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One-dimensional Molecular Crystal of Phthalocyanine Confined into Single-Walled Carbon Nanotubes

Laurent Alvarez^{1,}, Fallou Fall², Anouar Belhboub^{1,3}, Rozenn Le Parc¹, Yann Almadori^{1,2}, Raul Arenal^{4,5}, Raymond Aznar¹, Philippe Dieudonné-George¹, Patrick Hermet⁶, Abdelali Rahmani³, Bruno Jousset⁷, Stéphane Campidelli⁷, Julien Cambedouzou⁸, Takeshi Saito⁹ and Jean-Louis Bantignies¹*

⁽¹⁾ Laboratoire Charles Coulomb (L2C), UMR 5221 CNRS-Université de Montpellier, Montpellier, F-France.

Laurent.Alvarez@univ-montp2.fr, 33 467 14 35 41

⁽²⁾ Physikalisches Institut, Albert-Ludwigs-Universität, 79104 Freiburg, Germany

⁽³⁾ Laboratoire d'Etude des Matériaux Avancés et Applications (LEM2A), Faculté des Sciences, Université Moulay Ismail, Boîte Postale 11201, Zitoune, 50000 Meknes, Morocco

⁽⁴⁾ Laboratorio de microscopias avanzadas (LMA), Instituto de Nanociencia de Aragon (INA), U. Zaragoza, c/ Mariano Esquillor s/n, 50018 Zaragoza, Spain

⁽⁵⁾ Fundacion ARAID, 50018 Zaragoza, Spain

⁽⁶⁾ Institut Charles Gerhardt Montpellier, UMR 5253 CNRS-UM2-ENSCM-UM1, Université Montpellier 2, Place E. Bataillon, 34095 Montpellier, France

⁽⁷⁾ CEA-Saclay, IRAMIS, NIMBE, Laboratoire d'Innovation en Chimie des Surfaces et Nanosciences (LICSEN), 91191 Gif-sur-Yvette Cedex, France

⁽⁸⁾ Institut de Chimie Séparative de Marcoule, UMR 5257 CEA/CNRS/ENSCM/Université Montpellier 2, 30207 Bagnols sur Cèze, France

⁽⁹⁾ Nanotube Research Center, National Institute of Advanced Industrial Science and Technology (AIST), Tsukuba 305-8565, Japan

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ABSTRACT: New 1D hybrid nano-systems are elaborated with metallated or metal free phthalocyanine molecules encapsulated into the hollow core of single-wall carbon nanotubes. The x-ray diffraction experiments coupled to simulation allow evidencing the 1D structural organization of the molecules inside the nanotubes. The angle between the molecule ring and the nanotubes axis is close to 32° as determined from our DFT calculations. Confined molecules display Raman spectra hardly altered with respect to the bulk phase, suggesting a rather weak interaction with the tubes. For comparison, non-covalent functionalization at the outer surface of the tube is also investigated. The vibrational properties of the molecules functionalized at the outer surface of tubes display important modifications. A significant curvature of the phthalocyanine is induced by the interaction with the tube walls, leading to change of the central atom position within the molecular ring, in good agreement with our first principle calculations.

INTRODUCTION

Phthalocyanine (Pc) are 18 π -electron aromatic molecules which can accommodate a wide range of atoms in their central cavity to form metallocomplexes (MPc).¹ Initially used as dyes and pigments, MPc molecules have attracted an increasing interest as molecular building blocks for catalytic and opto-electronics.²⁻⁵ In addition, depending on the nature of the central atom, MPc can carry a magnetic moment so that the molecules can be thought as nanomagnets, opening new routes for spintronics⁶ and data storage devices at the molecular level.⁷ It has been demonstrated that the optical or magnetic properties can be tuned not only by changing the central atom but also by altering their structural properties,¹ providing practical routes to fit different requirements. More precisely, the molecular orientation plays a key role in charge transport, light absorption, energy level alignment at interfaces in organic semiconductors⁸ and magnetic properties.¹ For instance, edge-to-edge molecular organization favors in-plane charge transport whereas face-to-face arrangement favors out-of-plane charge transport.⁸

Thus, controlling the structural properties is a key point for potential applications. In this context, carbon nanotubes can be used to constrain phthalocyanine molecule configuration and create original 1D assemblies affording new and tunable magnetic and opto-electronic properties. New hybrid systems containing phthalocyanine derivatives and carbon nanotubes or graphene have been developed during the last years.^{6,9-21} Nevertheless, no endohedral functionalization of single-walled carbon nanotubes (NTs) with MPc has been reported. There are only few studies dealing with the insertion of MPc inside large multi-walled nanotubes with an average inner diameter around 25 Å.^{20,21} No internal order was evidenced.

Here, we report on the encapsulation of MPc (free-base (H₂) and metallated (Zn^{II} and Fe^{II})) into single-walled carbon nanotubes. These nanotubes display several advantages such as metallic or semiconducting properties and diameters (~14.5 Å) hardly larger than the molecule size. Thus, the molecules are expected to perfectly fit in the hollow core of NTs, allowing important confinement effects. We clearly evidence a 1D structure, showing that carbon NTs strongly influence the long range structural properties of MPc. The physical interactions in the hybrid system are discussed and we show that they are dominated by molecule-molecule interaction. For comparison, functionalization at the outer surface of the tube is also explored. In this case, molecule-nanotube interactions are dominant, affecting the local structure of the phthalocyanine. Two different single-walled carbon nanotubes are studied, exhibiting average diameters of 14.5 Å (NT14) and 17 Å (NT17).

EXPERIMENTAL METHODS:

Iron, Zinc and metal free phthalocyanine molecules (respectively FePc, ZnPc and H₂Pc) are purchased from Sigma Aldrich (*ar* for as received). They are purified (*pf*) by a temperature treatment at 500°C under dynamic high vacuum. The purification step consists in heating the molecules at their sublimation temperature. The objective is twofold: i) purification of the sample by separating the molecules from impurities and ii) the molecule after the purification step has undergone the same temperature treatment as for the encapsulation protocol (see below) and it is thus similar to the encapsulated ones. Consequently, it can be used as reference

with respect to the confined one. The different samples are named MPc_{ar} and MPc_{pf} (M=Fe, Zn or H₂). Single walled carbon nanotubes are either purchased from Carbon Solution Company or supplied by National Institute of Advanced Industrial Science and Technology (AIST), Japan. They are synthesized by the enhanced direct injection pyrolytic synthesis (eDIPS) method.²² The nanotubes of the two different sources display average diameter of 1.4±0.2 nm and 1.7±0.2 nm respectively. Therefore, the samples will be named respectively NT14 and NT17 in the following. As received (ar) closed nanotubes are used for the outer surface functionalization (π -stacking). Purified and annealed (pf) open nanotubes are used for molecule encapsulation into the hollow core of nanotubes. Thus, the different samples are named NT14_{ar}, NT14_{pf}, NT17_{ar} and NT17_{pf}.

Synthesis of the hybrid system: A mixture of nanotubes and MPc within ratio 1:2 was outgassed under 6×10^{-6} mbar at ambient temperature for 2 h followed by a thermal treatment at 200 °C. The glass-tube was then sealed and heated homogeneously in a furnace at 500°C during 24 hours. After this treatment, excess of phthalocyanine was removed by outgassing at 500°C at 6×10^{-6} mbar. Samples were then washed five times with N-Methyl-2-Pyrrolidone (NMP). As received nanotubes give rise to non-covalent functionalization at the outer surface of the nanotubes (MPc/NT_{ar}) whereas purified nanotubes lead to the encapsulation process (MPc@NT_{pf}). NT can be either NT14 or NT17.

Transmission electron microscopy and spatial-resolved electron energy loss spectroscopy:

The transmission electron microscopy (TEM) samples were prepared by dispersing the NTs powders in ethanol. The dispersions were ultrasonicated and subsequently deposited on holey carbon 3 mm copper grids.

High resolution TEM was performed employing an imaging-side aberration-corrected FEI Titan-Cube microscope working at 80 keV, equipped with a Cs corrector (CETCOR from CEOS GmbH).

Spatial-resolved electron energy loss spectroscopy (EELS) measurements were performed on probe-corrected scanning TEM (STEM) FEI Titan Low-Base 60-300 operating at 80keV (fitted with a X-FEG® gun and Cs-probe corrector (CESCOR from CEOS GmbH)). Furthermore, in order to avoid the effects of electron beam damage, these measurements have been performed using a liquid-nitrogen-cooled cryo-holder at -170⁰ C. EEL spectra were recorded using the spectrum-imaging (SPIM in 2D or spectrum-line (SPLI) in 1D) mode^{23,24} in a Gatan GIF Tridiem ESR 865 spectrometer. The convergent semi-angle was of 25 mrad, the collection semi-angle was of 35 mrad and the energy resolution ~ 1.2 eV.

X-ray diffraction: Small angle X ray scattering (SAXS) and wide angle X ray scattering (WAXS) experiments were performed with an in-house setup of the *Laboratoire Charles Coulomb, "Réseau X et gamma", Université Montpellier, France*. A high brightness low power X- ray tube, coupled with aspheric multilayer optic (GeniX^{3D} from Xenocs) was employed. It delivers an ultralow divergent beam (0.5mrad). Scatterless slits were used to give a clean 0.8 mm beam diameter (35 Mphotons/s) at the sample. We worked in a transmission configuration and scattered intensity was measured by a Schneider 2D imageplate detector prototype, at a

distance of 1,9 m from the sample for SAXS configuration, 0.2m from the sample for WAXS configuration. All Intensities were corrected by transmission and the empty cell contribution was subtracted.

The calculated diffraction pattern features bundles of 16 NTs with mean diameter of 1.7 nm. A diameter distribution of FWHM 0.17 nm was used. The nanotubes are filled with balls of diameter 0.7 nm and containing 290 electrons, mimicking Fe-phthalocyanines rotating around the long axis of carbon nanotubes. The balls were spaced of a distance compatible with the position observed for the diffraction peak that emerges in the pattern of FePc@NT17_{pf}. We chose to consider the inserted complexes as balls of smaller diameter than the complex for 2 reasons. First, the electron distribution is not homogeneous in the FePc complex, and is concentrated near the complex center. Second, we assume a rotation around the SWNT axis, which is the only direction possible considering the steric hindrance imposed by the surrounding nanotube. For such a rotation, and considering the symmetry of the electron distribution in the FePc complex, the mean electron distribution can be represented by a ball containing all the electrons of complex, i.e. 290 electrons.

Raman spectroscopy: Micro-Raman experiments have been performed in a back scattering geometry, using the fixed 633, and 660 nm excitation wavelengths. In order to prevent the heating of the tubes and oligomers, the laser power was adjusted at 100 μ W with a spot diameter of about 3 μ m using a 50X objective. The resolution is about 2 cm^{-1} . At least four spatially separated area of each sample were probed to ensure homogeneity. The precision in the peak position measurements is $\pm 1 \text{ cm}^{-1}$.

Computational details : The calculations were performed in the framework of the density functional theory (DFT) as implemented in the SIESTA code.²⁵ Two models have been studied: (i) an encapsulation model where we considered the guest molecule inside a (22,0) nanotube of 12.78 Å length and a diameter of 17.22 Å, and (ii) a π -stacked model where the guest molecule is π -stacked on a (17,0) or (22,0) nanotube of 29.82 Å length and a diameter of 13.31 and 17.22 Å, respectively. The periodic conditions were applied along the tube axis and the vacuum size was of 20 Å in the two other directions to avoid interactions between adjacent tubes.

Here, the guest molecules are metal-free phthalocyanine (H₂Pc) and zinc phthalocyanine (ZnPc). The atomic positions were relaxed at fixed lattice parameters using a conjugate gradient until the maximum residual force on the atoms was smaller than 0.02 eV/Å. Exchange-correlation effects were handled within the generalized gradient approximation (GGA) as proposed by Perdew, Burke and Ernzerhof.²⁶ Core electrons are replaced by nonlocal norm-conserving pseudopotentials.²⁷ A double zeta singly polarized basis set (DZP) of localized atomic orbitals was used for the valence electrons.²⁸ Their cutoff radii were determined from an energy shift of 50 meV by localization. A mesh cutoff of 300 Ry was used for the reciprocal-space expansion of the total charge density and the Brillouin zone was sampled using three k points along the nanotube axis. Van der Waals corrections between the nanotubes and the guest molecules were considered according to the Grimme approach.²⁹ This approach (DFT-D) consists in adding a semi-empirical dispersion potential (a simple pair-wise force field) to the conventional Kohn-Sham DFT energy.

RESULTS AND DISCUSSION:

Transmission electron microscopy: Figures 1.a corresponds to a high resolution (HRTEM) micrograph of single-walled nanotubes after the encapsulation step. All the nanotube seem filled with a very good efficiency. However, it is worth mentioning that from this kind of HRTEM micrographs it is not possible to deduce the nature of the confined material, neither to obtain any chemical information. Thus, in order to go further in the analysis of these materials and to know where the MPc molecules are localized, we have developed SR-EELS. This technique is a very powerful tool to investigate such nanomaterials at (sub-) nanometer scale.³⁰⁻³⁴ Structural and chemical properties of bundle of single-walled nanotubes have also been investigated. We have acquired a spectrum-image (SPIM) in the whole area showed in the high angle annular dark field (HAADF) image (Figure 1.b).

Figure 1.c displays two EEL spectra extracted in the blue and red areas marked on Figure 1.b, respectively. Each of these two EEL spectra corresponds to the sum of 9 spectra (3x3 probe positions of the SPIM). Carbon K edge at ~285 eV is present in both of these two EEL spectra.

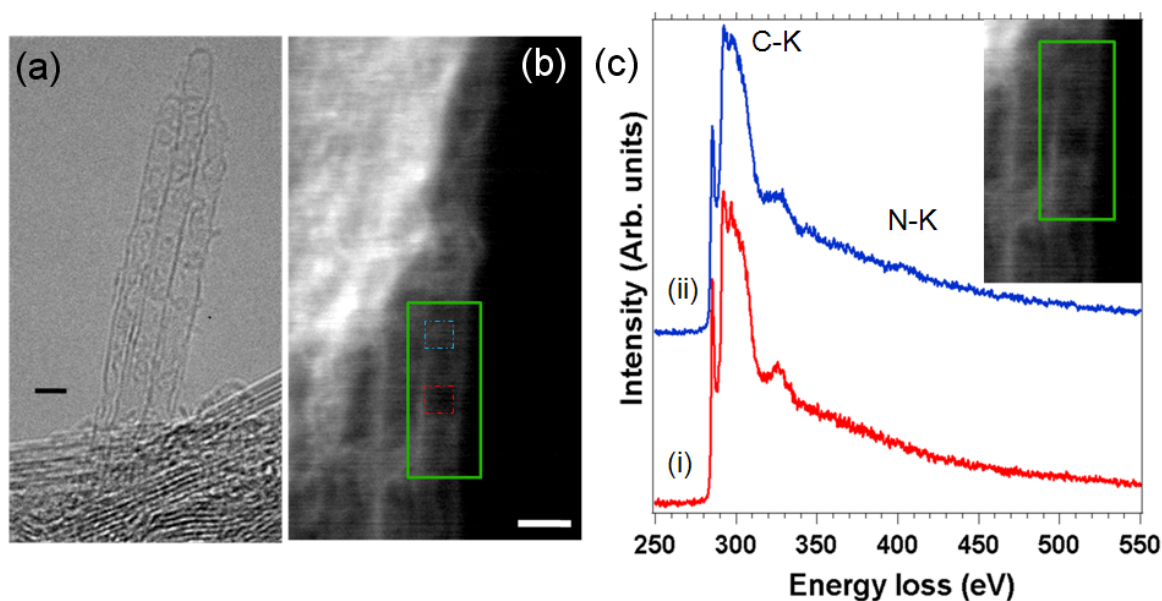


Figure 1: (a) HRTEM micrographs of bundles filled single-walled nanotubes. (b) HAADF image of another bundle of filled single-walled nanotubes. A 7x16 EELS SPIM has been recorded in the marked green area of this micrograph. (c) Sums of 9 selected EEL spectra collected from the two different marked areas in the SPIM, marked in Fig. 1.b. While the C-K edge is visible in both spectra, the N-K edge is only visible in the blue (Figure 1 (c-ii)) EEL spectrum acquired in the blue area of the HAADF image (Figure 1.b). The insert of this figure is a zoom of the HAADF image displayed in (b). The scale bars of the two micrographs (Figure 1.a and b) are 2 nm.

This C-K edge shows the characteristic peaks of sp^2 materials at ~285 eV and ~292 eV, corresponding to the π^* and σ^* contributions, respectively.³⁰⁻³² The most significant point

concerning these EELS results is that in the spectrum acquired over the blue region (Figure 1b), we additionally detect nitrogen (see Figure 1.c-ii). This nitrogen is clearly associated to FePc, as it is the only source of nitrogen in these samples. Thus, from the combination of the HRTEM images and local analytical findings, we conclude that these observed moieties correspond to the FePc, which are localized in the hollow core of NTs. However, these investigations do not allow to determine the long-range organization of the molecules confined into nanotubes.

X-ray diffraction: The x-ray diffraction experiments are first used to investigate the structural organization of the confined species into the hollow core of the nanotubes but also to probe the encapsulation efficiency of the phthalocyanine molecules. Indeed the main peak (referred as 10 peak) arising from the two-dimensional structural organization of the nanotubes into bundles is well known to vanish under encapsulation.^{35,36} Independently of the central atom, phthalocyanine molecules can organize into two main structures, respectively called the α and the β phases.^{2,37-46} The α structure is generally reported as metastable whereas the β one is thermally stable over a large temperature range.³⁷⁻⁴⁰ Both phases are stacked molecules forming columns with the ring tilted with respect to the vertical axis.^{2,41,45,47}

Figure 2 exhibits the diffraction pattern of raw and functionalized NT14 and NT17 nanotubes, the α and β FePc phases. The 10 peak of the nanotube bundles is located at $5.85 \pm 0.03^\circ$ for raw and purified NT14 and at $4.57 \pm 0.03^\circ$ for purified NT17. These positions correspond respectively to inter reticular distances of 15.1 ± 0.1 and 19.3 ± 0.15 Å, in good agreement with the tube diameter distributions. After outside functionalization (FePc/NT_{ar}), the peaks are shifted towards lower angle, respectively $5.7 \pm 0.03^\circ$ for NT14 and $4.2 \pm 0.03^\circ$ for NT17 (not shown). This modification could be explained by a very weak dilatation of the bundles. Another explanation is to assign the shift to form factor interferences as already described for iodine intercalation into nanotubes.³⁵ Anyway, the presence of the 10 peak clearly indicates that, as expected the nanotubes are still mainly empty after outside functionalization.

After FePc encapsulation, the 10 peaks are no longer observable for FePc@NT14 and FePc@NT17, confirming the nanotube filling, which is also evidenced by SR-EELS results. Furthermore, a new broad peak is observed below 7° for both samples. It turns out that both the α and β FePc phases exhibit a diffraction peak located respectively around 6.9° and 7° (figure 2), and corresponding to inter-reticular distances between the stacking axes, normal to the phthalocyanine rings.³⁹ The presence of only one peak around 6.9° is generally associated to the formation of an α -phase for FePc,^{42,44} ZnPc^{40,45,46,48} and H₂Pc⁴⁷ films or crystals and corresponds to an intermolecular distance along the [200] direction. Thus, the new peak features the structural organization of the FePc molecules inside the carbon nanotubes.

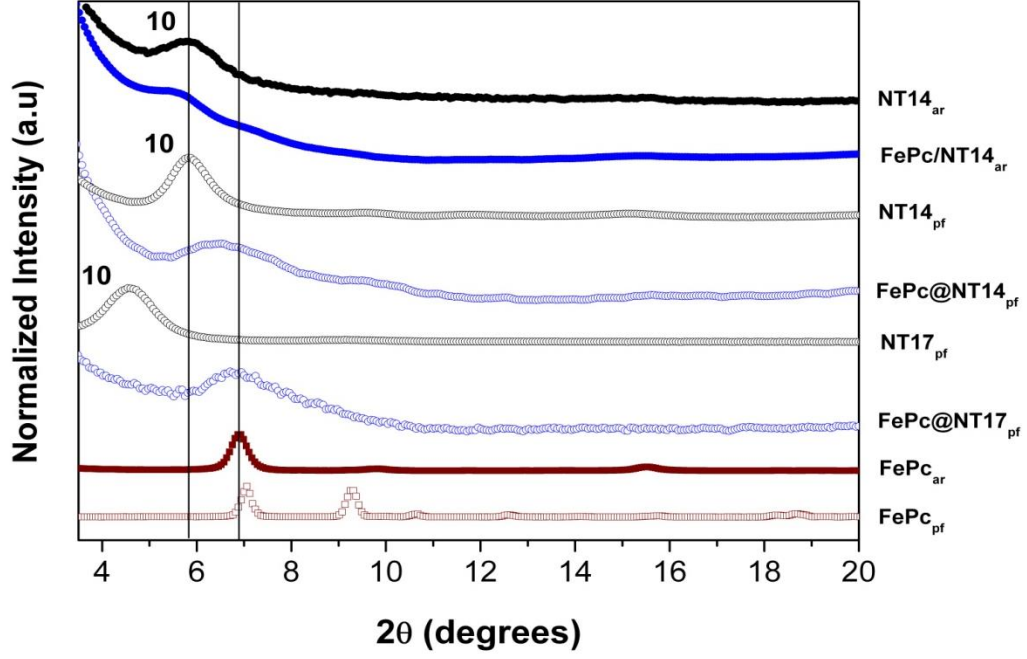


Figure 2: X-ray diffraction pattern recorded with a X-ray wavelength of 1.542 Å. (from top to bottom) as prepared NT14 (black full circle), FePc/NT_{ar} (full blue circle), purified NT14 (open black circle), FePc@NT14_{pf} (open blue circle), purified NT17 (open black circle), FePc@NT17_{pf} (open blue circle), α-phase FePc (full wine square), β-phase FePc (open wine square). The intensities are adjusted for clarity.

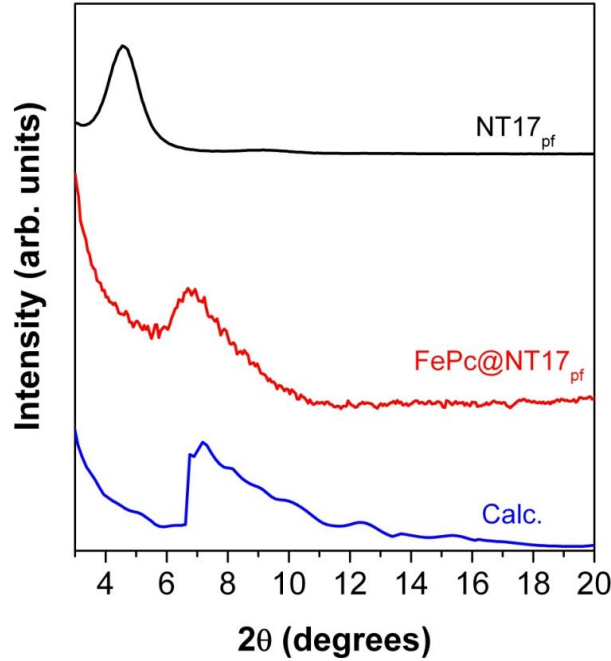


Figure 3: Experimental x-ray diffraction pattern of the reference NT17_{pf} powder (*top*) and the FePc@NT17_{pf} powder (*center*). Calculated x-ray diffraction pattern of a powder of filled NTs of diameter 1.7 nm, with a diameter distribution of FWHM 0.17 nm, (*bottom*; see experimental section for details).

To confirm our assumption on the structure of Fe-phthalocyanines inside the nanotubes, a numerical calculation of the diffraction pattern of the FePc@NT17 sample is shown on figure 3 and compared to the experimental profile. The formalism is the same as the one used in our

previous calculations of C₆₀ peapods⁴⁹ and iodine confined in NTs.⁵⁰ The intensity calculated within this model gives a very nice reproduction of the experimental diffraction profile. The 10 peak has disappeared because of the interference effects already discussed above. In the same time, an asymmetric peak of saw-tooth shape has grown, testifying for the 1D nature of the FePc stacking in NTs. Consequently, the new peak around 6.85° is definitively assigned to the FePc ordering inside the nanotubes. This supramolecular organization under nano-confinement is suggested to be a 1D array of molecules lying into the nanotubes. According to the Scherrer's equation, the mean crystallite size is about 5 nm, meaning an average cluster of around 4 molecules. It is worth mentioning that those results also stands for ZnPc and metal free molecules encapsulated into nanotubes.

To go further, we performed DFT-calculations to determine the molecule orientation with respect to the nanotube axis. With an intermolecular distance set at 12.8 Å, derived from reference⁴⁶ and in good agreement with our experimental data, the minimum of the bonding energy is obtained for an angle between the molecular ring and the nanotubes axis calculated at 32° as shown on figure 4. Thus, the corresponding stacking angle as defined in reference is 58°, quite different from the 36° proposed for multi-walled nanotubes.²⁰

Thus, nano-confinement inside single-walled nanotubes gives rise to a 1D supramolecular organization of the MPc molecules monitored by the small NT diameter.

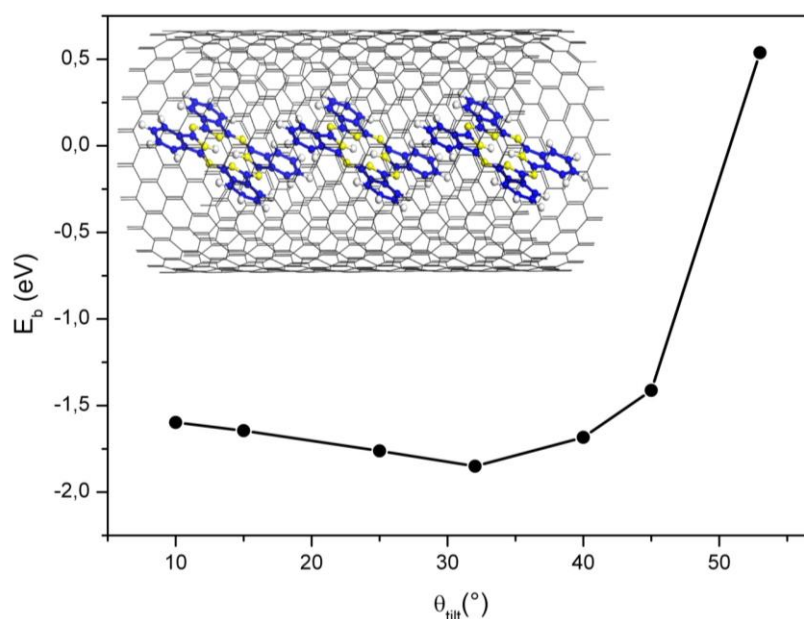


Figure 4 : Binding energy as a function of the tilt angle of the Pc molecule with respect to the nanotube (22,0) axis **Inset**: Scheme of the Pc structural organization inside single-walled carbon nanotubes (22,0)

Raman experiments:

The Raman spectra of the hybrid systems exhibit an important dependence with the excitation energy which is correlated to the optical absorption energy of the photo-active molecule.⁵¹ Figure 5 exhibits the Raman spectra recorded with an excitation wavelength of 633 nm of (a) the different purified phthalocyanine molecules MPc, (b) endohedral NT14 (MPc@NT14), and

(c) NT14 functionalized at their outer surface (MPc/NT14). The intensities are normalized either on the G-band peak or on the most intense peak of the MPc Raman signal.

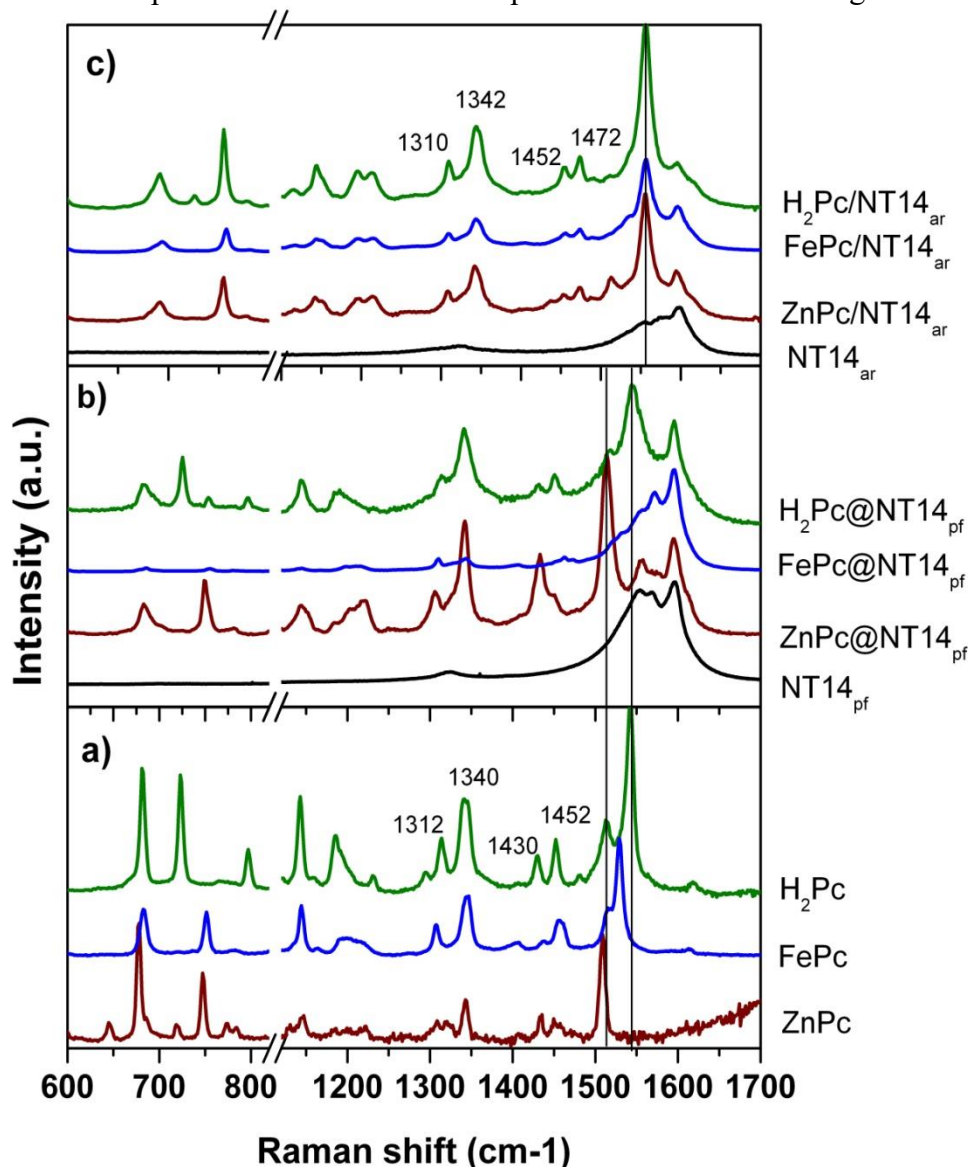


Figure 5: Normalized Raman spectra recorded with an excitation wavelength of 633 nm a) purified phthalocyanine molecules MPc -a strong background had to be subtracted- (b) purified NT14_{pf} and endohedral functionalized NT14_{pf} (MPc@NT14_{pf}) c) as received NT14_{ar} and surface functionalized NT14_{ar} (MPc/NT14_{ar})

Figure 5.a clearly shows that the central atoms display clear fingerprints in the Raman spectra as both profiles and peak positions change depending on the type of the MPc molecules (respectively between 600 and 800 cm⁻¹ and between 1400 and 1600 cm⁻¹). The peaks around 680 and 750 cm⁻¹ mainly correspond to breathing and deformation of the macrocycle whereas those lying between 1500 and 1550 cm⁻¹ involve mainly C-N-C stretching.⁵²⁻⁵⁴ All of them concern vibrations around the metal central atom, accounting for the difference in the Raman spectra for the three different phthalocyanine molecules. Most of the peaks between 1000 and 1500 cm⁻¹ involves mainly in-plane deformation and they are much less affected by the central atom size.⁵²⁻⁵⁶

Figure 5.b displays the Raman spectra recorded of NT14_{pf} nanotubes and after encapsulation (MPc@NT14_{pf}). The nanotube G-band exhibits a Breit-Wigner-Fano profile, characteristic of metallic nanotubes and in good agreement with the Kataura plot. The photoluminescence background of MPc is no longer observable after encapsulation into nanotubes and a Raman signal is observed, demonstrating the bleaching of the MPc photoluminescence effect. Nevertheless, the Raman spectrum of the hybrid system consists roughly in the superposition the Raman spectra of the nanotube and the confined molecule. These data strongly suggests that the encapsulated MPc behaves as a “free” molecule inside the nanotubes.

Figure 5.c displays the Raman spectra of the as-received NT14 nanotubes and after noncovalent functionalization at their outer surface with phthalocyanine molecules with different central atoms (MPc/NT14). The spectra no longer consist in the superposition of the MPc molecule and the nanotube Raman responses, indicating a significant interaction between both systems. In all cases, the profiles of the Raman spectra seem very similar, independently of the central atom, meaning that its Raman fingerprint cannot be observed anymore. These results strongly suggest that the central atom does not play any important role in the molecule vibrations. However, those Raman spectra do not match the Raman spectrum of the bulk H₂Pc molecules (figure 5) meaning that another effect has to be taken into account. For instance, the most intense peak is localized at 1556 cm⁻¹ for all MPc@NTs but shifted (14 cm⁻¹) compared to H₂Pc. Furthermore, important discrepancies are observed in the low frequency domain (below 800 cm⁻¹) where out-of-plane macrocycle vibrations are probed. Our hypothesis is to consider that the MPc molecules π -stacked at the outer surface fit the nanotube curvature, so that their structures are no longer flat. As a consequence, the central atom is out-of-plane and no longer contributes to the molecule vibrations. This assumption is strongly supported by the important shift of the peaks at 1430 and 1452 cm⁻¹ (respectively 1452 and 1472 cm⁻¹) for the bulk H₂Pc (respectively H₂Pc stacked onto NT). Indeed, those peaks arise from the vibrations of carbon atoms within the isoindole group⁵⁷ (localized near the molecule edge) and are thus strongly affected by the phthalocyanine curvature. Consequently, the differences in the Raman spectra of H₂Pc and H₂Pc/NT14 are assigned to the H₂Pc ring curvature due to the interaction with the nanotubes walls.

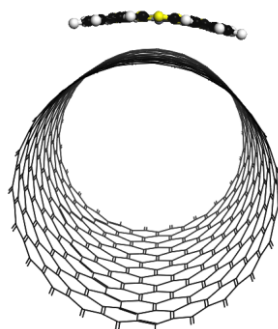


Figure 6: Structure of the ZnPc stacked at the outer surface of NT (17,0) from our DFT calculations.

This assumption is consistent with DFT calculations (figure 6), showing that the ring molecules switch from an angle of 180° (flat structure) to an angle of 154° (curved structure on the

nanotubes) for a (17,0) nanotube. In addition, taking van der Waals interactions into account, the central atom (Zn in our model) moves slightly towards the nanotube wall, being thus out of the molecule plane (figure 7). Let's consider the curved ZnPc stacked onto the nanotubes (up-left corner of figure 7) and only some of its atoms (those inside a black open circle on the up-right corner). Figure 7 displays the projection of those peculiar atoms onto the (xy) plane. The Zn atom is slightly but clearly shifted upon interaction with the tube. Thus, we assume that the central atom no longer plays a role in the molecule vibration, accounting for our Raman data.

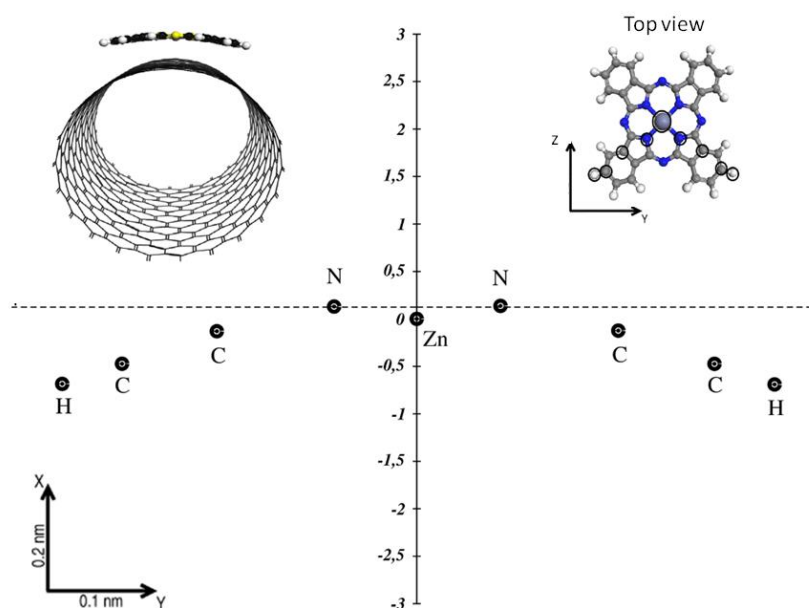


Figure 7: Projection onto the (x,y) plane of atoms of the ZnPc molecule stacked onto a nanotube .

To go further, we also studied hybrid systems with carbon structures displaying different curvatures (graphite, as received NT17 and NT14 nanotubes). Figure 8 exhibits the Raman spectra of bulk FePc and molecules deposited on these different carbon structures. As previously observed, stacking on NT14_{ar} gives rise to important changes in the FePc Raman response (peak shifted from 1510 to 1540 cm⁻¹ and splitting of the peak around 1450 cm⁻¹). By contrast, the FePc vibrations do not look altered by the interaction with graphite or NT17_{ar}. One can assume that the curvatures of the host carbon materials are not large enough to induce significant modifications of the FePc structure (remaining flat), preventing any change in the Raman vibrations.

One can notice that according to the Kataura plot, NT14 and NT17 nanotubes are respectively metallic and semiconducting for a 660 nm excitation wavelength (whereas graphite is a semi-metal). As similar results are obtained for both semiconducting and metallic NT17 with excitation wavelengths of respectively 660 and 785 nm, we assume that the electronic properties do not play significant role in the interaction and that the nanotube diameter is the relevant parameter.

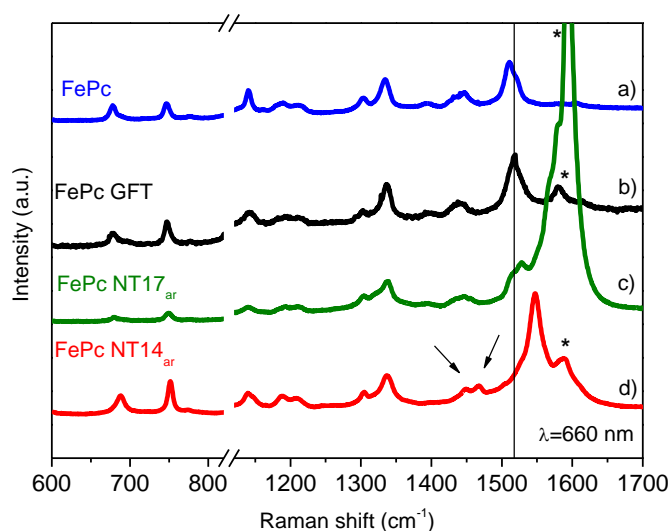


Figure 8: Raman spectra recorded with an excitation wavelength of 660 nm a) purified phthalocyanine molecules MPc (b) FePc/Graphite c) FePc/NT17_{ar} d) FePc/NT14_{ar} * Indicates the the G-band of the host carbon materials

Conclusions:

Phthalocyanine molecules have been noncovalently functionalized inside or at the outer surface of single-walled carbon nanotubes. HRTEM and SR-EELS analyses have confirmed that the MPc molecules are located inside of nanotubes. The structural 1D organization of the molecules inside the nanotubes is determined by x-ray diffraction coupled to DFT calculations. The MPc molecules are stacked along the tube axis, with an angle of 32°, giving rise to a 1D phase under nano-confinement. The Raman data suggest only a very weak interaction when the molecules are confined into the nanotubes. By contrast, localized at the outer surface, a strong interaction takes place. The strength of this last interaction depends on the nanotube diameter and induces a significant curvature of the molecule ring when decreasing the NT diameter. As a consequence, the central atom (M) gets out of plane, impacting the molecule vibration properties. These results demonstrate that carbon nanotubes can play the role of nano-support for organic molecules of interest in opto-electronic applications. As a matter of facts, they can guide their linear organization and can even bend them to an extent defined by the nanotubes radius. These results open exciting perspectives for the development of new hybrid systems with tunable optical properties.

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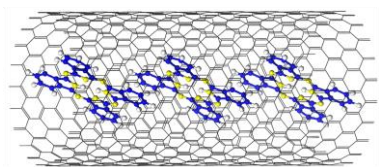


Figure 1: (a) HRTEM micrograph of bundles of filled single-walled nanotubes. (b) HAADF image of another bundle of filled single-walled nanotubes. A 7x16 EELS SPIM has been recorded in the marked green area of this micrograph. (c) Sums of 9 selected EEL spectra collected from the two different marked areas in the SPIM, marked in Fig. 1.b. While the C-K edge is visible in both spectra, the N-K edge is only visible in the blue (Fig. 1 (c-ii)) EEL spectrum acquired in the blue area of the HAADF image (Fig. 1.b). The insert of this figure is a zoom of the HAADF image displayed in (b). The scale bars of the two micrographs (Fig. 1.a and b) are 2 nm.

Figure 2: X-ray diffraction pattern recorded with a X-ray wavelength of 1.542 Å. (from top to bottom) as prepared NT14 (black full circle), FePc/NT_{ar} (full blue circle), purified NT14 (open black circle), FePc@NT14_{pf} (open blue circle), purified NT17 (open black circle), FePc@NT17_{pf} (open blue circle), α -phase FePc (full wine square), β -phase FePc (open wine square). The intensities are adjusted for clarity.

Figure 3: Experimental x-ray diffraction pattern of the reference NT17_{pf} powder (*top*) and the FePc@NT17_{pf} powder (*center*). Calculated x-ray diffraction pattern of a powder of filled NTs of diameter 1.7 nm, with a diameter distribution of FWHM 0.17 nm, (*bottom*; see experimental section for details).

Figure 4: Binding energy as a function of the tilt angle of the Pc molecule with respect to the nanotube (22,0) axis **Inset:** Scheme of the Pc structural organization inside single-walled carbon nanotubes (22,0)

Figure 5: Normalized Raman spectra recorded with an excitation wavelength of 633 nm a) purified phthalocyanine molecules MPc -a strong background had to be subtracted- (b) purified NT14_{pf} and endohedral functionalized NT14_{pf} (MPc@NT14_{pf}) c) as received NT14_{ar} and surface functionalized NT14_{ar} (MPc/NT14_{ar})

Figure 6: Structure of the ZnPc stacked at the outer surface of NT (17,0) from our DFT calculations.

Figure 7: Projection onto the (x,y) plane of atoms of the ZnPc molecule stacked onto a nanotube .

Figure 8: Raman spectra recorded with an excitation wavelength of 660 nm a) purified phthalocyanine molecules MPc (b) FePc/Graphite c) FePc/NT17_{ar} d) FePc/NT14_{ar} * Indicates the the G-band of the host carbon materials