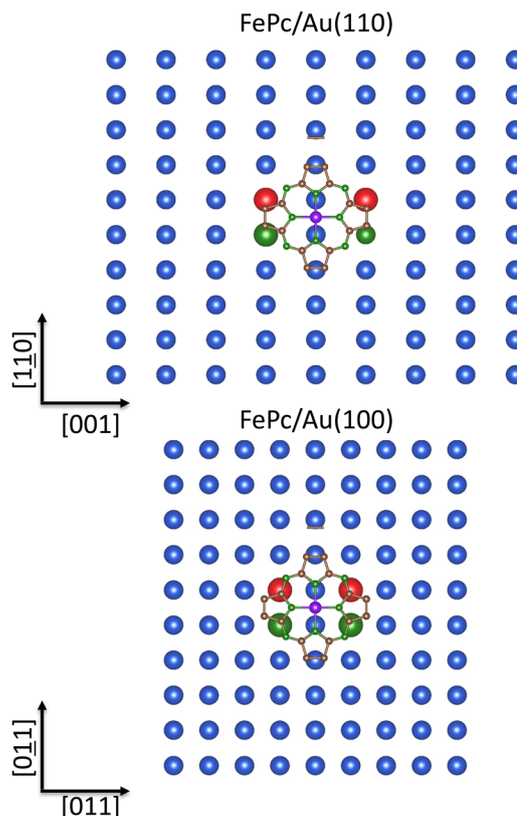


Fig. 9 Schematic representation of the $\text{S}_{\text{SB}}\text{Au}_{110}^{\text{nn}}$ and $\text{S}_{\text{B}}\text{Au}_{100}^{\text{nn}}$ 6s AOs SALCs of symmetry b on the Au(110) (upper panel) and Au(100) (lower panel) bulk-terminated surfaces superimposed to the optimized pmc^{FePc} placed at 2.85 Å above the substrate and oriented with Fe–N^{Py} bonds aligned to the [110] and [001] directions. Only Au(110) and Au(100) topmost layer atoms are displayed for clarity. Large spheres represent $\text{S}_{\text{SB}}\text{Au}_{110}^{\text{nn}}$ and $\text{S}_{\text{B}}\text{Au}_{100}^{\text{nn}}$ 6s AOs, while different colors account for their different phases.

SALCs displayed in Fig. 9 and the 3D CP of the FePc pmc^{FePc} π^* SOs (not herein reported because indistinguishable from the 3D CP of the CoPc pmc^{CoPc} π^* SOs reported in Fig. 7) allow us to assess that, for FePc at 2.85 Å¹¹¹ from the bulk terminated Au(110) and positioned at the SB site, the substrate → adsorbate charge transfer involving the pmc^{FePc} π^* SOs will mainly concern the N^{Py} and C^β 2p_z AOs. For FePc at the same distance from the bulk terminated Au(100) and positioned at the B site, the substrate → adsorbate charge transfer involving the pmc^{FePc} π^* SOs will mainly concern the N^{Py}, N^m, and C^α 2p_z AOs. The most stable chemisorption site of FePc on Au(110) and Au(100) should be then the SB and B one, respectively.

Strictly related to the just considered FePc/Au(110) interface, is the FePc/Ag(110) system; an interface able to catalyze the four-electron oxygen reduction reaction to form H₂O from O₂^{46,112} and closely related to the oxygen-binding active center of hemoglobin. Some of us have thoroughly investigated the adsorbate–substrate interactions taking place at the FePc/Ag(110) interface through the combined use of high-resolution STM measurements and DFT supercell periodic calculations including semiempirical dispersion interactions.¹¹¹ Experimental measurements unveiled that FePc molecules result



parallel to the Ag(110) surface and arranged in rows running along the [001] direction (see the central panel of Fig. 3); moreover, two closely related phases (c(10 × 4) and p(10 × 4) superstructures) observed after depositing FePc at RT and persisting after extensive annealing at 473 K, were revealed at the sub-ML regime.

DFT calculations, indicated that: (i) chemisorption sites of the c(10 × 4) and p(10 × 4) superstructures are different, they correspond to T and SB sites, respectively, with a height of Fe^{II} with respect to the Ag(110) topmost layer of 2.85 Å in the c(10 × 4) phase and 2.54 Å in the p(10 × 4) one; (ii) in the c(10 × 4) phase, all the FePc molecules are oriented with their C₂' axes (those aligned with Fe–N^{Pv} bonds) forming a 45° angle with the [110] direction of the Ag substrate; (iii) in the p(10 × 4) phase, FePc molecules are arranged in an alternated sequence of two linear arrays, both aligned along the [001] direction of the Ag substrate and differing for the angle formed by their C₂' axes and the [110] direction ((30 ± 2)° and (−30 ± 2)°, respectively).¹¹¹ Finally, even though the FePc IS state was mentioned, the authors provided no information about its symmetry.¹¹¹ All this evidence may be straightforwardly rationalized by referring, besides the symmetry and the geometry arguments invoked to assess the chemisorption site of FePc on Au(110), to the different ^{Au}χ and ^{Ag}χ.^{65,66} The ^{Ag}χ value, significantly lower than the ^{Au}χ one, and the high TSEA (2.70 eV) of the ³E_g⁽¹⁾ σ_{e_g-like σ_{a_{1g} SO, concur to favor a direct Fe–Ag bonding interaction between the Fe^{II} σ_{e_g-like σ_{a_{1g} and the ^SAg₁₁₀ⁿⁿ 5s AO and then a T chemisorption site. The FePc sitting at the T site does not prevent the possibility of an alternative chemisorption site, the SB one, with both the Fe^{II} π_{t_{2g}}-like π₁e_g¹ and σ_{e_g}-like σ_{a_{1g}} SOs contributing to the adsorbate–substrate grafting. We remark that the Fe atom optimized eight of 2.54 Å characterizing the p(10 × 4) phase¹¹¹ implies a ^{SB}Ag₁₁₀ⁿⁿ–Fe internuclear distance of 2.92 Å, very similar to the ^SAg₁₁₀ⁿⁿ–Fe one (2.85 Å). Finally, the results herein reported for the FePc/Ag(110) interface can be straightforwardly transferred to the FePc/Cu(110) one.¹¹³}}}}

3.3.4 CoTPP and NiTPP on CM surfaces. The choice of CoTPP/CM and NiTPP/CM interfaces as two further case studies is determined by the evidence that, despite the huge amount of experimental and theoretical data available in the literature for CoTPP/Cu,^{76,114–117} CoTPP/Ag,^{76,114,117–122} CoTPP/Au,^{76,115,118,120,123–125} NiTPP/Cu,^{50,51,60,61,117} NiTPP/Ag,¹¹⁷ and NiTPP/Au,^{52,123} controversies about the actual MTPP adsorption geometry are still present. Starting from the Co^{II} tetrapyrroles, the following issues must be considered: (i) both Cu and Ag surfaces act as electron donors able to induce a Co^{II} (d⁷) → Co^I (d⁸) surface reduction,^{114,115,118–121} while the Au substrate does not affect the Co oxidation state;^{118,123,124} (ii) the Co^{II} (d⁷) → Co^I (d⁸) surface reduction,^{114,115,118–121} systematically implies the fulfillment of the σ_{e_g}-like σ_{a_{1g}} MO almost completely localized on the Co^{II} 3d_{z²} AO and its concomitant loss of magnetization;¹¹⁴ (iii) the CoTPP first affinity level corresponds to the Co σ_{e_g}-like 3d_{z²}-based 25a_{1g}¹ SO (see Table 2 and Fig. 2); (iv) tiny differences in the chemisorption energy (~85 meV) characterize different chemisorption sites on Ag(111).¹¹⁹

Experimentally, Schwarz *et al.*¹²⁶ ultimately stated the SB adsorption geometry and the adsorption height (2.25 ± 0.04 Å) of CoP on Cu(111) by combining STM, high-resolution XPS, XSW measurements, and DFT calculations. However, when TPPs are concerned, intermolecular interactions allowed by the rotational degrees of freedom of Ph fragment decorating the pmc may overcome those of M and pmc with the substrate. Some of us revealed by PED the coexistence of H_{hpc} and H_{fcc} sites for a saturated CoTPP layer on Ag(111), as driven by the Ph intercalation among adjacent molecules (the so-called T-type interaction).¹²⁷

At first sight, the experimental and theoretical results collected by Schwarz *et al.*¹²⁶ are liable to affect the proposed approach; nevertheless, a thorough analysis of data reported in Table 2 and displayed in Fig. 2 demonstrates the opposite. The CoTPP highest TSEA (2.13 eV) is the largest among MTPP complexes (alike to CoPc among MPCs); however, it is noteworthy that the CoPc first affinity level (3.00 eV) significantly exceeds the CoTPP one. In addition, the TSEA of the CoTPP σ_{e_g}-like 3d_{x²-y²}-based 13b_{1g}¹ SO (1.06 eV), is the highest among lighter MTPP (ΔTSEA between CoTPP σ_{e_g}-like SOs = 1.07 eV). Incidentally, the ΔTSEA between the CoP σ_{e_g}-like SOs amounts to 1.00 eV.††† Now, the participation of the σ_{e_g}-like 3d_{x²-y²}-based SO to the substrate → adsorbate charge transfer is prevented by symmetry when the adsorbate is chemisorbed at the T site, while the contribution of both σ_{e_g}-like MOs is symmetry allowed when the adsorbate occupies a B site (SB for the CM(110) surface). This is ultimately stated both experimentally and theoretically¹²⁶ for the least electronegative CM;^{65,66} thus, it sounds reasonable that the chemisorption site of CoTPP on CM surfaces is, in the presence of a relevant substrate → adsorbate charge transfer involving the Co^{II} (d⁷) → Co^I (d⁸) reduction, the B one (SB for the CM(110)).

We recently addressed the adsorption configuration of NiTPP on Cu(100) by symmetry arguments as follows. The isolated NiTPP has a closed-shell nature and a ¹A_{1g} GS determined by the π_{t_{2g}}⁶-σ_{e_g}²-like configuration; moreover, the empty Ni^{II} 3d-based σ_{e_g}-like MO corresponds to the GS 12b_{1g} LUMO (see Fig. 10) whose TSEA (1.47 eV, see Table 2 and Fig. 2) is closely spaced to that of the ^{pmc}π* 13e_g MOs (1.53 eV, see Table 2 and Fig. 2) and significantly higher than the TSEA of 9b_{1u} ^{pmc}π* MO (the forth low-lying NiTPP empty orbital; see Table 2, Fig. 2, and Fig. 10).

Experimental and theoretical outcomes pertinent to the NiTPP/Cu(100) interface revealed that: (i) both the substrate and the adsorbate have a local C₄ symmetry;¹²⁸ (ii) the huge Cu(100) → NiTPP charge transfer taking place at the interface determines the partial occupation of all four low-lying unoccupied NiTPP MOs (the σ_{e_g}-like 12b_{1g}, as well as the 13e_g and 9b_{1u} ^{pmc}π* MOs; see Fig. 10);¹²⁸ (iii) the pmc of the chemisorbed NiTPP has a flat geometry,¹²⁸ it lies at 1.93 Å from the topmost layer of the surface,⁶⁰ and is characterized by the presence of the highly reactive 3d⁹ Ni^I species,⁶¹ which excludes adsorption at a T site by symmetry constraints.⁶⁰ With specific

††† The TSEA of the CoP 3d_{z²}-based σ_{a_{1g}}¹ and 3d_{x²-y²}-based σ_{b_{1g}}¹ SOs are 1.73 eV and 0.73 eV, respectively.



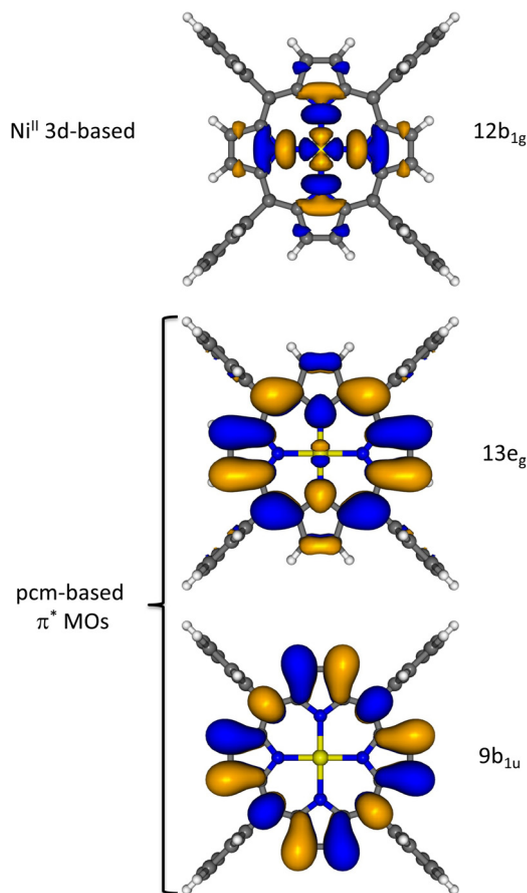


Fig. 10 3D CPs of low-lying NiTPP empty MOs (only one component of the $13e_g$ orbital is reported); displayed isosurfaces correspond to $\pm 0.02 e^3 \text{ \AA}^{-3/2}$ values.

reference to the last point, the $\text{Ni}^{\text{II}} (3d^8) \rightarrow \text{Ni}^{\text{I}} (3d^9)$ surface reduction necessarily involves the only Ni^{II} 3d empty AO ($3d_{x^2-y^2}$), basis for the $i_r b$ in the local C_4 symmetry, whose overlap with the $7\text{Cu}_{100}^{\text{II}}$ 4s AO (basis for the $i_r a$ in the same fourfold symmetry) is identically zero. Thus, the substrate \rightarrow adsorbed charge transfer rules the NiTPP adsorption on a specific chemisorption site; *i.e.*, the fourfold hollow site characterizing the Cu(100) surface (see the top panel of Fig. 3).⁶⁰ Incidentally, STM and STS results recently collected by Okuyama *et al.* indicate that CuPc (a single vacancy in the $\text{Cu}^{\text{II}} 3d_{x^2-y^2}$) on Cu(100) maintains its flat geometry and occupies a C_4 H site.¹²⁹

As far as the NiTPP/Cu(110) interface is concerned, MOT¹³⁰ results indicated a relevant adsorbate–substrate interaction characterized by a significant Cu(110) \rightarrow NiTPP charge transfer and involving both the $13e_g$ and $9b_{1u}$ $^{\text{pmc}}\pi^*$ MOs of the isolated NiTPP. Zamborlini¹¹⁷ also emphasized that, similarly to NiTPP/Cu(100), no contribution from the σ_{e_g} -like $12b_{1g}$ MO was revealed in the NiTPP/Cu(110) μ -ARPES maps. Incidentally, the absence of such a contribution in the NiTPP/Cu(100) μ -ARPES maps, successively revealed by NEXAFS measurements at the Ni L_3 -edge,⁶¹ was tentatively ascribed to the fact that the σ_{e_g} -like $12b_{1g}$ MO momentum map shows only four narrow

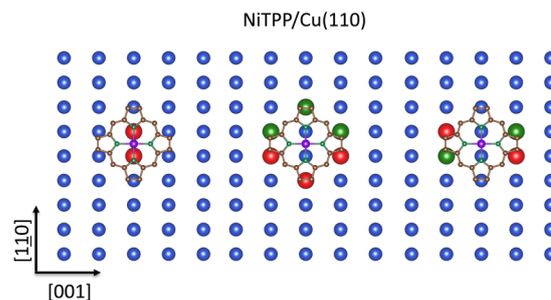


Fig. 11 Schematic representation of the $5_{\text{SB}}\text{Cu}_{110}^{\text{II}}$ 4s AOs SALC of symmetry a in the local C_2 symmetry of the Cu(110) bulk-terminate surface (left panel), $5_{\text{SB}}\text{Cu}_{110}^{\text{II}}$ 4s AOs SALC of symmetry b in the local C_2 symmetry of the Cu(110) bulk-terminate surface (central panel), $5_{\text{SB}}\text{Cu}_{110}^{\text{II}}$ 4s AOs SALC of symmetry a in the local C_2 symmetry of the Cu(110) bulk-terminate surface superimposed to the optimized $^{\text{pmc}}\text{NiTPP}$ placed at 2 Å above the substrate and oriented with Ni–N^{Py} bonds aligned along the [110] and [001] directions. Only Cu(110) topmost layer atoms are displayed for clarity. Large spheres represent $5_{\text{SB}}\text{Cu}_{110}^{\text{II}}$ and $5_{\text{SB}}\text{Cu}_{110}^{\text{I}}$ 4s AOs, while different colors account for their different phases.

lobes, presumably located outside the experimentally probed k -space range.¹¹⁷

To the best of the authors' knowledge, NEXAFS measurements at the Ni L_3 -edge of the NiTPP/Cu(110) interface able to unequivocally determine the Ni oxidation state are not present in the literature. Nevertheless, it has been claimed that the Cu(110) \rightarrow NiTPP charge transfer taking place at the NiTPP/Cu(110) interface is similar to that characterizing the NiTPP/Cu(100).¹¹⁷ Thus the frontier electronic structure of the free NiTPP prevents a T chemisorption site and favors a SB site in agreement with periodic DFT calculations.¹¹⁷ In fact, the empty Ni 3d-based MO reminiscent of the σ_{e_g} -like $12b_{1g}$ orbital may interact, in the local C_2 symmetry, with the $5_{\text{SB}}\text{Cu}_{110}^{\text{II}}$ 4s AOs SALC of symmetry a (see the left panel of Fig. 11). The inspection of Fig. 11, where the SALCs of symmetry b and a of the $5_{\text{SB}}\text{Cu}_{110}^{\text{II}}$ 4s AOs are displayed in the central and right panels, makes evident that they are very well suited for transferring electronic charge into the $^{\text{pmc}}\pi^*$ MOs reminiscent of the free NiTPP $13e_g$ and $9b_{1u}$ orbitals, respectively.

The crystal structure shared by Cu and Ag, their very similar χ values (1.90 and 1.94 Pauling's units, respectively),^{65,66} and the unoccupied frontier electronic structure of NiTPP would prompt to foresee the same SB chemisorption site for NiTPP/Cu(110) and NiTPP/Ag(110) interfaces, eventually imaging a weaker adsorbate–substrate interaction in the latter case determined by the larger size of Ag atoms ($r_{\text{Ag}} = 1.60 \text{ \AA}$) compared to that of the Cu ones ($r_{\text{Cu}} = 1.35 \text{ \AA}$).⁶⁵

3.3.5 ZnTPP on CM surfaces. The last case study herein considered concerns the Zn^{II} tetrapyrroles/CM interfaces. The $3d^{10}$ closed shell configuration of the Zn^{II} ion carries with it both formal and substantial consequences. As far as the former is concerned, Zn^{II} tetrapyrroles are better described as complexes of a post-transition element (see IUPAC gold book definition),¹³¹ on the other hand, the Zn^{II} configuration determines, even though indirectly, the chemisorption site of these



complexes on CM surfaces. More specifically, the substrate \rightarrow adsorbate charge transfer, if present, can only involve the low-lying $^{pmc}\pi^*$ MOs as the interaction between the occupied πt_{2g} -like/ σe_g -like Zn^{II} 3d-based MOs and the SCM ns SALCs of suitable symmetry is repulsive in character. A sizable charge transfer and a tough substrate-adsorbate grafting of Zn^{II} tetrapyrroles to CM surfaces will then imply the minimization (maximization) of the interaction of the Zn^{II} 3d-based (low-lying $^{pmc}\pi^*$) MOs with SCM ns SALCs of suitable symmetry. The chemisorption T site on any CM surface will be ruled out because of the repulsive interaction between the Zn^{II} σe_g -like σa_{1g} MO pointing directly toward the ns AO of symmetry a localized on the $^SCM^{nn}$. In addition, the inspection of the top panel of Fig. 3 reveals at glance that the CM(100) H site is also prevented because of the possible repulsive interaction between the Zn^{II} σe_g -like σb_{1g} MO or, depending on the alignment of the chemisorbed pmc , πt_{2g} -like πb_{2g} MO and the CM ns SALC of symmetry b localized on $^SCM^{nn}$ (see Fig. S14 of the ESI†). The only possible chemisorption site of Zn^{II} tetrapyrroles on CM(100) should be then the B one. Moving to CM(111) substrates, the T site has already been excluded; moreover, the incompatibility of the local C_3 symmetry of the H site with the adsorbate local C_4 symmetry prevents the occupation of this site too. Again, the only possible chemisorption site of Zn^{II} tetrapyrroles on CM(111) should be the B one with $Zn-N^{Py}$ bonds aligned with the $[01\bar{1}]$ and $[21\bar{1}]$ directions.

The last substrate to be considered is the CM(110) one. Again, the inspection of Fig. 3 (middle panel) is enlightening. T and SB sites must be rejected because they are unfitted to minimize the repulsive interaction between Zn^{II} πt_{2g} -like/ σe_g -like completely occupied MOs and the underneath SCM atoms. As far as the CM(110) H and LB sites are concerned, a schematic representation of the Ag(110) with the optimized $^{pmc}ZnTPP$ placed at 2.8 \AA^{132} above the substrate and oriented with $Zn-N^{Py}$ bonds aligned along the $[110]$ and $[001]$ directions is reported in Fig. 12. The inspection of the figure reveals at once that the overlap between the $^S_HAg_{110}^{nn}$ 5s SALC of symmetry b and the $13e_g$ $^{pmc}\pi^*$ MOs (see Fig. 10; NiTPP and ZnTPP 3D plots of the $13e_g$ MOs are indistinguishable) is poorer for the $^{pmc}ZnTPP$ at the H site than for the $^{pmc}ZnTPP$ at the LB one. In this regard, it must be mentioned that, with the $^{pmc}ZnTPP$ at 2.8 \AA from the unreconstructed surface,¹³² the shortest (longest) $^S_HAg_{110}^{nn}-N^{Py}$ internuclear distance is 3.15 \AA (3.52 \AA), while the $^S_HAg_{110}^{nn}-C^m$ and $^S_HAg_{110}^{nn}-C^\alpha$ are 3.00 and 2.94 \AA , respectively. At variance, N^{Py} atoms lying along the $[001]$ direction are perfectly on top $^S_{LB}Ag_{110}^{nn}$ ($^S_{LB}Ag_{110}^{nn}-N^{Py}$ internuclear distance is 2.80 \AA); moreover, the $^S_{LB}Ag_{110}^{nn}-C^m$ and $^S_{LB}Ag_{110}^{nn}-C^\alpha$ are 2.86 and 2.95 \AA , respectively. In the presence of a significant substrate \rightarrow adsorbate charge transfer, which, as already mentioned implies the maximization of the overlap between low-lying $^{pmc}\pi^*$ MOs and SCM ns SALCs of suitable symmetry, the favored CM(110) chemisorption site should be the LB one.

Available experimental evidence well agrees with the above-reported considerations. Baklanov *et al.*¹³² investigated the adsorption configuration of ZnP on Cu(111) and Ag(111) by

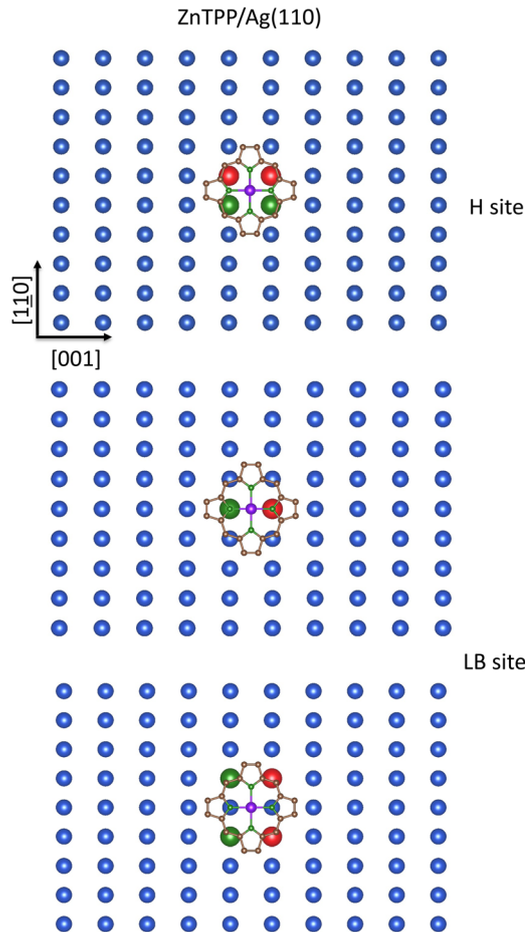


Fig. 12 Schematic representation of the $^S_HAg_{110}^{nn}$ (upper panel), $^S_{LB}Ag_{110}^{nn}$ (middle panel), and $^S_{LB}Ag_{110}^{nn}$ (bottom panel) 5s AOs SALCs transforming as the ir b in the local C_2 symmetry of the Ag(110) bulk-terminate surface superimposed to the optimized $^{pmc}ZnTPP$ placed at 2.8 \AA above the substrate and oriented with $Ni-N^{Py}$ bonds aligned along the $[110]$ and $[001]$ directions. Only Ag(110) topmost layer atoms are displayed for clarity. Large spheres represent $^S_HAg_{110}^{nn}$ and $^S_{LB}Ag_{110}^{nn}$ 5s AOs, while different colors account for their different phases.

combining XSW, XPS, STM, bond-resolved AFM, LICAD, and DFT-based supercell periodic calculations (limited to the ZnP/Cu(111) interface) to conclude that the ZnP species chemisorbed on Cu(111) occupies the B site with two $Zn-N^{Py}$ bonds oriented along the primary axis (see Scheme 1 of ref. 132). In addition, De Luca *et al.*,¹³³ based on STM and PES measurements on the ZnTPP/Au(111) interface, proposed the same B chemisorption site, superseding former claims of adsorption on T site.¹³⁴ Finally, Amsalem *et al.*¹³⁵ investigated the electronic and vibrational properties of the ZnPc/Ag(110) interface by LEED, STM, and HREELS measurements, whose outcomes were consistent with a quite strong substrate \rightarrow adsorbate charge transfer, implying the partial filling of the ZnPc π^* LUMO. Amsalem *et al.*¹³⁵ also proposed a tentative model for the geometrical structure of 1 ML ZnPc/Ag(110), although arbitrarily placing the Zn^{II} ions on T and SB sites instead of the LB one.¹³⁵



4. Conclusions

Structural and electronic reasons determining a crystal field stronger in the Pc^{2-} ligand than in the TPP^{2-} one have been identified, and the impact of this difference on the frontier electronic structure of free MTPP and MPc ($Z^M = 23-30$; the largest series so far ever considered) has been investigated by DFT calculations with the ADF package. Easily accessible information about group 11 elements (crystal structure,⁹⁵ nobleness,⁶³ surface geometry¹³⁶) combined with a thorough description of the free MTPP and MPc frontier electronic structure offer a cost-effective approach to foresee the chemisorption site of MTPP and MPc on low-index Cu, Ag, and Au surfaces, provided that two requirements are fulfilled: an established substrate \rightarrow adsorbate charge transfer; the absence of significant distortion of the macrocycle upon adsorption. A thorough comparison of the electronic structures of isolated MTPP and MPc and their building blocks indicates a more efficient substrate \rightarrow adsorbate charge transfer in MPc than in MTPP. The stronger MPc ligand field, determined by the shrinking of the coordinative pocket and the higher π acceptor capability of MPc (see the ESI^\dagger), yields an overall electronegativity larger in MPc than in MTPP, which brings the MPc LUMOs closer in energy to the ^5CM SALCs. Because of their very similar electronegativities,^{65,66} Cu and Ag surfaces show similar behaviors and predictable chemisorption sites, while the Au nobleness⁶³ prevents weighty charge transfer and only the chemisorption site of FePc (in its $^3\text{E}_g^{(1)}$ state) and CoPc on Au surfaces^{59,137,138} may be shortly, easily, and cheaply foreseen. This information also provides an useful input to avoid time-expensive trial-and-error computations, when the full interfacial structural and electronic details details must be retrieved through costly numerical experiments. The predictive capability of the semi-quantitative approach herein proposed reinforces the idea that the adsorbate–substrate interaction, if strong, is a phenomenon mainly local in character¹³⁹ where the accuracy of the local description of the potential is more relevant than the inclusion of long-range effects and then it can be suitably treated through the molecular cluster approximation.¹³⁹⁻¹⁴¹

Acronyms

ADF	Amsterdam density functional	DFT	Density functional theory
AFM	Atomic force microscopy	DMRG	Density matrix renormalization group
AO	Atomic orbital	DV- $X\alpha$	Discrete variational $X\alpha$
ARPES	Angular resolved photoelectron spectroscopy	EA	Electron affinity
ASPCS	Atomic subshell photoionization cross-sections	ESI	Electronic supplementary material
B	Bridge chemisorption site	fcc	Face centered cubic
B3LYP	Becke, 3-parameter, Lee–Yang–Parr	FMO	Frontier molecular orbital
BP86	Becke–Perdew 86	GED	Gas phase electron diffraction
CASPTn	Multi reference complete active space perturbation theory	GGA	Generalized gradient approximation
CASSCF	Complete active space self-consistent field	GS	Ground state
CM	Coinage metal	H	Hollow chemisorption site
C_n	Clockwise rotation through $\frac{2\pi}{n}$ radians	H_2P	Porphine
CP	Contour plot	H_2Pc	Phthalocyanine
		H_2Pz	Porphyrazine
		H_2TPP	<i>meso</i> -Tetraphenylporphyrin
		hcp	Hexagonal close packed
		HOMO	Highest occupied molecular orbital
		HREELS	High resolution electron energy loss spectroscopy
		HS	High spin
		i-Ind	Isoindole
		IE	Ionization energy
		IETS	Inelastic electron tunneling spectroscopy
		IPS	Inverse photoemission spectroscopy
		<i>ir</i>	Irreducible representation
		IR	InfraRed
		IS	Intermediate spin
		IUPAC	International union of pure and applied chemistry
		LB	Long bridge
		LEED	Low energy electron diffraction
		LICAD	Ligand-induced central atom displacement
		LS	Low spin
		LUMO	Lowest unoccupied molecular orbital
		<i>m</i>	<i>meso</i>
		M	Transition metal
		ML	Monolayer
		MO	Molecular orbital
		MOT	Molecular orbital tomography
		MP	Porphinato transition metal complex
		MPc	Phthalocyaninato transition metal complex
		MTPP	Tetraphenylporphyrinato transition metal complex
		<i>n</i>	Principal quantum number
		NEXAFS	Near edge X-ray absorption fine structure
		nn	Nearest neighbour
		nnn	Next nearest neighbour
		OEP^{2-}	2,3,7,8,12,13,17,18-Octaethylporphyrinato ion
		OMTS	Orbital-mediated tunneling spectroscopy
		P^{2-}	Porphinato ion
		Pc^{2-}	Phthalocyaninato ion
		PED	Photoelectron diffraction
		PES	Photoelectron spectroscopy
		Ph	Phenyl group
		pmc	Planar aromatic macro cycle core
		PPhMe_2	Dimethylphenyl phosphine
		Py	Pyrrrole
		RASPT2	Multiconfigurational second-order perturbation theory restricted active space



RT	Room temperature
S	Total spin quantum number
SALC	Symmetry adapted linear combination
SB	Short bridge
^s CM	Surface coinage metal
SO	Spin orbital
STM	Scanning tunneling microscopy
STS	Scanning tunneling spectroscopy
T	Top chemisorption site
TS	Transition state
TSEA	Transition state electron affinity
TSIE	Transition state ionization energy
THF	Tetrahydrofuran
TPP ²⁻	meso-Tetraphenyl porphyrinato ion
TTP ²⁻	meso-Tetra-p-tolylporphyrinato ion
UV	Ultraviolet
VASP	Vienna <i>ab initio</i> Simulation Package
VdW	van der Waals
XC	Exchange–correlation
XMCD	X-Ray magnetic circular dichroism
XPS	X-Ray photoelectron spectroscopy
XSW	X-Ray standing wave
Z	Atomic number
χ	Electronegativity
μ	Magnetic moment
ν	Frequency
3D	Three dimensional

Author contributions

S. C.: investigation, data curation, writing – review & editing;
 I. C.: investigation, data curation, writing – review & editing;
 V. F.: investigation, data curation, writing – review & editing;
 L. S.: investigation, data curation, writing – review & editing;
 L. F.: investigation, data curation, writing – review & editing; M. C.:
 conceptualization, investigation, methodology, resources, supervision,
 writing original draft, writing – review & editing.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

There are no conflicts to declare.

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