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^1H and ^{195}Pt NMR prediction for inclusion compounds formed by cisplatin and oxidized carbon nanostructures†

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Prediction of NMR chemical shifts can assist experimentalists in the characterization of drug delivery systems based on carbon nanocomposites. Chemical shifts are strongly correlated to the nucleus position and its chemical neighborhood. Therefore, to predict structures and NMR properties of complex chemical models, choosing a more consistent theoretical level capable of providing more realistic results and moderate computational demand is a major challenge. In this work, we predicted the NMR spectra of inclusion compounds formed by cisplatin (cDDP) and an oxidized carbon nanotube (CNTox) and nanocone (CNCox) considered by specialists as potential drug delivery systems. The ^{195}Pt NMR chemical shifts calculated at the DFT level with the new relativistic NMR-DKH basis set were -2314 ppm and -2192 ppm for cDDP@CNTox and cDDP@CNCox complexes, respectively, which are both high-field shifted relative to the free cDDP (-2110 ppm). ^1H NMR chemical shifts are also sensitive to the inclusion process. The H (NH_3) signals are found on average at $+4.3$ (cDDP), -5.1 (cDDP@CNTox) and $+6.6$ ppm (cDDP@CNCox). Interestingly, despite the similar inclusion modes in CNTox and CNCox cavities, the ^1H NMR shifts were in opposite directions. A possible reason might be the higher stability of cDDP@CNTox ($\Delta E_F = -19.9$ kcal mol $^{-1}$) than that of cDDP@CNCox ($\Delta E_F = -5.7$ kcal mol $^{-1}$), which suggests a short guest–host contact in the former and consequently, a more efficient shielding of hydrogen atoms due to the electron-rich carbon structure. These results may be helpful as comparison data in the NMR spectra assignment in solution and the inclusion compounds' structural elucidation.

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Introduction

Antitumor agent cisplatin¹ and its derivatives,^{2,3} when used alone or in combination with other chemotherapeutics^{4,5} can be effective against the advancement of several clinical cases of cancer. However, there are several known factors such as low selectivity, resistance and consequently, high toxicity that are limiting for their clinical use.⁶ Thus, alternative formulations of

platinum compounds have been studied, in particular, drug delivery systems (DDSs) formed by organic macromolecules,^{7–9} liposomes,^{10,11} nanoparticles^{12,13} and oxidized carbon nanocomposites.^{14–20} Once in the bloodstream, it is expected that these systems are able to protect the drug against undesirable side reactions as well as provide efficient release in a controlled biological target. Therefore, the understanding of how these systems are provided as inclusion compounds can contribute to design improved complexes with desirable properties.

Nuclear magnetic resonance (NMR) spectroscopy is an important tool for investigating the structural and dynamic aspects of many bioactive compounds. Ravera *et al.*²¹ showed the synthesis and characterization by ^1H , ^{13}C and ^{195}Pt NMR of Pt(IV)–bis(benzoato) complexes with potential anticancer activity. The authors observed that the lipophilicity, cellular accumulation and cytotoxicity on chemoresistant MPM cell lines were considerably increased in relation to their Pt(II) analogs, cisplatin and oxaliplatin. However, it was observed that the structure–activity relationship of Pt(IV) complex derived-oxaliplatin is possibly affected by the interaction of the aromatic ligand with intracellular components. In order to minimize these undesirable reactions, the authors studied the

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inclusion of the drug in cyclodextrins (CDs) and the inclusion complex formed was characterized by ^1H NMR spectra. The results showed that the signals of protons H-3, H-5, and H-6, located inside the β -CD cavity, are shifted, supporting the identification of inclusion compound. Levet *et al.*²² developed a dry powder type formed by liposomes and/or PEGylated nanoparticles with cisplatin for controlled release of the drug after inhalation during adjuvant chemotherapy in the treatment of lung cancer. The integrity and quantification of cisplatin, reduced to submicron sizes, were assessed by electrothermal atomic absorption spectrometry (ETAAS) and ^{195}Pt NMR spectroscopy. Through the integration of the ^{195}Pt NMR signal, it was observed a slightly larger amount of the drug in the final product compared to the amount found by the ETAAS technique.

Our research group^{23–27} has reported theoretical calculations of structure, stability and characterization by ^1H , ^{13}C and ^{15}N NMR spectroscopy of inclusion nanocomposites. Recently, we used density functional theory (DFT) methods to study inclusion complexes formed between cisplatin and oxidized carbon nanotube and nanocone molecules.²⁶ Our results showed through the comparison between the B3LYP and B97-D density functional that the dispersion correction significantly improves the stabilization energy and thermodynamic parameters of the compounds studied. ^1H NMR spectra calculations in aqueous solution showed that experimental detection of the inclusion compounds can be promptly attained through analysis of chemical shifts for cisplatin NH_3 protons, with variations around -12 and -5 ppm due to the formation of complexes, compared to the free cisplatin molecule. Hosni *et al.*²⁰ used the gauge invariant atomic orbitals (GIAO) method to calculate the ^{195}Pt chemical shielding tensors in inclusion complex models formed by various types of carbon nanotube and cisplatin. The authors used the Hartree–Fock (HF) method with aug-cc-pVDZ-PP and STO-3G basis set for the Pt and H, C, N, Cl atoms, respectively, for the geometries optimization and ^{195}Pt NMR chemical shifts calculations in gas phase. Their results showed that the $\delta^{195}\text{Pt}$ is able to inform about the CNT diameter and also to lead directly to confinement energy of drug. A decreasing variation from 533 to 38 ppm were observed for the $\delta^{195}\text{Pt}$ of the drug into nanotubes models with approximately 6 to 12 Å in diameter. Even though ECP was used for Pt in ref.²⁰, it is of great importance for the computational study proposed for this type of system. Moreover, to date, this has been one of the most current studies in attempt to perform theoretical calculations of ^{195}Pt NMR chemical shifts in inclusion compounds formed by carbon nanotubes and cisplatin.

Theoretically, the accurate prediction of NMR parameters involving heavy nuclei must take into account several aspects such as structure, solvent and relativistic effects, basis sets and electronic correlation, as can be seen in the works of Vicha *et al.*,^{28,29} Truflandier *et al.*³⁰ and Pawlak *et al.*³¹ However, to study very large systems, such as carbon nanocomposites, the inclusion of relativistic effects can become very expensive and the use of non-relativistic protocols might be viewed as an affordable alternative. In this sense, Paschoal *et al.*³² performed non-relativistic and relativistic calculations to analyze the ^{195}Pt

NMR chemical shifts of 258 Pt(II) complexes, including the anticancer drug cisplatin. The authors proposed a protocol that includes all-electron Gaussian basis sets to describe the $\delta^{195}\text{Pt}$ and all elements commonly found as Pt-ligands. The new basis sets, named NMR-DKH, were partially contracted as a triple-zeta doubly polarized with all coefficients obtained from a Douglas–Kroll–Hess (DKH) second-order scalar relativistic calculation. Accordingly, the chemical shifts of ^{195}Pt were calculated through empirical models fitted to reproduce experimental NMR data ranging from -1000 to -6000 ppm. Among other theoretical protocols proposed, the best one employed the B3LYP/LANL2DZ/def2-SVP/IEF-PCM(UFF) level for the geometry optimization and PBEPBE/NMR-DKH/IEF-PCM(UFF) level for $\delta^{195}\text{Pt}$ calculation. The authors obtained results comparable with relativistic DFT calculation with the mean absolute deviation (MAD) and the mean relative deviation (MRD) around to 168 ppm and 5%, respectively. Very recently, Carvalho *et al.*³³ applied the same protocol to predict the $^1J(^{195}\text{Pt}-^{15}\text{N})$ coupling constant in 71 Pt(II) complexes. For a total of 98 coupling constants, the authors found a MAD and MRD of only 36 Hz and 10.4%, respectively, showing the quality of the NMR-DKH basis set and theoretical protocol employed.

In the present work, we used the very same protocol proposed by Paschoal *et al.*³² for prediction of the ^{195}Pt NMR spectrum for the inclusion complexes formed between cisplatin (cDDP) and oxidized carbon nanotubes (CNTox)/nanocones (CNCox) as potential DDS. Our results provide relevant spectroscopic data of ^1H and ^{195}Pt NMR that can assist experimentalists interested in the preparation and characterization of these systems.

Theoretical methodology

The calculations were carried out using the GAUSSIAN 09 package.³⁴ The methodology used for the prediction of the NMR chemical shifts follows the computational protocol proposed by Paschoal *et al.*³² The geometries of the cDDP@CNTox and cDDP@CNCox inclusion complexes were fully optimized at DFT³⁵ B3LYP³⁶ level using the LANL2DZ³⁷ effective core potential for Pt atom and the def2-SVP³⁸ basis set for the other atoms. The solvent effect was accounted for within the continuum aqueous solution approach using the IEF-PCM³⁹ formalism with dielectric constant adjusted for water solvent ($\epsilon = 78.3553$) and UFF atomic radii, namely, B3LYP/LANL2DZ/def2-SVP/IEF-PCM(UFF) level. The complexes formation energies (ΔE_{F}) were calculated as

$$\Delta E_{\text{F}} = E_{\text{complex}} - (E_{\text{CNox}} + E_{\text{cDDP}}) \quad (1)$$

where E_{complex} , E_{CNox} and E_{cDDP} correspond to the total energy of the molecular complex, free oxidized carbon nanostructures and cisplatin molecules fully optimized structures, respectively.

The NMR spectra of inclusion complexes were calculated using the GIAO⁴⁰ method at DFT level with the GGA PBE^{41,42} functional with the NMR-DKH³² basis set, namely, PBEPBE/NMR-DKH/IEF-PCM(UFF) level, with the fully optimized geometries at B3LYP/LANL2DZ/def2-SVP/IEF-PCM(UFF) level.



For the ^1H and ^{15}N NMR chemical shifts, the tetramethylsilane (TMS) and NH_4Cl compounds were used as internal reference, respectively. NMR chemical shifts were obtained by eqn (2), where the σ_{ref} is the calculated shielding constant for a reference compound and the σ_{calc} is the shielding constant calculated for the nuclei of interest.

$$\delta_{\text{calc}} (\text{ppm}) = \sigma_{\text{ref}}(\text{TMS}) - \sigma_{\text{calc}} \quad (2)$$

The NMR chemical shift for ^{195}Pt nucleus was calculated using eqn (3), where the σ_{calc} is the ^{195}Pt shielding constant calculated with the NMR-DKH basis set, as proposed by Paschoal *et al.*³²

$$\delta_{\text{calc}} (\text{ppm}) = -2065.7558 - 0.9250\sigma_{\text{calc}} \quad (3)$$

The explicit solvent effect on the calculation of NMR chemical shifts was also investigated by analyzing a set of 20 solvated structures generated from molecular dynamics (MD) simulations of the cDDP@CNTox inclusion complex. Regarding the parameterization of the CNTox model, while both intramolecular and Lennard-Jones parameters (LJ 12-6) were selected from the General Amber Force Field (GAFF),⁴³ the atomic charges were computed at HF/6-31G(d,p) level in aqueous solution (PCM) with the ChelpG fitting procedure.⁴⁴ In relation to the cDDP molecule, the intramolecular parameters were extracted from its optimized geometry in aqueous solution (IEF-PCM) at MP2(FC)/LANL2DZ/6-31G(d,p) level.^{45–47} In particular, the force constants of both bond stretching (K_b) and bending angle (K_a) were derived by using the Visual Force Field Toolkit (VFFDT).⁴⁸ With regard to the intermolecular parameters, we used the LJ 12-6 parameters set for the Pt atom reported by our group,⁴⁹ the LJ 12-6 parameters for the nonmetal atoms (Cl, N, and H) included in the GAFF, and the atomic charges calculated at HF/LANL2DZ/6-31G(d,p) level in aqueous solution (IEF-PCM) with the ChelpG scheme.

When it comes to the computational details of this MD simulation, we firstly placed our the cDDP@CNTox complex at the center of a truncated octahedral box (average length of 71.2 Å) containing water molecules described by the TIP3P model⁵⁰ and the ff99SB force field.⁵¹ By using periodic boundary conditions with the isothermal–isobaric ensemble (*NPT*), the density was equilibrated aiming to achieve an average value of 0.993 g cm⁻³ at $T = 310.15$ K and a pressure of 1.0 bar. The Langevin thermostat⁵² and the Berendsen barostat⁵³ were applied in order to keep temperature and pressure constant. While the van der Waals interactions were computed within a cutoff of 10 Å, the long-range electrostatic interactions were treated by means of the particle mesh Ewald method.⁵⁴ The Leap Frog algorithm was used to numerically integrate the Newtonian equations of motion considering a time step of 2.0 fs and the SHAKE algorithm⁵⁵ to constrain all bonds involving hydrogen atoms.

At last, the simulation protocol firstly involved the energy minimization of the solvent molecules followed by the entire system. While the first 1000 cycles of this stage were performed by using the steepest descent method,⁵⁶ the last 1000 cycles were

computed with the conjugate gradient method.⁵⁷ Next, the system was heated from 0 to 310 K with a weak restraint (10 kcal mol⁻¹ Å⁻²) and equilibrated for 540 ps. Finally, the production dynamics was conducted for 100 ns with the *NPT* ensemble. By selecting a snapshot of the cDDP@CNTox model every 5 ns from this trajectory (100 ns), we organized a set of 20 solvated structures of this inclusion complex that was employed to evaluate the explicit solvent effect on the calculation of NMR chemical shifts. This MD simulation was carried out with the AMBER 12 software⁵⁸ and the trajectory analysis were carried out with the CPPTRAJ tools.⁵⁹

Results and discussions

In this work we re-optimize the geometries of the molecular models of cDDP@CNTox or complex I ($\text{C}_{168}\text{H}_{24}\text{O}_{15}\text{Cl}_2\text{N}_2\text{Pt}$) and cDDP@CNCox or complex II ($\text{C}_{250}\text{H}_{38}\text{O}_{20}\text{Cl}_2\text{N}_2\text{Pt}$) inclusion complexes reported in our previous studies²⁶ at a higher level of theory: B3LYP/LANL2DZ/def2-SVP/IEF-PCM(UFF). The oxidized nanostructures topologies correspond to the final geometries obtained by De Souza *et al.*⁶⁰ in a study of the oxidation mechanism of the surfaces of carbon nanotubes and nanocones. Thus, here we shall describe the changes in the geometry of each inclusion complex and the ^{195}Pt and ^1H NMR chemical shifts, not reported in the literature yet.

Fig. S1 (ESI)[†] and 1 show the potential energy curve for the geometry optimization process of inclusion complexes and their fully optimized geometries, respectively. It can be seen that the energy difference between the starting structure and fully optimized geometries is only 2.44 kcal mol⁻¹ (cDDP@CNTox) and about 20 kcal mol⁻¹ (cDDP@CNCox). In complex I the cisplatin molecule is further from the oxidized region of CNTox, and this carbon structure is more rigid structurally than the CNCox model. Thus, in Fig. S1a,[†] ΔE reflects the slight adjustment of the structural parameters for the complex I. In complex II we observed that the drug molecule is closer to the oxidized region of CNCox and a larger ΔE (Fig. S1b[†]) is found between the initial and final geometries. CNTox (Fig. 1a) and CNCox (Fig. 1c) structures are about 13.4 Å and 18 Å in length, respectively, and represent the region where the drug is located in the nanostructure cavity. According to the literature, isolated units of carbon nanotubes¹⁵ and nanocones¹⁶ are obtained in the range of 40–800 nm in length. Free cisplatin geometry and its structure in the inclusion complexes were found to be square planar, as expected, with average Pt–Cl and Pt–N bond lengths of 2.35 Å and 2.07 Å, respectively, which compare well with the X-ray data: Pt–Cl equal to 2.33(9) Å and Pt–N equals to 1.95(4)/2.05(4) Å.⁶¹ Some hydrogen-bond interactions can contribute to the inclusion complexes stabilization. Both inclusion complexes have a short intramolecular hydrogen-bond interaction between hydroxyls of carboxyl groups (cDDP@CNTox) and, between a hydroxyl of carboxyl group and vicinal carbonyl (cDDP@CNCox), as shown in Fig. 1b and d. The intramolecular interaction in the complex I (Fig. 1b) is not present in the isolated CNTox (Fig. 1a). Comparing Fig. 1c and d we observe that after the formation of complex II the intramolecular interaction between two neighboring hydroxyl



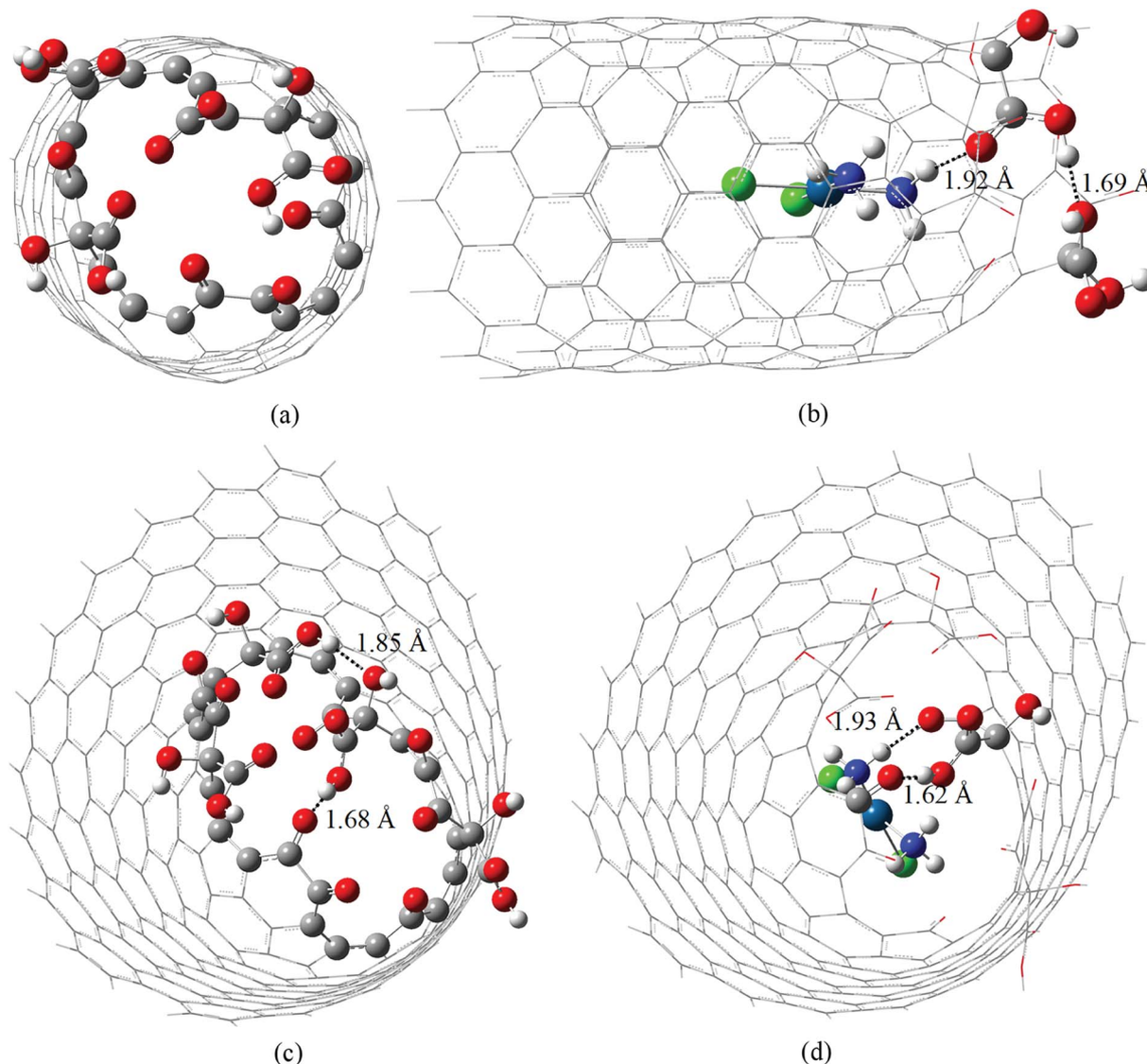


Fig. 1 B3LYP/LANL2DZ/def2-SVP/IEF-PCM(UFF) fully optimized structures of CNTox (a), cDDP@CNTox – complex I (b), CNCox (c) and cDDP@CNCox – complex II (d). Intra and intermolecular hydrogen interactions are highlighted as dotted lines.

groups of CNCox is broken. The intermolecular hydrogen-bonds in the two inclusion complexes are assigned as medium to long-distance, on average 1.92 Å (see Fig. 1b and d). They are between the NH₃ group (cDDP molecule) and the carbonyl group (carboxyl) of oxidized nanostructures.

Local ChelpG charges⁴⁴ in the interaction region between cisplatin and carbon atoms of the oxidized nanostructures are highlighted in Fig. 2. The scales represented by the color range show that the cisplatin molecule causes a more pronounced perturbation in the local charge distribution on the CNTox surface than on the CNCox surface when compared to the isolated monomers. The interatomic distances and local ChelpG charge analysis for Pt atom (cisplatin) and the nearest C atoms of the oxidized nanostructures shown in Fig. 2 are given in Table S1 (ESI).[†] The calculated average distance between the Pt atom and the CNTox surface is slightly smaller than that of the CNCox. The sum of the charge of carbon atoms closer to

cisplatin molecule shows a negative value of $\sim -0.2e$ (complex I) and $\sim -0.03e$ (complex II) which represents an equivalent local charge accumulation of $\sim -0.14e$ and $\sim -0.06e$ in relation to the isolated CNTox and CNCox monomers, respectively. Table S1[†] also shows that the negative charge concentration on the Pt atom in complex I is increased and in complex II is slightly decreased when compared to the calculated charge for the Pt atom in the isolated cisplatin molecule. These results suggest that the implicit effect of solvent (PCM-water), besides inferring an electrostatic interaction that stabilizes the inclusion complexes, it may also induce an extra interaction represented by stronger host-guest charge transfer from CNTox to metal in complex I than that from CNCox to metal in complex II.

Inclusion complexes formation energies (ΔE_F) are given in Table 1. We compared the values obtained in this work with those for the same complexes reported in ref. ²⁶. B3LYP/LANL2DZ/def2-SVP/IEF-PCM(UFF) energies of formation



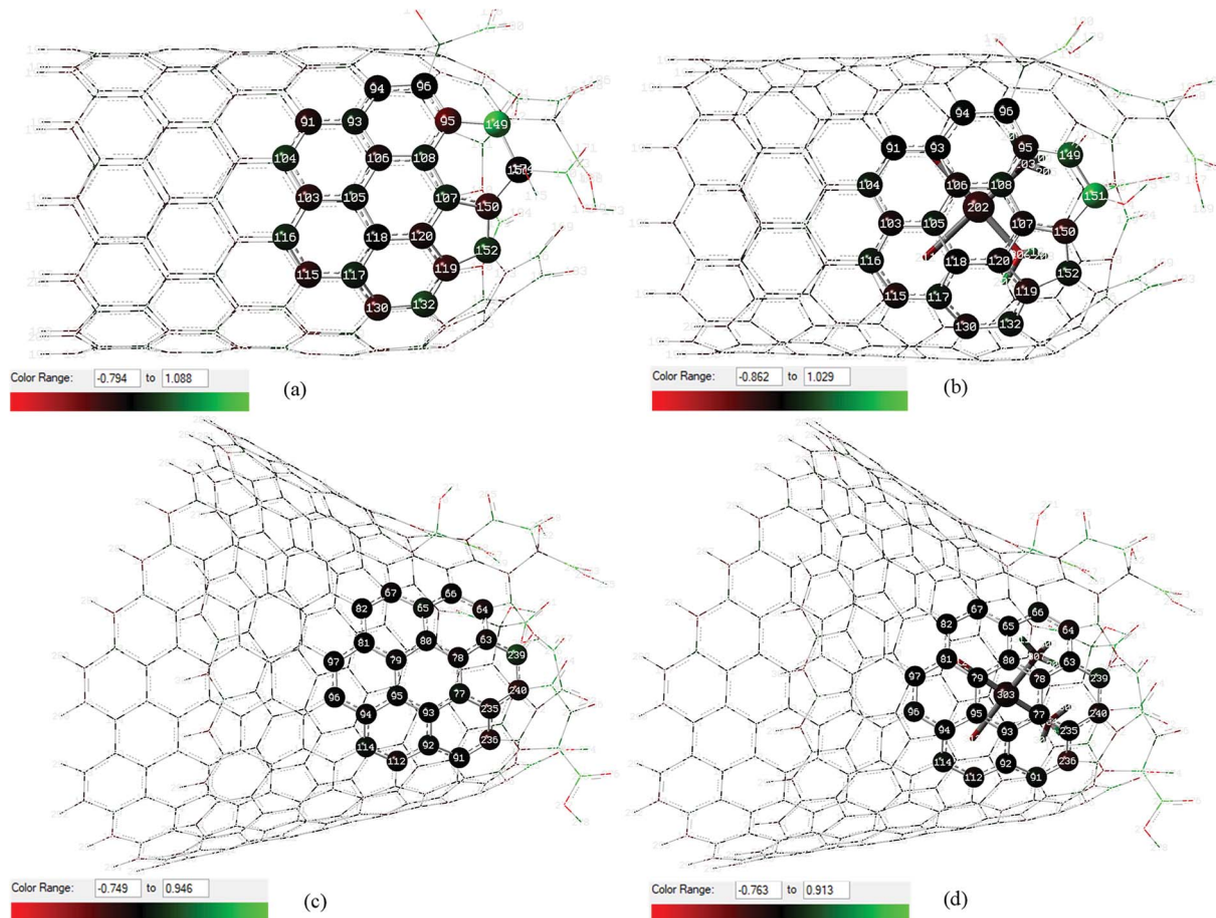


Fig. 2 Local ChelpG charges distribution calculated at B3LYP/LANL2DZ/def2-SVP/IEF-PCM(UFF) level in the interaction region between cisplatin molecule and carbon atoms of the oxidized nanostructures: CNTox (a) and inclusion complex cDDP@CNTox – complex I (b), CNCox (c) and inclusion complex cDDP@CNCox – complex II (d). Green and red represent positive and negative atoms according to the scale shown.

showed that complex I is more strongly bound than complex II. For the cDDP@CNTox complex I, it was observed that the ΔE_F calculated in this work ($-19.9 \text{ kcal mol}^{-1}$) is similar as that calculated in ref. ²⁶ where the BSSE was considered for the correction of ΔE_F with the 6-31G basis set. For the cDDP@CNCox complex II, the difference between ΔE_F values is more significant when we consider these two levels of theory ($-5.7 \text{ kcal mol}^{-1}$ in the present work – see Table 1). Kruse *et al.*⁶² showed examples for geometry optimizations calculations of larger systems with inter- and intramolecular BSSE using DFT

calculations and def2-SVP basis sets, where the BSSE is typically below 10% of the interaction energy. Here, we do not perform the BSSE calculation since we believe that it is small for def2-SVP basis sets. We also believe that using a post-Hartree-Fock level of theory, which different from the DFT calculations includes explicitly electron correlation effects, such as MP2 calculation,⁶³ and the use of larger basis sets with highly polarized and diffuse functions are unfeasible from a quantum and computational point of view for large molecular systems with 250–350 atoms, such as those studied in this work.

Table 1 Energy of formation (ΔE_F) for the cDDP@CNTox and cDDP@CNCox inclusions complexes. All values in units of kcal mol^{-1}

Theory of level	cDDP@CNTox	cDDP@CNCox
B3LYP/LANL2DZ/def2-SVP/IEF-PCM(UFF) ^a	-19.9	-5.7
PBEPBE-D3/LANL2DZ/def2-SVP/IEF-PCM(UFF) ^a	-99.9	—
B3LYP/LANL2DZ/6-31G/gas-phase – BSSE ^b	-19.8	-19.2
PBEPBE-D3/LANL2DZ/def2-SVP/IEF-PCM(UFF) (single point) ^c	-73.2	-42.0
B97D/LANL2DZ/def2-SVP/IEFPCM(UFF) (single point) ^c	-82.0	-38.0

^a Calculated from eqn (1) in this work. ^b Results from ref. 26. ^c Calculated using the B3LYP/LANL2DZ/def2-SVP/IEF-PCM(UFF) fully optimized structures of inclusion complexes.



However, we performed the geometry optimization of complex I with dispersion corrected including the D3 Grimme's empirical dispersion⁶⁴ to GGA PBE functional.^{36,37} In addition, through single point (SP) calculations, the PBEPBE-D3 and B97D⁶⁵ functional, it was used to evaluate the effect of the level of theory on the both inclusion complexes formation energies. A comparison of the B3LYP with PBEPBE-D3 and B97D SP results showed that the dispersion correction leads a much larger stabilization of the complexes. Table 1 show that the both PBEPBE-D3 and B97D ΔE_F is about 75% (cDDP@CNTox) and 86% (cDDP@CNCox) lower, respectively, than the B3LYP values. ΔE_F of the cDDP@CNTox complex optimized at PBEPBE-D3/LANL2DZ/def2-SVP/IEF-PCM(UFF) level is even lower (80%) than the B3LYP level. The small difference between the PBEPBE-D3 values (5%) suggests that the calculation of geometry optimization may not compensate the computational cost obtained. There is a slight change in the geometry of the oxidized region of CNTox when comparing the calculated structures with the B3LYP and PBEPBE-D3 functional. The equilibrium point for the cisplatin molecule in the cDDP@CNTox complex is practically the same at both levels of theory. The dispersion terms that describe better the van der Waals interactions and hydrogen-bonds, which are important for the stabilization of inclusion complexes. In spite of the effect of dispersion corrections, our results suggest that both inclusion complexes are energetically favorable. The higher stability of complex I might be related to the formation of intramolecular hydrogen-bonds in complex I and the disappearance of this type of interaction between neighboring hydroxyl groups in the oxidized region of CNCox in complex II geometry (see Fig. 1c and d). Moreover, the possibility of charge transfer occurring between the CNTox surface and the Pt atom (cDDP) might strength the interaction between monomers in complex I. Harmonic frequency calculations were not attempted due to the size of the molecular system and high level of theory used, since it is a quite demanding computational task. Discussions about the thermodynamic parameters of these inclusion complexes and similar ones can be seen in our previous works.^{23–27,32}

Fig. 3a–c show the PBEPBE/NMR-DKH/IEF-PCM(UFF) ¹H NMR spectra for free cDDP, and cDDP@CNTox and cDDP@CNCox inclusions complexes, respectively. The solvent effect (water) was accounted for using single point NMR calculations on the fully optimized geometries following the computational protocol proposed by Paschoal *et al.*³² Our calculations show that the chemical shifts (δ) for the two distinct hydrogen atoms of cisplatin molecule, opposite (H_a) and near (H_b) the chlorine atoms, are 4.5 and 4.1 ppm (4.3 ppm on average – Table 2), respectively. The experimental value,⁶⁶ measured in a 5 mmol dm⁻³ solution containing cisplatin in 95% H₂O and 5% D₂O, pH 4.72, is 4.06 ppm, which is a fair agreement. The calculated ¹H $\Delta\delta$ values may be trusted to ± 0.5 ppm, which is adequate for this work in what changes due to complex formation is concerned. Fig. 3b and c show that in the two inclusion complexes formed, the hydrogen atoms of cisplatin (NH₃ groups) exhibit a different behavior for the chemical shift variation ($\Delta\delta$). When the cisplatin spectrum (Fig. 3a) is compared to that of complex I (Fig. 3b), the ¹H NMR

signals are shifted to high-field region (more shielded), -5.1 ppm on average (see Table 2). The electron-rich carbon nanostructure and the oxygen groups on the CNTox surface, in addition to the close proximity of the drug molecule to the inner wall of the tube, increase the shielding of the hydrogens of cDDP in complex I; therefore, these protons tend to absorb energy on higher intensity fields. These shifts reach up to -6.0 ppm for H_a and -8.2 ppm for H_b nuclei (see definition in Fig. 3a). This result can also be discussed in terms of the ring current effect that affects the chemical NMR shifts of molecules adsorbed on carbon nanostructures. Forse *et al.*⁶⁷ carried out a theoretical-experimental study of the nuclei-independent chemical shift (NICS) calculations for molecular species adsorbed on sp²-hybridized carbon systems. The authors observed that the larger graphene-like systems and their derived concave surfaces result in greater chemical shielding of atomic nuclei, even at distances relevant for molecular adsorption. According to Casabianca,⁶⁸ the largest ring current effect on concave carbon surfaces (such as carbon nanotubes) on NICS, is due to two competing factors: (i) increased curvature breaks ring aromaticity and decreases the magnitude of the NICS, concomitantly; (ii) the proximity of the rings faces increases the shielding overcoming the decrease due to loss of surface aromaticity. Thus, the proximity of the cisplatin molecule to the concave surface of the nanotube in the cDDP@CNTox model, more strongly shields the ¹H nuclei shifting them to high-field region in the NMR spectrum (Fig. 3b). On the other hand, we observed the inverse effect for the protons of cDDP in complex II. In complex II, H_a nuclei have $+1.5 < \Delta\delta < +4.3$ ppm and H_b nuclei have a symmetric shift of only $+0.1$ ppm (see Table 2), both for low-field region. The average hydrogen chemical shift was $+6.6$ ppm. This result shows that cisplatin protons are less shielded in complex II. The reason of the distinct trends for complexes I and II might be in the interaction mode of cDDP guest. For the narrower CNTox, the cDDP is closer to the electron-rich carbon surface increasing the shielding of hydrogen atoms (high-field shift), whereas, for CNCox the cDDP is closer to the oxygen groups at the oxidized tip, interacting through hydrogen bond and de-shielding the H nuclei (low-field shift). These results agree with the ChelpG charge analysis and the degree of interaction between the monomers in the two inclusion complexes studied.

The experimental ¹⁵N shifts are referenced to 1.5 mol dm⁻³ NH₄Cl in 1 mol dm⁻³ HCl and equal to -68.7 ppm.⁶⁶ In this work, PBEPBE/NMR-DKH/IEF-PCM(UFF) ¹⁵N NMR spectra for free cDDP (Table 2) is -63.3 ppm. Our interest is to observe the $\Delta\delta^{15}\text{N}$ of cDDP molecule into carbon nanostructures which is much larger than the deviation ($\sim 7\%$) between theoretical and experimental values for free cDDP. Table 2 shows large variations upon complex I formation, on average -18.6 ppm for high-field region; for complex II, an average shift about 7 ppm is observed for the low-field region. These results show that in the cDDP@CNTox complex, the ¹⁵N nuclei (cisplatin NH₃ groups) are also more shielded due to the ring current effects.

Fig. 3d–f show PBEPBE/NMR-DKH/IEF-PCM(UFF) ¹⁹⁵Pt NMR spectra for free cDDP, and the cDDP@CNTox and cDDP@CNCox inclusions complexes, respectively. Table 2



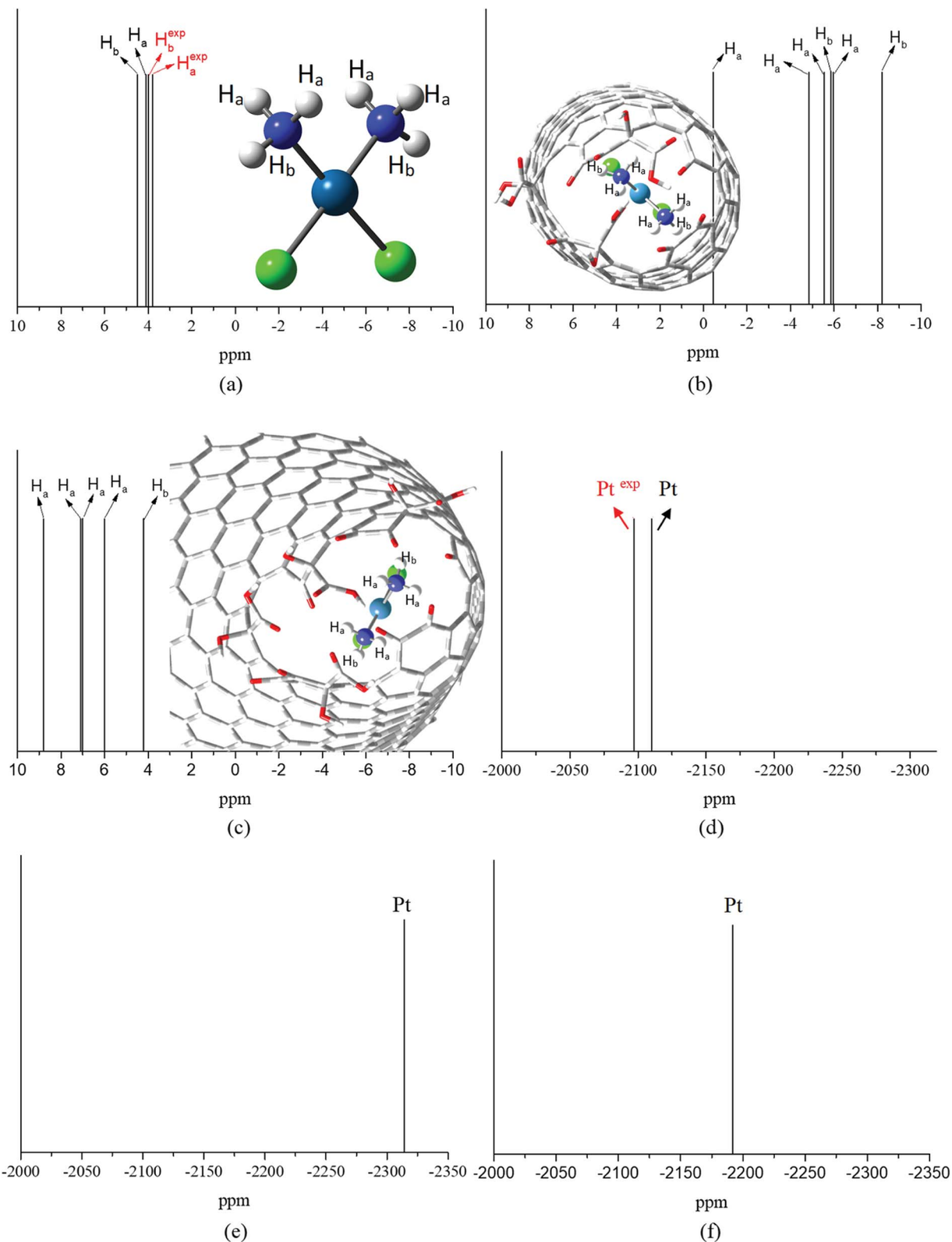


Fig. 3 PBE/PBE/NMR-DKH/IEF-PCM(UFF) ^1H and ^{195}Pt NMR spectra for free cDDP (a and d), cDDP in the cDDP@CNTox – complex I (b and e) and in the cDDP@CNCox – complex II (c and f) inclusion complexes. All values in ppm.

shows that the use of the relativistic NMR-DKH Gaussian basis sets³² for Pt and other atoms, commonly used as Pt-ligands, leads to δ (calculated by eqn (3)) equals to -2110 ppm for

^{195}Pt nucleus of free cisplatin. This result is in excellent agreement with the experimental⁶⁹ value for cisplatin in aqueous solution (D_2O) that is equals to -2168 ppm, a deviation of only



Table 2 PBEPBE/NMR-DKH/IEF-PCM(UFF) ^1H , ^{15}N and ^{195}Pt NMR chemical shift values for the free cisplatin and its inclusion complexes. ^1H NMR chemical shift values for the oxidized region of free nanostructures and its inclusion complexes are also shown. All values in ppm

Structures	Chemical shifts (δ)/ppm						
	$^1\text{H}_a$	$^1\text{H}_b$	^{15}N	^{195}Pt			
Free cDDP	4.5 (4.06) ^a	4.1 (4.06) ^a	-63.3 (-68.7) ^a	-2110 (-2168) ^b			
cDDP@CNTox	-0.4, -4.9, -5.5, -6	-5.9, -8.2	-79.4, -84.5	-2314			
cDDP@CNCox	8.8, 7.1, 7, 6	4.2	-57.9, -54.7	-2192			

Structures	Chemical shifts (δ)/ppm					
	Carboxyls			Hydroxyls		
	H_c	H_d	H_e	H_f	H_g	H_h
CNTox	8	8.7	8.4	5.2	7.7	6.2
cDDP@CNTox	14.2	10.7	10.7	7.1	9.7	8.1

Structures	Chemical shifts (δ)/ppm							
	Carboxyls				Hydroxyls			
	H_i	H_j	H_k	H_l	H_m	H_n	H_o	H_p
CNCox	9.8	15.2	14.7	9.4	5.7	8.2	9.2	7
cDDP@CNCox	10	16.6	13.5	9.5	5.6	8	7.9	6.5

^a Experimental data from ref. 66. ^b Experimental data from ref. 69.

2.7%. Upon the formation of inclusion complexes, we observed a significant ^{195}Pt signal shift to high-field region for the cDDP@CNTox, found at -2314 ppm. For cDDP@CNCox, the ^{195}Pt chemical shift was at -2192 ppm, also found at high-field region in relation to free cisplatin. These results shown that the cDDP molecule is in closer contact with electron-rich tube surface in complex I than in complex II. Thus, ^{195}Pt nucleus in complex I is more shielded by the ring current effects of the concave surface of the nanotube than in complex II where the internal surface of the nanocone is less concave.

Fig. S2 (ESI)[†] show the PBEPBE/NMR-DKH/IEF-PCM(UFF) ^1H NMR spectra calculated for the organic groups on the oxidized tip of CNTox (Fig. S2a[†]) and CNCox (Fig. S2b[†]) and in the inclusion complexes geometries, Fig. S2c and d,[†] respectively. Analyzing the Fig. S2a and c,[†] we note that the proton signals of CNTox shift to low-field region (less shielded) when the complex I is formed (Table 2). According to Fig. S2b and d,[†] due to the proximity of cisplatin to the oxidized surface, most proton signals (H_i , H_m , H_n , H_o and H_p) of CNCox slightly shift to high-field region upon the complex II formation, with average values for both species around 9.8 ppm (Table 2).

The solvent effect on the calculation of NMR chemical shifts is discussed here. As has been pointed out by Benzi *et al.*⁷⁰ the continuum solvation models may not be very adequate for the description of solvent effect on DFT calculations of chemical shifts, in what N-H and O-H protons are concerned, due to hydrogen-bond and other solute-solvent interactions. It has been shown recently by Da Silva and De Almeida⁷¹ that N-H chemical shifts calculated using the GIAO method and TMS as

reference for a series of amine compound (in chloroform) at DFT level using the PCM is systematically underestimated independent of the specific functional employed. The agreement with experimental N-H chemical shifts (in CHCl_3) is significantly improved when explicit CHCl_3 solvent molecules are included in the NMR calculations, reaching a deviation of -0.1 to -0.2 ppm. The same result was found for cisplatin molecule⁷¹ with deviation around 2 ppm being predicted at B3LYP/LANL2DZ/6-31G(d,p)/IEF-PCM(UFF) and PBEPBE/LANL2DZ/6-31G(d,p)/IEF-PCM(UFF) levels of calculation. In the present work using the PBEPBE/NMR-DKH/IEF-PCM(UFF) level the deviation from experiment for N-H cisplatin protons drops to approx. 0.5 ppm, which is a good improvement over the corresponding DFT calculations with the standard 6-31G(d,p) basis set. We expect a similar behavior for the O-H (organic groups on the oxidized surface) chemical shifts, and so O-H $\Delta\delta$ values can be trusted to ± 0.5 ppm.

In order to analyze the explicit solvent effect on the calculation of NMR chemical shifts, we performed a MD simulation of the cDDP@CNTox complex in explicit solvent (water molecules). Specifically, the cDDP@CNTox model was selected as a prototype for this analysis which involved the same parameterization procedure and simulation protocol recently employed in our works^{27,72} and, moreover, in this work it represents the most energetically favorable model.

The simulation protocol proved to be effective in providing the equilibration of the complex I by verifying a convergence in the temporal variation plots of the temperature, kinetic and potential energy, density, volume and pressure through the 100



ns (Fig. S3, ESI[†]), as also concluded in our recent works with similar systems.^{27,72} The RMSD value of 0.32 ± 0.05 Å suggests a structural stability of this complex.

Since the NMR chemical shift is sensitive to the chemical environment, we calculated the radial distribution function (RDF) designated as $g(r)$ CM-O_w (see Fig. 4a), which were defined between the center of mass (CM) of the CNTox model and the oxygen atom (O_w) of the water molecules, aiming to define the solvent molecules distribution closest to the encapsulated cDDP drug. In Fig. 4a, the first peak at 2.55 Å indicated the presence of water molecules inside the cavity of nanotube forming an inner solvation shell as previously reported by our group for the carbon nanocones models.^{27,72} By computing the integral over this first peak, we found a coordination number equals to 2 from the analysis of the red dot line. This means that, on average, there are 2 water molecules distributed around the CM of CNTox. Although this number seems to be small compared to the 13 water molecules found in our previous cDDP@CNHox model,⁷² it is worth to emphasize that the model studied here (C₁₆₈H₂₄O₁₅Cl₂N₂Pt) contains a substantially small nanostructure than that model (C₃₆₀H₃₈O₂₀Cl₂N₂Pt) recently studied by our group. When it comes to the solvation structure referring to the oxidized region of the CNTox model, we also calculated the RDF named as $g(r)$ O_{CNT}-O_w (see Fig. 4b, which was defined between the oxygen atoms (O_{CNT}) of the functional groups inserted on the CNTox structure and the O_w of the water molecules. By evaluating Fig. 4b, we noticed that while the first solvation shell was centered at 2.85 Å involving, on average 1 water molecule, the third solvation shell was located at 6.45 Å (third peak in Fig. 4) including, on average, 27 solvent molecules.

The temporal variation of the three species (cDDP molecule and two water molecules) encapsulated inside the CNTox cavity (Fig. 4) is presented in Fig. 5a–d, which involves the overlay of 200 frames collected from the 100 ns production run. From

Fig. 5a–d, it was possible to conclude that, on average, the plane defined by the square planar geometry of the cDDP molecule remained parallel to the main axis of the carbon nanostructure, as well reported in our recent MD simulations results.^{27,72} In addition, NH₃ groups of cisplatin also remained close to the oxidized region of the CNTox. The same spatial configuration was also observed in our inclusion complexes involving pristine,²⁷ oxidized, and reduced CNH topologies.⁷² Regarding the inner solvation shell, we observed that the water molecules did not access the space between the cDDP molecule and the oxidized region of the CNTox structure, probably due to steric hindrance of the van der Waals radii. In spite of involving the same overlay of 200 frames selected from the production trajectory (100 ns), Fig. 5e and f focus on the temporal variation of the 10 water molecules closest to the oxidized region of the cDDP@CNTox. These solvent molecules integrated a part of the third solvation shell described by the $g(r)$ O_{CNT}-O_w (see Fig. 4b). Fig. 5e and f clearly depicted the almost spherical distribution of the water molecules around the oxidized region of the CNTox model during the 100 ns simulation.

With the inner solvation shell data of the cDDP@CNTox complex, a set of 20 structures (frames) of this model was collected from the 100 ns MD simulation taking into account only the two explicit water molecules. Specifically, these topologies (see the snapshots in Fig. S4–S7, ESI[†]), named as **Structure N** (with N in the range of 0 to 19), were selected at 5 ns intervals from the total simulation time. It is worthy to mention that the 20 geometries of the cDDP@CNTox model were not reoptimized at the DFT level of theory before the aforementioned NMR calculations, since the Pt-ligand distances were, on average, close to the experimental results.⁶¹

Table S2 (ESI[†]) shows the PBEPBE/NMR-DKH/IEF-PCM(UFF) ¹H, ¹⁵N and ¹⁹⁵Pt NMR chemical shifts for selected structures (cDDP@CNTox) obtained from the MD simulation. Some dynamic effects can be considered when discussing the

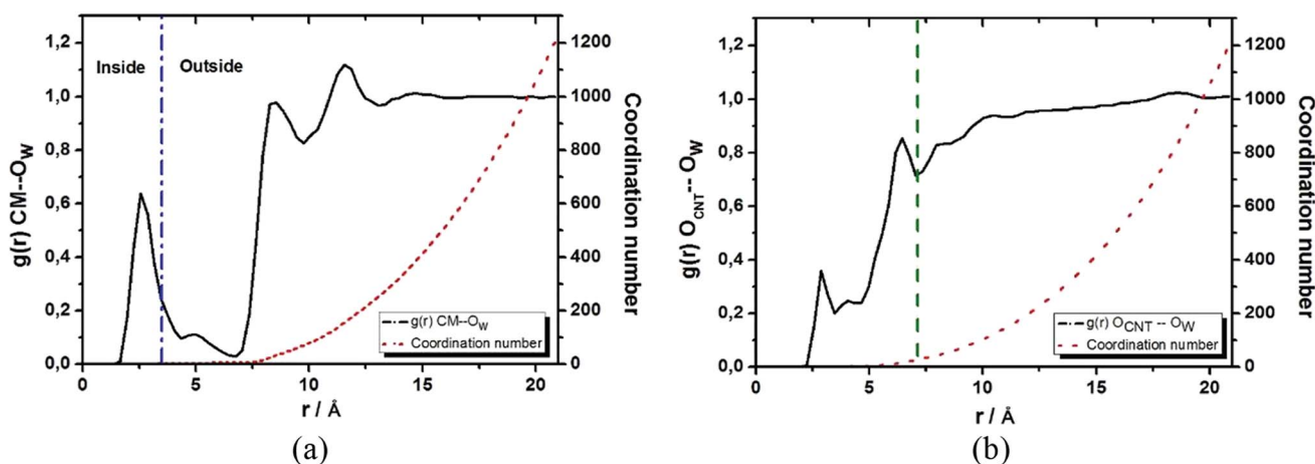


Fig. 4 Radial distribution functions: $g(r)$ CM-O_w defined between the center of mass of the CNTox structure and the oxygen atom of the water molecules (a) and $g(r)$ O_{CNT}-O_w referring to the oxygen atoms inserted in the CNT structure with respect to the water molecules (b). While the red dot line indicates the coordination number, the blue dash dot line demarcates the water molecules presented in the internal and external region of the CNTox model, and the green dash line indicates the third solvation shell with respect to the oxidized region of the carbon nanostructure.

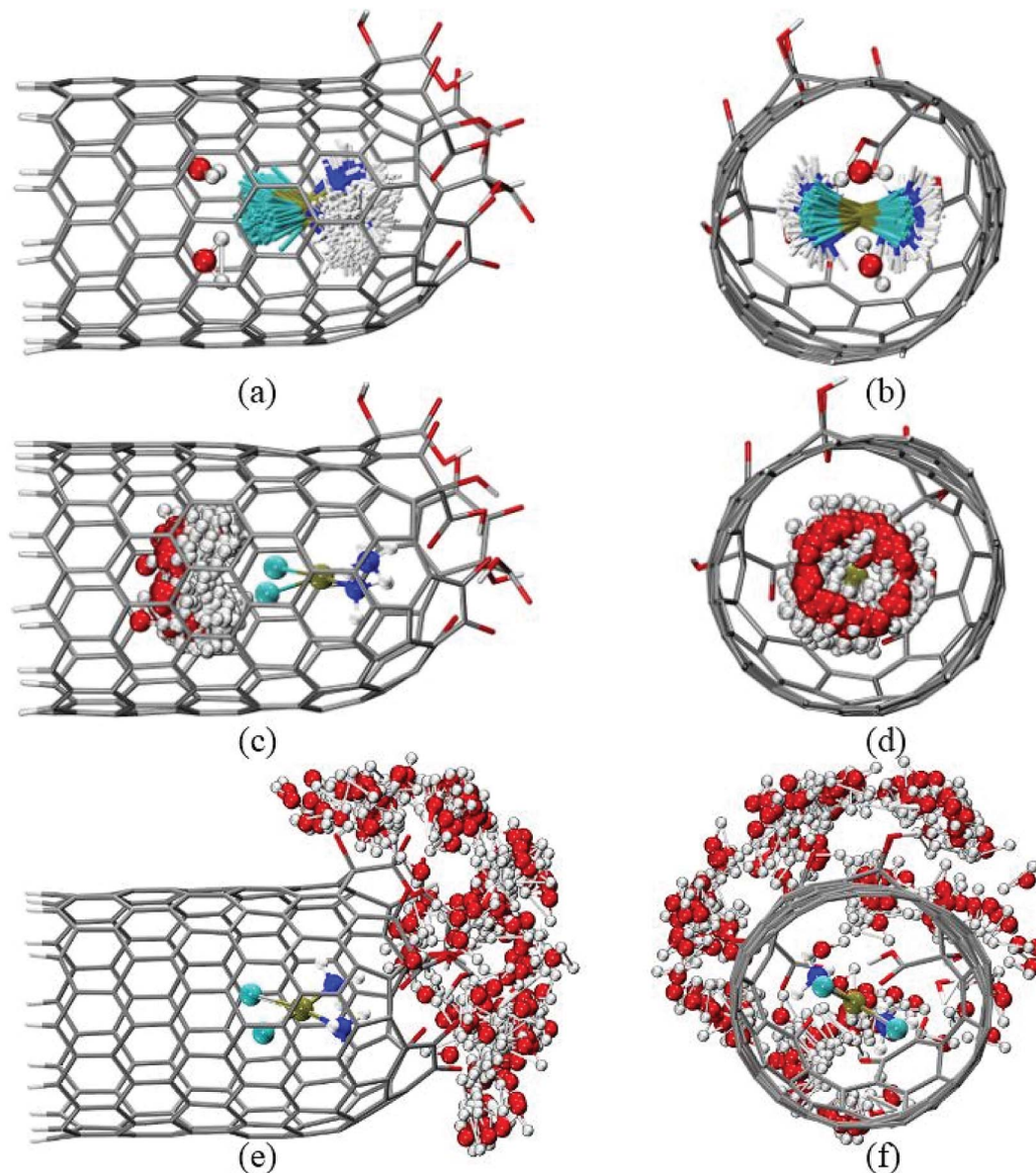


Fig. 5 Temporal variation of the encapsulated cDDP molecule, the two encapsulated water molecules, and the ten water molecules around the oxidized region of the CNTox: side view (a, c, and e) and front view (b, d, and f), respectively.

chemical shifts, such as: the temporal variation of cisplatin (Fig. 5a and b) and the two water (Fig. 5c and d) molecules inside the nanotube cavity and, the organic groups of the CNTox oxidized region (see Fig. 5e and f). ^1H δ values of cDDP (NH_3 groups) are more sensitive with respect to temporal variation of the drug itself and organic groups of the CNTox. Among the selected frames, ^1H signals can be observed at -188 ppm (**Structure 12**) and -110.3 ppm (**Structure 17**), both high-field shifted, respectively. For **Structure 3**, for example, H_a and H_b signals can be observed at 115 ppm and 96 ppm, both low-field shifted, respectively. From Fig. S8 to S10 (ESI),[†] the dispersion of the ^1H signals of the cisplatin are observed. ^1H chemical shifts are influenced by the rotations of the NH_3 groups around the Pt-N bond axis and also by the temporal variation of the organic groups of the CNTox model during the MD simulation.

^1H nuclei are more strongly shielded (high-field shifted) when the organic groups are closer to them; by contrast, the shielding effect is less (low-field shifted) when such groups are more distant from the cisplatin protons. Thus, considering the dynamic nature of these rotations and the selected frames, our NMR calculations revealed that the protons (NH_3 groups) of cisplatin can be identified in the range 115 to -188 ppm. ^{15}N chemical shifts (Table S3[†]) appear to be more dependent on the temporal variation of the cisplatin molecule that is described by incomplete rotations around the center of mass of the drug in the tube cavity. The ^{15}N δ values are found in the 272–102 ppm range (low-field region). Fig. 5a and b shows that for the selected frames, the translational movement of cDDP molecule inside the CNTox cavity is limited. The intermolecular distances between the two water molecules revealed by the RDF analysis



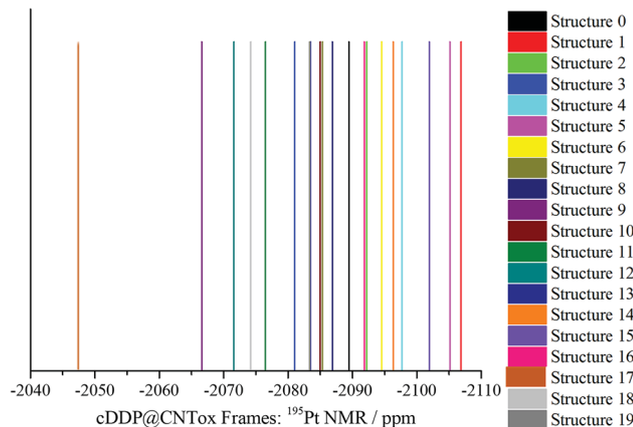


Fig. 6 PBEPBE/NMR-DKH/IEFPCM(H₂O) ¹⁹⁵Pt NMR spectrum of structures 0 to 19 (frames) obtained by MD simulation is shown in Fig. 5a and b.

and the drug molecule vary in the 3.86–4.84 Å range (see Table S3, ESI).[†] Fig. 6 shows the ¹⁹⁵Pt NMR spectra for the 20 selected structures obtained from MD trajectory (see also Table S2[†]). The average chemical shift (−2086 ppm) in the low-field region compared to the experimental value (−2168 ppm, see Table 2) suggests that the two water molecules have no direct influence on the shielding of the ¹⁹⁵Pt nucleus. In relation to the cDDP@CNTox model optimized at B3LYP/LANL2DZ/def2-SVP/IEF-PCM(UFF) level, a difference of approximately +228 ppm for the low-field region can be estimated. Thus, as observed for the hydrogens, the ¹⁹⁵Pt NMR chemical shift is also sensitive to the dynamics of the cDDP molecule inside the CNTox cavity. We selected a structure from the MD simulation to investigate the effect of 10 water molecules on the ¹H NMR spectrum of the CNTox oxidized region, the ¹H δ are shown in the Table S4 (ESI).[†] For the isolated CNTox in the inclusion complex geometry, all ¹H δ values are shifted to the low-field region, on average 8.4 ppm (carboxyls groups) and 9.4 ppm (hydroxyls groups). For the cDDP@CNTox, all ¹H δ values of carboxyls groups are shifted to the high-field region while the majority of proton signals of the hydroxyl groups are shifted to the low-field region. These results show that the field effect due to the presence of the cDDP molecule is greater than that caused by the 10 water molecules.

In light of the reported results, we recommend the new NMR-DKH basis sets and the computational protocol proposed by Paschoal *et al.*³² as a good alternative to predict the ¹⁹⁵Pt and ¹H NMR spectra of inclusion complexes formed by cisplatin and carbon nanostructures. Our results combined with experimental NMR investigations can demonstrate an efficient and promising strategy for the application of oxidized carbon nanostructures (CNTox and CNCox) in the area of drug delivery and controlled release systems.

Conclusions

In this work we used the relativistic NMR-DKH basis sets for the calculation of the ¹H, ¹⁵N and ¹⁹⁵Pt NMR spectra of inclusion complexes formed between cisplatin and oxidized carbon

nanotube and nanocone molecules. The structural results at B3LYP/LANL2DZ/def2-SVP/IEF-PCM(UFF) level showed that intermolecular hydrogen-bonds contribute to the complexes stabilization. The complex I formed by cDDP and CNTox was around 70% more favorable than that with CNCox. According to the reported results, the experimental detection of the inclusion compound could be attained through the analysis of the ¹H and ¹⁵N (NH₃ groups), and ¹⁹⁵Pt NMR spectra for cisplatin guest molecule. ¹H NMR results showed a shift in the NMR signals from 4.3 ppm (cDDP) to −5.1 and +6.6 ppm (cDDP@CNTox and CDDP@CNCox complexes, respectively). Our ¹⁹⁵Pt NMR also shifted upon inclusion, found at −2314 and −2192 ppm for inclusion complexes with CNTox and CNCox nanostructures, respectively compared to the free CDDP (−2110 ppm). We investigated the explicit solvent effect (water molecules) on the calculation of ¹H, ¹⁵N and ¹⁹⁵Pt NMR chemical shifts by MD simulation of the cDDP@CNTox complex. The results showed that the two water molecules inside the nanotube cavity do not directly influence the ¹H, ¹⁵N and ¹⁹⁵Pt chemical shifts of the cisplatin molecule. The calculated spectra are more sensitive to the dynamics of the drug relative to the inner surface of the tube. Lastly, in addition to the chemical evidences which might guide the experimentalists to characterize the cDDP@CNTox (or CNCox) DDS, the computational protocol used here including the NMR-DKH basis sets might be useful to predict ¹⁹⁵Pt NMR chemical shift of big molecules such as the inclusion complexes formed by cisplatin and carbon nanostructures with a reasonable computational cost.

Conflicts of interest

There are no conflicts to declare.

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