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# Magnetic dimeric metal—porphyrin rings mechanically bonded around carbon nanotubes: the role of nanotube defects modulating magnetic properties†

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Carbon nanotubes constitute an excellent option to connect molecular nanomagnets (MNMs) to solidstate devices, allowing the electrical control of the molecule spin state. The main challenge in this area is to obtain stable nanohybrid structures that preserve the magnetic properties and functionality of the molecule. One of the strategies developed to build these structures combining MNMs and carbon nanotubes involves the encapsulation of the nanotube within magnetic organic macrocycle(s), leading to magnetic mechanically interlocked derivatives of carbon nanotubes (mMINTs). Recently, mMINTs with dimeric metal-porphyrin rings containing Cu<sup>2+</sup> or Co<sup>2+</sup> attached to carbon nanotubes have been synthesized. Different structural and spectroscopic techniques confirm that the metal centers in both mMINTs preserve the coordination sphere and structure of the free macrocycles, with only small deviations from the square planar geometry. However, different magnetic behaviors are observed depending on the metal. The Cu derivative has well-preserved spin geometry, with quantum coherence times on the microsecond scale, as in the free molecule ( $\tau = 25 \mu s$ ). In contrast, the continuous-wave electron paramagnetic resonance (EPR) spectrum measured on the Co derivative does not show any clear contribution from Co, although EXAFS unambiguously shows the presence of Co<sup>2+</sup>. In order to understand this different magnetic behavior, we studied mMINTs by means of periodic DFT calculations. The analysis of the optimized geometries, the spin density of the free macrocycles and the mMINTs, the adsorption energies, the extension of the charge transfer between the macrocycle and the nanotube and the density of states reveals a stronger macrocycle-nanotube interaction for the Co-mMINT. Our results point to carbon vacancies in the nanotubes as responsible for the different macrocycle-nanotube interaction and the observed loss of the Co<sup>2+</sup> EPR signal in the Co-mMINT.

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# Introduction

Molecular nanomagnets (MNMs), such as porphyrins, organic radicals, and single-ion magnets like polyoxometalates or phthalocyanines, are hot candidates for spintronic and quantum technology applications.<sup>1–6</sup> In particular, in recent years, metalloporphyrins and their derivatives have been extensively studied as potential qubits.<sup>7–11</sup> The use of MNMs in solid-state devices requires their immobilization on a substrate, which can produce changes in their magnetic properties affecting quantum tunneling, magnetic hysteresis or quantum coherence times. These properties determine the quality of the systems to be used as a qubit.<sup>12–14</sup>

The most commonly studied substrates for molecular nanomagnets are nanoparticles, metallic thin films, carbon-based nanostructures (graphene, fullerenes, and carbon nanotubes), metal–organic frameworks, and porous silicas.<sup>13</sup> Among them,

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carbon nanotubes have emerged as an excellent option to connect magnetic molecules to real devices, allowing the electrical control of the spin state of the molecule, as some nanotubes are excellent electrical conductors. <sup>15,16</sup> Several strategies have been explored to build nanohybrid structures combining magnetic molecules and carbon nanotubes, including both grafting the molecule on the external surface of the nanotube or encapsulating the molecule into the nanotube. <sup>17–20</sup> The

main challenge is to obtain stable structures that preserve the

magnetic properties and functionality of the molecule.

Recently, a new clipping strategy to bind single-walled carbon nanotubes (SWNTs) to macrocycles to form rotaxanelike species has been developed.21 The product is a system with a SWNT encapsulated within organic macrocycle(s), called mechanically interlocked derivatives of carbon nanotubes (MINTs). The formation of MINTs has been shown to be a promising strategy to immobilize functional molecules to SWNT surfaces in a kinetically stable fashion, but without covalent modification of the SWNTs.<sup>22</sup> The proposed method has been employed to several types of macrocycles which have led to species showing promising properties for various applications, such as for reinforcing polymers and designing new catalysts.22-25 Motivated by previous work, magnetic dimeric metal-porphyrin rings with copper or cobalt, both with S = 1/22, have been attached to SWNTs, to obtain magnetic MINTs (mMINTs).<sup>26</sup> Different structural and spectroscopic techniques confirm that the metal center in both mMINTs preserve the coordination sphere and structure with respect to the free metal-porphyrins, with only small deviations from the square planar geometry. The Cu derivative has a well-preserved spin geometry and quantum coherence times (25 µs for the free molecule) remain in the microsecond scale in the mMINT derivative. In contrast, the continuous-wave electron paramagnetic resonance (EPR) spectrum measured on the Co derivative does not show any clear contribution from Co<sup>2+</sup>, although EXAFS unambiguously shows the presence of Co<sup>2+</sup> in a slightly distorted square planar coordination sphere. It has been suggested that the proximity and interaction of the spin with the SWNT, and therefore its EPR spectrum, could be determined by the molecular orbital occupied by the unpaired electron on the metal, which is different for cobalt  $(d_{z^2})$  and copper  $(d_{x^2-y^2})$  dimeric metal-porphyrin rings.

In this work, we have employed periodic DFT calculations to perform a detailed study of the electronic structure and magnetic properties of mMINTs containing dimeric metalporphyrin rings with Co<sup>2+</sup>or Cu<sup>2+</sup> and mixed mMINTs containing both metals. We have considered pristine nanotubes and analyzed the effect of the presence of carbon vacancies on the nanotubes. Carbon vacancies are intrinsic defects on carbon nanotubes, and their presence has been shown to determine the reaction tendency of these systems. <sup>18,27</sup> For instance, they have been found to be responsible for the anomalously strong binding of NO<sub>2</sub> molecules. <sup>28</sup> For the studied systems, our results reveal that carbon vacancies play a key role in the macrocycle–nanotube interaction and, therefore, their inclusion is essential to get a realistic model of the system. We

also found that in the presence of carbon vacancies the macrocycle–nanotube interaction is significantly stronger for the Co<sup>2+</sup> derivative that for the Cu<sup>2+</sup> one. This strong macrocycle–nanotube interaction modifies the electronic structure and magnetic properties of the Co<sup>2+</sup> derivative and is responsible for the loss of the Co<sup>2+</sup> EPR signal for Co atoms directly interacting with defects. Finally, our results allow us to predict a similar effect on the EPR spectra of mixed mMINTs, containing both Co<sup>2+</sup> and Cu<sup>2+</sup>, where Co<sup>2+</sup> would be EPR-silent and only the Cu<sup>2+</sup> signal would be registered.

# Computational details

The electronic structure and magnetic properties of mMINTs have been studied within periodic density functional theory (DFT) using the VASP (Vienna *ab initio* simulation package) code.  $^{29-32}$  We employed the generalized gradient approximation (GGA) with the revised Perdew–Burke–Ernzerhof functional (rPBE) $^{33}$  and projector-augmented wave (PAW) potentials.  $^{34,35}$  Valence electrons are described using a planewave basis set with a cutoff of 500 eV and the  $\Gamma$ -point of the Brillouin zone is used.  $^{36}$  van der Waals interactions are taken into account through the DFT-D3 method of Grimme, with Becke–Johnson damping function.  $^{37,38}$  We have considered mMINTs containing two Co-porphyrins, two Cu-porphyrins as well as mixed mMINTs containing one Co-porphyrin and one Cu-porphyrin.

We have used a 6,5-SWNT with optimized lattice parameters a = b = 7.50 Å and c = 40.96 Å, containing 364 carbon atoms. The nanotubes are placed along the z axis. 42.5 Å of vacuum are added in x and y directions to avoid the interaction between nanotubes (50.00 Å × 50.00 Å × 40.96 Å cell). To evaluate the effect of the presence of a defect on the nanotube, one C atom has been removed from the supercell, leading to a vacancy density of 0.1 nm<sup>-2</sup>.

To analyse the effect of the encapsulation of the SWNT on the electronic structure and magnetic properties of the macrocycles, we have also optimized isolated  $\mathrm{Co^{2^+}}$  and  $\mathrm{Cu^{2^+}}$  dimeric metal–porphyrin rings, in the same cell of the mMINTs (50.00 Å × 50.00 Å × 40.96 Å). Moreover, to assess the effect of dimer and mMINT formation on the porphyrin ring structure, the isolated Co and Cu porphyrin structures have also been optimized.

Electronic relaxation has been performed until the change in the total energy between two consecutive steps is smaller than  $10^{-5}$  eV and the ionic relaxation has been performed until the Hellmann–Feynman forces were lower than 0.03 eV Å $^{-1}$ . For each structure we have optimized different spin-polarized magnetic solutions, considering ferromagnetic or antiferromagnetic interactions between the two metal atoms in the macrocycle. Several starting geometries have been used for geometry optimization runs.

Adsorption energies,  $E_{\rm ads}$ , are calculated as  $E_{\rm ads} = E_{\rm mMINT} - (E_{\rm nanotube\_opt} + E_{\rm dimer\_opt})$ . The energies of the nanotube  $(E_{\rm nanotube\_opt})$  and the dimeric porphyrin rings  $(E_{\rm dimer\_opt})$  are

obtained from two independent geometry optimization calculations using the same supercell as the adsorbed molecule, in order to minimize numerical errors. Thus, negative adsorption

energies represent bound states.

Finally, interaction energies,  $E_{int}$ , are calculated as  $E_{int}$  =  $E_{\rm mMINT}$  – ( $E_{\rm nanotube\_sp}$  +  $E_{\rm dimer\_sp}$ ). The energies of the nanotube  $(E_{\text{nanotube sp}})$  and the dimeric porphyrin rings  $(E_{\text{dimer sp}})$ are obtained from a single point calculation of the carbon nanotube or the macrocycle with the optimized geometry of the whole mMINT system and using the same supercell as the whole system. Interaction energies are included in Table S2.†

### Results and discussion

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### Free dimeric metal-porphyrin macrocycles

The structures of both Co<sup>2+</sup> and Cu<sup>2+</sup> dimeric metal-porphyrin rings have been optimized in two different magnetic states (Fig. 1).  $Co^{2+}$  and  $Cu^{2+}$  are both  $S = \frac{1}{2}$  centers and, therefore, the interaction between the two metals on each macrocycle can be ferromagnetic or antiferromagnetic, leading to one structure with total magnetization 0 (Cu-dimer-0 and Codimer-0 in Table 1) and another structure with total magnetization 2 (Cu-dimer-2 and Co-dimer-2 in Table 1). The energy difference between both magnetic solutions depends on the amplitude of the magnetic coupling between the metal centers.

No structural differences have been found between both spin states. After the formation of the macrocycle, the porphyrin structure preserves the preexisting planar geometry (Table S1†). Inside the macrocycle, the two porphyrin rings are almost parallel. Distances between metallic centres are 12.49  $\mbox{\normalfont\AA}$ and 12.51 Å for Co<sup>2+</sup> and Cu<sup>2+</sup> derivatives, respectively. Moreover, for both metals, we found no meaningful energy difference between both magnetic solutions (the energy difference being smaller than 0.2 kJ mol<sup>-1</sup>, Table 1). That is, the metal centers are not coupled, and the magnetic interaction between them is zero.

Fig. 1 shows the spin density for all the considered structures and spin states, calculated as the difference between the  $\alpha$  and  $\beta$  electronic densities. Interestingly, there are appreci-

**Table 1** Relative energies (kJ mol<sup>-1</sup>), total magnetization ( $\mu_B$ ) and magnetization on different atoms ( $\mu_B$ ) of the isolated dimeric metal-porphyrin rings

		Magnetization								
Structure	$\Delta E$	Total	M1 (Por	Ring1 ph1)	M2 (Por	Ring2 ph2)				
Co-dimer-0	0.05	0.0	-1.026 (-0.	0.092 .934)	1.026	-0.092 934)				
Co-dimer-2	0.00	2.0	1.026	-0.092 934)	1.026	-0.092 934)				
Cu-dimer-0	0.00	0.0	0.564	0.382 946)	-0.564	-0.382 .946)				
Cu-dimer-2	0.15	2.0	0.564 0.382 (0.946)		0.564	0.382 946)				

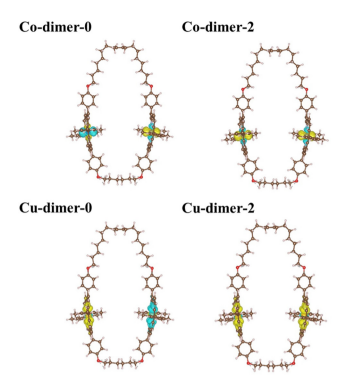


Fig. 1 Spin density for the optimized geometries of Co<sup>2+</sup> (top) and Cu<sup>2+</sup> (bottom) dimeric porphyrin rings with total magnetization 0 (left) and 2 (right). Plots are obtained with an isodensity value of  $10^{-3}$  e bohr<sup>-3</sup>. Brown, light blue, red, white, dark blue and orange spheres represent C, N, O, H, Co and Cu, respectively.

able differences in spin density depending on the metal. In the case of the Cu-dimer, the spin density is mainly localized on the porphyrin plane. The unpaired electron occupies the  $d_{x^2-y^2}$  orbital, with a non-negligible participation of the  $\sigma$  orbitals of the surrounding N porphyrin atoms (as shown in Table 1, the magnetization of the porphyrin rings is relevant for the Cu-dimer). In the case of the Co-dimer, the spin density is mainly out of the plane of the porphyrin ring. The unpaired electron occupies the d<sub>22</sub> orbital, with a minor participation of the porphyrin (Table 1). In both cases there is no spin density in the rest of the organic macrocycle.

### Co<sup>2+</sup> and Cu<sup>2+</sup> mMINTs with pristine nanotubes

To study the encapsulation of the SWNT within the dimeric metal-porphyrin rings geometry optimizations have been run from several starting structures (Fig. 2). In the most stable structures metal-nanotube distances are 3.51-3.58 Å for the Co<sup>2+</sup> dimer and 3.47–3.53 Å for the Cu<sup>2+</sup> dimer (Table 2). There are no significant structural differences for the different metal derivatives and they also present almost identical adsorption energies (Table 2). In agreement with previous experimental work,<sup>26</sup> we found that the metal centers in the mMINT porphyrins preserve the coordination sphere and structure when cycled, with only slight distortions of the square planar geometry. Moreover, optimized geometries, with the macrocycle embracing the nanotube, are consistent with the registered

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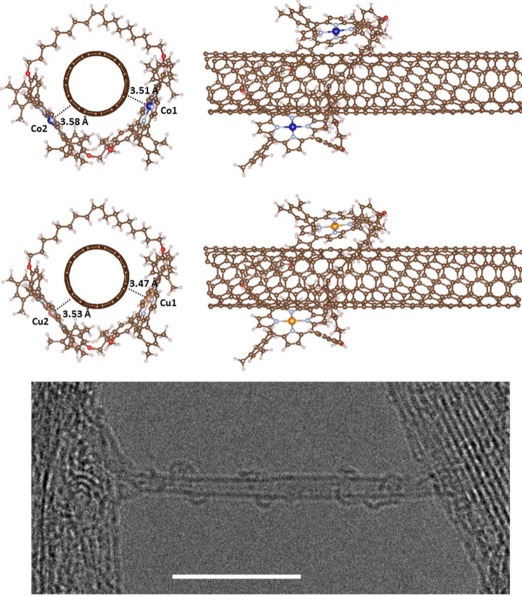


Fig. 2 Optimized structures of  $Co^{2+}$  (top) and  $Cu^{2+}$  (middle) mMINTs with pristine nanotubes with total magnetization 0 (Co-mMINT-0 and Cu-mMINT-0 in Table 2). There are no differences between structures with ferromagnetic and antiferromagnetic interactions of the unpaired electrons. Brown, light blue, red, white, dark blue and orange spheres represent C, N, O, H, Co and Cu, respectively. HR-TEM image showing a single Co-mMINT (bottom), where four  $Co^{2+}$  dimeric metal-porphyrin rings can be seen embracing the SWNT. Scale bar: 400 nm. Adapted with permission from *J. Am. Chem. Soc.*, 2021, **143**(50), 21286–21293. Copyright 2021 American Chemical Society.

high-resolution microscopy (HR-TEM) images, as shown in Fig. 2 (bottom).  $^{26}$ 

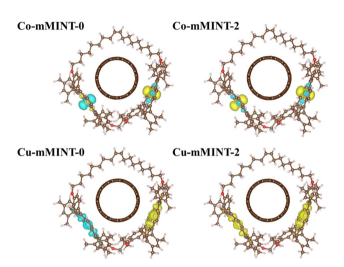
As in the case of free dimers, we found that structures with ferromagnetic (M-mMINT-2) and antiferromagnetic (M-mMINT-0) interactions between the unpaired electrons of the metal centers are degenerated for mMINTs. Interestingly, spin density remains unaltered after mMINT formation, with no spin density observed in the carbon nanotube, as shown in Fig. 3. Moreover, Fig. 4 (top) illustrates the charge density difference,  $\Delta \rho$ , between the mMINT, the nanotube, and the dimer, denoted as  $\Delta \rho = \rho(\text{mMINT}) - [\rho(\text{nanotube}) + \rho(\text{dimer})]$ .

Notably, there is no significant charge transfer between the dimers and the nanotube in any of the systems.

The magnetism of Co<sup>2+</sup> and Cu<sup>2+</sup> dimeric metal-porphyrin rings has been analyzed by electron paramagnetic resonance (EPR) spectroscopy. Interestingly, while in Cu-mMINT spectral features coming from the Cu porphyrin centers appear, no EPR signature of the Co-mMINT has been observed, although EXAFS unambiguously shows the presence of Co<sup>2+</sup> in a slightly distorted square planar coordination sphere. Based on this observation, it has been suggested that the different molecular orbitals occupied by the unpaired electrons of Cu and Co,

**Table 2** Adsorption energies (kJ mol<sup>-1</sup>), selected distances (Å), total magnetization ( $\mu_B$ ) and magnetization on different atoms ( $\mu_B$ ) of mMINTs with pristine nanotubes

		Distances M-C		Magnetization						
Structure	$E_{ m ads}$	M1-C	М2-С	Total	M1 (Po	Ring1 rph1)	M2 (Por	Ring2 ph2)	NT	
Co-mMINT-0	-241.2	3.51	3.58	0.0	1.023	-0.089	-1.024	0.090	0.000	
					(0.934)		(-0.934)			
Co-mMINT-2	-241.2	3.51	3.58	2.0	1.023	-0.089	1.024	-0.090	0.000	
					(0.934)		(0.934)			
Cu-mMINT-0	-244.1	3.47	3.53	0.0	0.563	0.382	-0.562	-0.381	0.000	
					(0.945)		(-0.943)			
Cu-mMINT-2	-244.1	3.47	3.53	2.0	0.563	0.382	0.562	0.381	0.000	
					(0.945)		(0.943)			



**Fig. 3** Spin density for the Co-mMINT (top) and Cu-mMINT (bottom) with pristine nanotubes with total magnetization 0 (left) and 2 (right). Plots are obtained with an isodensity value of  $10^{-3}$  e bohr<sup>-3</sup>. Brown, light blue, red, white, dark blue and orange spheres represent C, N, O, H, Co and Cu, respectively.

 $d_{x^2-y^2}$  and  $d_{z^2}$ , respectively, may determine the mean distance of the porphyrin to the nanotube and the strength of the interaction of the spin with the SWNT, and therefore its EPR spectrum. Interestingly, our results confirm that the choice of the metal determines the orbital occupied by the unpaired electron, the in-plane  $d_{x^2-y^2}$  orbital of  $Cu^{2+}$  derivatives and the out-of-plane  $d_{z^2}$  orbital in the case of Co-mMINTs. However, the analysis of metal–NT distances, the adsorption energies, the spin density and the extension of the charge transfer between the macrocycle and the nanotube reveals a similar interaction between the macrocycle and the nanotube for both studied metals, which will not affect the magnetic properties of the porphyrin dimeric ring. Therefore, this first model is not able to explain the different behaviors for the Co-mMINT and Cu-mMINT observed in experiments.

### Co<sup>2+</sup> and Cu<sup>2+</sup> mMINTs with defective nanotubes

Carbon vacancies are intrinsic defects on carbon nanotubes, and their presence has been shown to determine their reaction

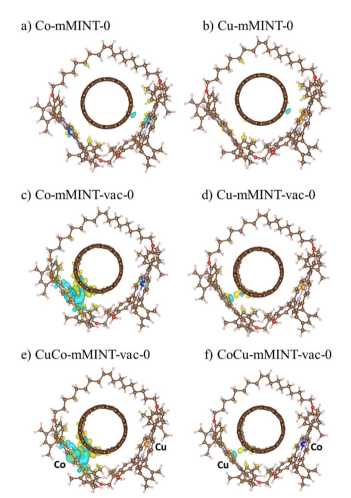


Fig. 4 Charge density difference  $\Delta\rho$  between the mMINT, the nanotube and the dimer for the mMINTs with pristine nanotubes, (a and b), mMINTs with defective nanotubes, and (c and d) mixed mMINTs, both with  $\mathrm{Co}^{2+}$  (e) or  $\mathrm{Cu}^{2+}$  (f) close to the C vacancy. Plots are presented with an isodensity value of  $10^{-3}$  e bohr<sup>-3</sup>. Yellow (positive difference) and cyan (negative difference) surfaces represent the charge accumulation and loss of electron charge density upon adsorption, respectively. Brown, light blue, red, white, dark blue and orange spheres represent C, N, O, H, Co and Cu, respectively.

tendency. 18,27,28 In fact, discrepancies between theoretical simulations and experimental results in studies concerning the adsorption of small molecules on carbon nanotube sidewalls have been ascribed to the use of non-realistic models, neglecting common topological and structural defects on nanotubes. 28,39,40 In order to model our system in a more realistic way, we have analyzed the effect of introducing a carbon vacancy on the nanotube.

Removing a C atom from the nanotube generates four unpaired electrons and produces a noticeable distortion of the structure near the vacancy, with the formation of a new carbon-carbon bond, leading to the creation of a pentagonal ring, and an out-of-plane shift of the dangling carbon atom opposite to the pentagon (labeled as C<sub>vac</sub>) (Fig. S1†). The two remaining electrons can either couple ferromagnetically (resulting in a total magnetization of 2) or form a diamagnetic solution with a total magnetization of 0, with the diamagnetic solution being 21.1 kJ mol<sup>-1</sup> more stable.<sup>28</sup> Fig. 5 shows the spin density for both solutions, calculated as the difference between the electronic density of  $\alpha$  and  $\beta$ -electrons. In the ferromagnetic solution, two electrons occupy a different region of space: one electron is strongly localized on a carbon atom beside the vacancy ( $C_{vac}$ ), having a  $\sigma$ -character, while the other electron strongly delocalizes on the  $\pi$ -states of the nanotube. In the diamagnetic solution there is only a residual spin density on the nanotube. Interestingly, the creation of a C vacancy on the nanotube has an important effect on the density of states (Fig. 6, top); it introduces states around the Fermi level, decreasing the band gap, and leading to more conductive species.

Fig. 7 shows the optimized structure of Co<sup>2+</sup> and Cu<sup>2+</sup> mMINTs with defective nanotubes. For both metals we observe that, after the removal of one C atom of the nanotube, the macrocycle structure changes in such a way that one of the metal centers (M2) moves closer to the vacancy. This fact produces an increase of one of the metal-nanotube distances  $(d_{\rm M1-C})$  from 3.5 Å to 4.1-4.2 Å and the decrease of the other

metal-nanotube distance ( $d_{M2-C}$ ). However, this reduction of the distance between the metal that is close to the vacancy and the nanotube is different for both mMINTs: the  $d_{Cu2-C}$  distance is 2.8 Å and the  $d_{\text{Co2-C}}$  distance is 2.0 Å (Table 3). These different distances constitute the first evidence of different interactions between the dimer and the defective nanotube depending on the metal. Adsorption energies also show a stronger interaction between the nanotube and the ComMINT. While for the Cu-mMINT the adsorption energy remains almost unaltered after C removal, for the Co-mMINT with defective nanotube the adsorption energy is around 33 kJ mol<sup>-1</sup> larger than that for the Cu-mMINT (Table 3).

Important differences between Co<sup>2+</sup> and Cu<sup>2+</sup> derivatives are found when analyzing total magnetization. Two different magnetic solutions, which are energetically degenerated, are obtained for each mMINT. For the Cu-mMINT total magnetization values are 0 (Cu-mMINT-vac-0) and 2 (Cu-mMINT-vac-2). Spin density plots for these two structures (Fig. 8) show that the spin density on the Cu-dimer remains unaltered after vacancy formation, with one unpaired electron on each  $d_{x^2-y^2}$ orbital and the participation of the porphyrin ring. The nanotube presents a residual spin density, exactly the same situation observed for the isolated nanotube. In contrast, total magnetization is strongly modified after vacancy creation for the Co derivative. We intended to get solutions with total magnetization of 0 and 2, resulting from antiferromagnetic or ferromagnetic interactions of Co<sup>2+</sup> unpaired electrons, imposing initial magnetization 1 or -1 on each  $Co^{2+}$ . However, after geometry optimization we obtained two equivalent magnetic solutions, with total magnetization around 1 (-0.853 for ComMINT-vac-0 and 1.154 for Co-mMINT-vac-2). In both structures spin density plots (Fig. 8) show one unpaired electron occupying the  $d_{z^2}$  orbital of Co1 (the cobalt atom that is far away from the vacancy) while for Co2 (close to the vacancy) there has been a strong reduction of the magnetization (from -1.024 with the pristine nanotube to -0.135 with defects). These results are consistent with the charge density differ-

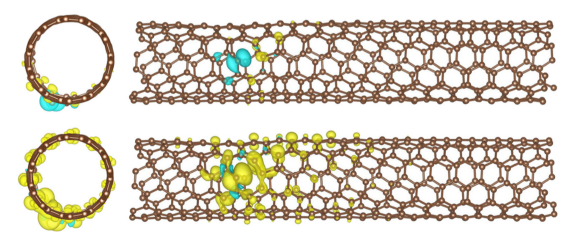


Fig. 5 Spin density for the defective nanotube with total magnetization 0 (top) and 2 (bottom). Plots are obtained with an isodensity value of  $10^{-3}$  e

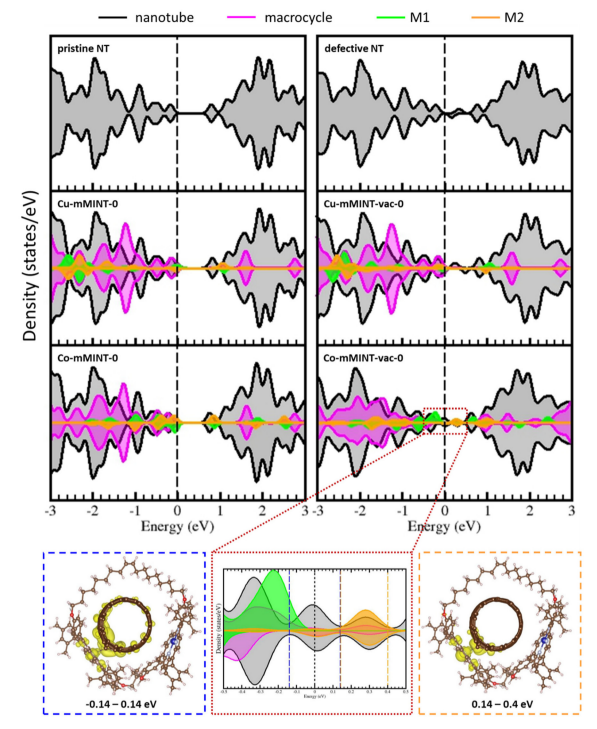


Fig. 6 Projected density of states for the SWNT and for mMINTs with a pristine nanotube (left) and for the mMINTs with a defective nanotube (right). Zoomed-in view of the region around the Fermi level for Co-mMINT-vac-0 (red box), the electronic density plots of the states in the region [-0.14, 0.14] eV (blue box) and the lowest empty states in the region [0.14, 0.4] eV (orange box). Plots are obtained with an isodensity value of  $10^{-3}$  e bohr<sup>-3</sup>. Brown, light blue, red, white and dark blue and orange spheres represent C, N, O, H and Co, respectively.

ences shown in Fig. 4(c), where we can appreciate a significant charge transfer from Co2 to the nanotube in the presence of a carbon vacancy.

To gain a deeper insight into the electronic structure of mMINTs we compared the projected density of states (pDOS)

for the mMINTs with and without carbon defects on the nanotube (Fig. 6). For both mMINTS with pristine nanotubes the highest occupied states are located on the metal atoms. The lowest unoccupied states correspond to nanotube states in the case of Cu-mMINT-0 while for Co-mMINT-0 they have contriNanoscale **Paper** 

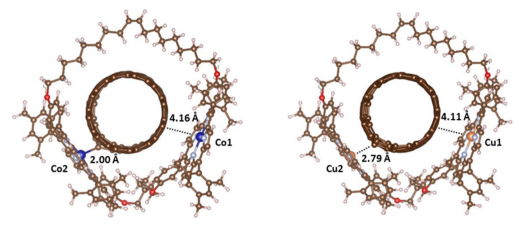


Fig. 7 Optimized structures of Co<sup>2+</sup> (left) and Cu<sup>2+</sup> (right) dimeric metal-orphyrin rings around defective carbon nanotubes (mMINTs). Brown, light blue, red, white, dark blue and orange spheres represent C, N, O, H, Co and Cu, respectively.

**Table 3** Adsorption energies (kJ mol<sup>-1</sup>), selected distances (Å), total magnetization ( $\mu_B$ ) and magnetization on different atoms ( $\mu_B$ ) of mMINTs with defective nanotubes. Cvac refers to the carbon atom adjacent to the carbon vacancy, which is the C atom closest to M2. NT refers to the sum of the magnetization over all the NT atoms, including  $C_{\text{vac}}$ 

Structure	$E_{ m ads}$	Distance		Magnetization								
		M1-C	M2-C <sub>vac</sub>	Total	M1 (Por	Ring1 ph1)	M2 (Por	Ring2 ph2)	$C_{\rm vac}$	NT		
Co-mMINT-vac-0	-259.6	4.162	1.995	-0.853	-1.026 (-0.	0.092 934)	-0.135 (-0.	0.008	0.093	0.157		
Co-mMINT-vac-2	-259.6	4.162	1.995	1.154	1.026	-0.092 934)	-0.135 (-0.	0.008 .127)	0.093	0.162		
Cu-mMINT-vac-0	-226.8	4.111	2.785	0.044	-0.552	-0.377 929)	0.568	0.375 943)	0.123	0.038		
Cu-mMINT-vac-2	-226.8	4.105	2.794	-2.008	-0.553 (-0.	-0.377 930)	-0.568 (-0.	-0.374 .942)	-0.113	-0.033		

bution from both the nanotube and Co. In both cases pDOS show a band gap of about 0.7 eV, which disappears in the case of the defective nanotube, with the presence of some states in the (original) band gap region (range: 0.0-0.7 eV). The C vacancy does not affect the Cu states, as they appear exactly at the same energy as in the case of the pristine nanotube. The states related to the vacancy only have contribution from the nanotube for the system containing Cu. In contrast, we appreciate important changes in the states of the Co atom close to the vacancy, as the band gap states have contribution of both the nanotube and Co2 metal, indicating a strong interaction between the metal center and the nanotube. The states with macrocycle contribution are also influenced by the defect, becoming wider and stabilized (all states appear at lower energies) for the defective nanotube, because of the dimer-nanotube interaction. As Co1 remains far away from the vacancy, the states projected on this atom remain unaltered after the defect formation. Looking closer, the states around the Fermi level of Co-mMINT-vac-0 show a significant mixing of the Co2  $d_{z^2}$  orbital and those of the nanotube. Fig. 6 (bottom) shows the electronic density of the states immediately above and below the Fermi level, within the range -0.14 to 0.14 eV (blue

box in Fig. 6) and the lowest unoccupied states in the range 0.14 to 0.4 eV (orange box in Fig. 6). The former are partially occupied, with a large contribution from nanotube states and a minor contribution of Co2 d<sub>22</sub> states. Interestingly, the lowest fully empty states show an important contribution from Co2 d<sub>22</sub> states. This indicates some extent of electron transfer from the Co2 atom to the nanotube and explains the total magnetization of about 1 obtained for both solutions. The unpaired electron of the Co2 center is transferred to the nanotube, and then the total magnetization of the mMINTs is 1, that is, the system contains a single unpaired electron located on the 3d<sub>2</sub><sup>2</sup> orbital of the Co1 atom, the one more distant to the carbon defect.

All the studied properties, geometries, adsorption energies, magnetization on different atoms, spin density representations, charge density differences and projected density of states, confirm stronger dimer-nanotube interactions for the Co-mMINT than for the Cu-mMINT when carbon vacancies are present on the nanotube. As a consequence of this strong interaction on Co-mMINT, magnetization on one Co atom is reduced, which could explain the loss of the Co<sup>2+</sup> EPR signal for  $Co_2$ .

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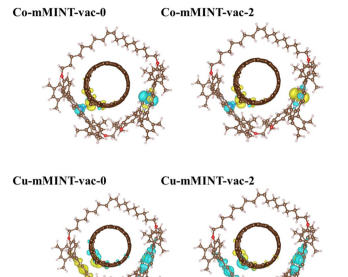


Fig. 8 Spin density for the Co-mMINT (top) and Cu-mMINT (bottom) with defective nanotubes. Two geometries are shown for each mMINT-vac, optimized with different spin solutions. Plots are obtained with an isodensity value of  $10^{-3}$  e bohr<sup>-3</sup>. Brown, light blue, red, white and dark blue and orange spheres represent C, N, O, H and Co, respectively.

### Mixed mMINTs with defective nanotubes

Finally, we extended the study to mixed mMINTS containing both one Co<sup>2+</sup> and one Cu<sup>2+</sup> atom. As the importance of the inclusion of defects on the nanotube to model the macrocycle–nanotube interaction has been proven, one carbon vacancy has been introduced in the nanotube, and two different structures have been optimized, one with the Cu atom close to the defect (CoCu-mMINT-vac) and the other one with the Co atom close to the defect (CuCo-mMINT-vac). For each structure two almost degenerated magnetic solutions have been found. Selected distances, relative energies, adsorption energies, total magnetizations and magnetization on selected atoms for the four structures have been included in

Table 4. Moreover, plots of the charge density differences are presented in Fig. 4 (bottom) and pDOS for the mixed mMINTs (with Cu<sup>2+</sup> or Co<sup>2+</sup> near the vacancy) are shown in Fig. S2.†

In general, in the presence of a carbon vacancy, all the analyzed properties of the mixed mMINT with a Co atom close to the vacancy are very similar to those of the homonuclear Co derivative, while the properties of the mixed mMINT with Cu close to the vacancy are almost identical to those of the homonuclear Cu derivatives.

The heteronuclear dimeric ring is stabilized by about 23.2 kJ mol<sup>-1</sup> when the vacancy is close to Co-porphyrin (CuCo-mMINT-vac). The adsorption energy for the most favorable mixed mMINT-vac structure is the same as that of the homonuclear Co-mMINT-vac. Hence, the CoCu-mMINT-vac adsorption energy is about 23 kJ mol<sup>-1</sup> smaller, and similar to the adsorption energy of Cu-mMINT-vac.

Our results reveal that in mixed mMINTs Co<sup>2+</sup> will approach carbon vacancies on the nanotube, and a significant charge transfer from Co<sup>2+</sup> to the nanotube will occur. We can predict that in the EPR spectrum of mixed mMINTS containing both Cu<sup>2+</sup> and Co<sup>2+</sup>, Co<sup>2+</sup> will be silent if it is close to the vacancy. In other words, only the Cu signal would be registered, as strong Co–vacancy interactions produce an important modification of the macrocycle electronic structure and a decrease on cobalt magnetization, with consequent loss of the Co<sup>2+</sup> EPR signal.

# Conclusions

In this work we have performed a detailed theoretical study of mechanically bonded magnetic Co<sup>2+</sup> and Cu<sup>2+</sup> dimeric metalporphyrin rings around carbon nanotubes (mMINTs) employing periodic DFT calculations. Our aim was to understand the differences found in the electron paramagnetic resonance (EPR) spectra of mMINTs containing different metals (Co<sup>2+</sup> and Cu<sup>2+</sup>). While in the Cu-mMINT the spectral features coming from the Cu<sup>2+</sup> porphyrin centers are observed once the

Table 4 Relative energies (kJ mol<sup>-1</sup>), magnetization ( $\mu_B$ ), selected distances (Å) and adsorption energies (kJ mol<sup>-1</sup>) of mixed mMINTs.  $C_{vac}$  refers to the carbon atom adjacent to the carbon vacancy, which is the C atom closest to M2. mag NT refers to the sum of the magnetization over all the NT atoms, including  $C_{vac}$ 

		Magnetization							Distances		
Structure	$E_{ m rel}$	Total	Co (Co-	$ m Ring_{Co}$ porf)	Cu (Cu	Ring <sub>Cu</sub> -porf)	$C_{\rm vac}$	NT	Со-С	Cu-C	$E_{ m ads}$
Cu <sup>2+</sup> close to the defect											
CoCu-mMINT-vac-0	23.18	0.001	-1.037 (-0.	0.087 950)	0.567 (0.	0.376 .943)	0.075	0.015	4.064	2.883	-237.4
CoCu-mMINT-vac-2	23.23	2.008	1.032	-0.088 944)	0.566 (0.	0.377 .943)	-0.002	-0.003	4.063	2.883	-237.4
Co <sup>2+</sup> close to the defect											
CuCo-mMINT-vac-0	0.01	0.861	0.138	-0.005 133)	0.562 (0.	0.381 .943)	-0.092	-0.158	1.993	4.160	-260.5
CuCo-mMINT-vac-2	0.00	1.131	-0.138 (-0.	0.004 134)	0.562 (0.	0.381 .943)	0.091	0.153	1.993	4.160	-260.5

macrocycle encapsulates the nanotube, no EPR signature of Co<sup>2+</sup> was observed in the case of the Co-mMINT.

In a first stage, we considered mMINTs formed with pristine nanotubes. For these systems, our results reveal that the interaction between the macrocycle and the nanotube will not affect the magnetic properties of the dimeric metal-porphyrin rings. Our model does not show any significant differences between the Co<sup>2+</sup> and Cu<sup>2+</sup> derivative properties that could explain the observed differences in registered spectra. We can conclude that the inclusion of defects on the nanotube is essential to get a realistic model and to simulate the dimernanotube interaction.

After the inclusion of a carbon vacancy in the systems, optimized geometries, spin density, magnetization on different atoms, adsorption energies, extension of the charge transfer between the macrocycle and the nanotube and density of states have been analyzed. Our results reveal that the macrocycle-nanotube interaction is stronger for the Co-mMINT than for the Cu-mMINT. As a consequence of this stronger interaction on Co-mMINT the electronic structure of the macrocycle is modified and magnetization on one Co<sup>2+</sup> atom is reduced, which could explain the loss of the Co2+ EPR signal for Co atoms directly interacting with defects. As a consequence, the Co<sup>2+</sup> EPR signal would decrease, and could be buried under the SWNT contribution in experimental measurements. Furthermore, a stronger interaction between the metal centers and the nanotubes could reduce coherence and broaden the signal, thereby contributing to a lower signal-to-noise ratio.

In all of the systems considered, the metal centers are not magnetically coupled, and the solutions with a ferromagnetic interaction and those with an antiferromagnetic interaction between them are degenerated.

Finally, we extended the work to include mixed mMINTs containing both one Co<sup>2+</sup> and one Cu<sup>2+</sup> atom. These mixed mMINTs have not yet been synthesized. We found that mixed mMINTs are more stable when the Co-porphyrin approaches a carbon vacancy on the nanotube. All the studied properties for such a structure are very similar to those of the system containing only Co2+ as the metal. The formation of mixed mMINTs involves a significant charge transfer from Co<sup>2+</sup> to the nanotube as in the homonuclear Co derivative. We can predict that in the EPR spectrum of mixed mMINTs containing both Cu<sup>2+</sup> and Co<sup>2+</sup> only the Cu signal would be registered, as the strong Co-vacancy interaction causes an important modification of the macrocycle electronic structure and a decrease of cobalt magnetization, with a consequent loss of the Co<sup>2+</sup> EPR signal.

Our work highlights the importance of incorporating defects into theoretical models to achieve more realistic representations of carbon nanotube derivatives. We have considered single carbon vacancies, as they are the most common defects in carbon nanotubes. Evaluating the impact of other types of defects, such as double vacancies, interstitials, bond rotations, or non-hexagonal rings, on the electronic structure and magnetic properties of carbon nanotube derivatives, although involving an extremely high computational cost, would also provide valuable insights and make a significant contribution to the field.

# **Author contributions**

Conceptualization: R. Sánchez-de-Armas, C. J. Calzado, E. Burzurí, E. M. Pérez, and N. Montenegro-Pohlhammer; methodology: C. J. Calzado, R. Sánchez-de-Armas, and N. Montenegro-Pohlhammer; funding acquisition: C. J. Calzado, E. Burzurí, and E. M. Pérez; investigation: R. Sánchez-de-Armas, C. J. Calzado, and N. Montenegro-Pohlhammer; formal analysis: Sánchez-de-Armas, C. J. Calzado, N. Montenegro-Pohlhammer, and E. Burzurí; resources: E. M. Pérez, E. Burzurí, and C. J. Calzado; writing-original draft preparation: R. Sánchez-de-Armas and C. J. Calzado; writing—review and editing: R. Sánchezde-Armas, C. J. Calzado, E. Burzurí, E. M. Pérez, and N. Montenegro-Pohlhammer.

# 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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