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Development of fluorescent carbon nanoparticles from *Madhuca longifolia* flower for the sensitive and selective detection of Cr⁶⁺: a collective experimental–computational approach†

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Herein, blue-emitting carbon nanoparticles (CNP) were synthesized using the *Madhuca longifolia* flower for the highly selective and sensitive detection of Cr⁶⁺ ions in aqueous media using a simple, green, and cost-effective approach, and computational experiments were also performed. The prepared CNPs were well-dispersed in water with an average diameter of 12 nm and functionalized with carbonyl, hydroxyl and carboxylic acid groups. The decrease in the fluorescence intensity of the CNPs with an increase in the content of Cr⁶⁺ provided an important signal for the sensitive and selective detection of Cr⁶⁺ in aqueous media. The limit of detection for Cr⁶⁺ in an aqueous medium was found to be 103 ppb, which is more sensitive in comparison with the previously reported study. Furthermore, the validation of the proposed higher sensing feature and more selective nature of the CNPs towards Cr⁶⁺ was also explained using an *in silico* approach. The results from the theoretical calculations based on the DFT approach demonstrated that the binding energy (BE) of the CNPs with three transition metal (TM) cations (Cr⁶⁺, Fe³⁺, and Hg²⁺) follows the order of Cr⁶⁺ > Fe³⁺ > Hg²⁺, where the Cr⁶⁺ TM cation associated with the CNPs possesses the highest valence state, showing the highest sensing feature and highest selectivity among the investigated ions, as expected. The metal ions associated with the CNPs having a higher charge and a smaller radius indicated a higher BE and larger degree of deformation of the CNPs. Moreover, to achieve new insights into the structural, stability/energetics, and electronic features, some useful tools, such as NCI-plot, HOMO–LUMO gap, MESP, and QTAIM analysis were employed, which facilitated noteworthy outcomes.

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1. Introduction

Heavy metals are required for normal metabolism at specific concentrations; however, the presence of an excess amount of heavy metal ions causes severe disorders in humans. Chromium is one of the significant heavy metals that is discharged into the environment through industrial and anthropogenic activities.¹ Among the prevalent and stable forms of chromium, hexavalent chromium (Cr⁶⁺) such as dichromate (Cr₂O₇²⁻) and chromate (CrO₄²⁻) is considered to be the most toxic due to its

high mobility and carcinogenic and mutagenic effects in both living organisms and the environment.^{2,3} The human nervous and cardiovascular systems are susceptible to the accumulation of Cr⁶⁺ and it is very difficult to detoxify its toxic effect. In this case, Cr⁶⁺-contaminated drinking water and air even at low concentrations cause major health issues, such as lung cancer, edema, pulmonary congestion, gastrointestinal ulceration, dermal irritation and liver damage.^{4,5} Therefore, the detection of chromium [Cr⁶⁺] has received significant attention given that it is one of the most toxic heavy metal ions in recent years, the use of which has been significantly curtailed globally. Its concentration in water used for drinking is rigorously regulated and must not exceed the micromolar range. The US Environmental Protection Agency placed it at a danger threshold of 0.1 mg L⁻¹ (100 ppb) for total Cr and beyond this limit, it causes carcinogenicity and genotoxicity to humans.⁶⁻⁸

Therefore, the quantitative and qualitative detection of Cr⁶⁺ is essential to protect human health. In this case, various fluorescent sensors have been developed for the convenient and rapid detection of Cr⁶⁺ heavy metal in water.⁹⁻¹⁴ Among the various

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fluorescent materials, carbon nanoparticles (CNPs) have attracted significant attention owing to their green synthesis, biocompatible nature, water dispersibility, tunable fluorescence, ease of surface functionalization, *etc.*^{15–18} Thus, CNPs are considered alternative fluorescent materials to organic dyes and semiconductor quantum dots (QDs). The properties of CNPs have been extensively used in various applications such as biomedicine,^{19,20} optoelectronics,^{21,22} fluorescence sensing,^{23,24} photocatalysts,^{25,26} water purification,^{27,28} and visible ink.^{29,30} Carbon-rich precursors are used for the synthesis of fluorescent CNPs on a feasible scale using the top-down and bottom-up approaches.³¹

In recent years, vegetables,³² fruits,³³ agricultural waste,³⁴ biomass,³⁵ and hydrocarbons³⁶ have been widely used for the development of non-toxic fluorescent CNPs. For example, blue fluorescent, water-soluble, and nitrogen-doped CNPs were prepared *via* the calcination of diethylenetriaminepentaacetic acid³⁷ and hydrothermal treatment of methyl glycine diacetic acid trisodium salt³⁸ for the determination of Cr⁶⁺ ions with a calculated detection limit of 0.15 mM and 2.1 μ M, respectively, through turn-off of fluorescence due to the inner filter effect process. X. Guo and co-workers developed a fluorescent aerogel with a 3D net-like structure based on carboxymethylated cellulose nanofibrils and carbon dots as a highly effective adsorbent and sensitive optical sensor of Cr⁶⁺ and its limit of detection was found to be 17.6 mg L⁻¹ using the calibration plot. A fluorescent microcline-based nanoprobe was developed by incorporating nitrogenous carbon dots in the natural microcline nanostructure for the sensitive detection of the Cr⁶⁺ with a limit of detection of 4.0 μ M and it was found that photoinduced electron transfer is responsible for the fluorescence quenching.³⁹ J. Goswami *et al.* prepared banana stem-derived blue emissive carbon dots using phosphoric acid treatment as a novel sensor for the detection of Cr⁶⁺ *via* a photoinduced electron transfer quenching mechanism.⁴⁰

According to the literature and the above-mentioned examples, it can be observed that green carbon-rich precursors and chemical-free synthetic approaches for the preparation of fluorescent CNPs for the detection of Cr⁶⁺ have not been explored to date. Furthermore, the detection limit for Cr⁶⁺ using fluorescent CNPs in the above-mentioned reports appears to be slightly high. Therefore, the development of environmentally benign approaches for the sensitive detection of Cr⁶⁺ in aqueous medium using natural carbon-rich precursors is necessary. Therefore, in the present study, the *Madhuca longifolia* flower (a highly rich source of carbohydrates and vitamin C) was chosen,^{41,42} which mainly contains carbon, hydrogen, and oxygen. The presence of a high content of carbon and oxygen-containing groups in the *Madhuca longifolia* flower provides high potential for the synthesis of blue fluorescent CNPs in aqueous medium. Herein, *Madhuca longifolia* flowers were used as a sustainable source of carbon for the preparation of blue-emitting CNPs. Subsequently, the CNPs were used as a nanosensor for the detection of Cr⁶⁺ in an aqueous medium *via* a static quenching mechanism. A detailed experimental investigation was carried out to understand the sensing mechanism for the detection of Cr⁶⁺, which was further supported employing an *in silico* approach.

A few sensing features were established by the non-covalent interactions (NCIs) between the carbon network of the CNPs and external stimuli. Furthermore, useful insights into the sensing features [in terms of the supramolecular approach-based binding energy (BE) or interaction energy (IE)], especially the metal–nonmetal bonding interaction between the metal cation and nonmetals of the associated CNP complex were explored by performing an extensive computational experiment including the involved NCIs and van der Waals (vdW) (extremely weak) interaction(s). To complement the experimental results of the selective sensing of some metal cations (Cr⁶⁺/Fe³⁺/Hg²⁺), the interaction between the metal ions (Cr⁶⁺/Fe³⁺/Hg²⁺) and fluorescent CNPs and some other interesting and important features were studied theoretically in the framework of the widely used dense generalized density functional theory (DFT) approach. Quantum theory of atoms in molecules (QTAIM), NCI-plot-based reduced density gradient (RDG), electrostatic surface potential (ESP), and frontier molecular orbital (highest occupied orbital–lowest unoccupied molecular orbital) analyses were also performed to inspect some other useful structural, stability, and electronic features (*vide infra*). Moreover, in the present work, the quantum mechanical calculations were not only consistent with the experimental findings for the selective sensing of three metal ions (Cr⁶⁺, Fe³⁺, and Hg²⁺) interacting with a small unit of functionalized coronene as the CNP, but also facilitated a deep understanding and visualization of the CNP complexes associated with the probed metals.

2. Experimental

2.1. Materials

Madhuca longifolia flowers were collected from our institute CSIR-CIMFR Digwadih Campus, Dhanbad. Metal ions including cobaltous chloride, lead acetate trihydrate, manganese chloride, ferric chloride, mercury chloride, chromium oxide, aluminium chloride, zinc acetate, cadmium chloride, nickel chloride, silver nitrate, and magnesium acetate were purchased from Merck India. Whatman 42 filter paper was purchased from Sigma Aldrich.

2.2. Instrumentation

The size and morphology of the as-synthesized nanomaterials were determined using a Thermo Fisher Scientific Talos F200X G2 high-resolution transmission electron microscope (HRTEM). The morphology and elemental composition mapping were studied using a Zeiss Merlin VP compact field-emission scanning electron microscope (FESEM) and energy dispersive X-ray spectroscopy (EDS), respectively. The determination of various functional groups was done using a Shimadzu Iraffinity-1s Fourier transform infrared spectrophotometer. X-ray photoelectron spectroscopy for the characterization of surface functionality and elemental composition was carried out using a PHI 5000 versa probe III instrument. The count per second of different functional groups as a function of the BE (eV) was measured using a standard monatomic argon ion gun,



which was capable of generating 5 eV to 5 keV Ar ion beams. UV-vis electronic absorption and fluorescence emission spectra of the water-dispersed carbon nanomaterials were analyzed using an Agilent Carry 5000 UV-Vis-NIR and Hitachi F-7000 (150 W xenon lamp) fluorescence spectrophotometer, respectively. The zeta potential measurement was carried out using an Anton Paar Litesizer 500 system.

2.3. Preparation of carbon nanoparticles (CNPs) from *Madhuca longifolia* flower

The collected *Madhuca longifolia* flowers were washed with water to remove dust particles and allowed to dry for 30 min at 100 °C. Then 5.0 g of *Madhuca longifolia* flowers was mixed with 30 mL of double distilled water and transferred to a 50 mL Teflon-linked hydrothermal pressure vessel and heated at a temperature of 160 °C for 10 h. After completion of the reaction, the autoclaved was left to cool to room temperature and the reaction mixture was filtered through Whatman 42 filter paper. Further, the filtrate solution was centrifuged at 10 000 rpm for 5 min. Then, the supernatant solution was purified using a 1 kDa dialysis membrane for 24 h. The purified CNPs [6.0 mg mL⁻¹] were kept at 4 °C for further characterization and application.

In this green, economical and simple approach, no passivating agent or substrate was used. The CNPs were freely dispersed in an aqueous medium (Scheme 1).

2.4. Computational methodology

The capability of accurately describing metal-ligand interactions in various fields is demonstrated by the large number of calculations using the generalized DFT (B3LYP) method. The B3LYP functional of the generalized density functional theory was employed to acquire information on the geometrical optimization of the complexes formed by Cr⁶⁺, iron (Fe³⁺) and mercury (Hg²⁺) ions with the CNPs (*i.e.*, in this work, three types of complexes showing the selective features were examined theoretically), respectively. The initial configuration of all the CNP-based complexes was first passed through the minima (equilibrium) in the potential energy surface (PES), and then their frequencies were analyzed using the same level of theory, which resulted in all positive frequencies and showed the stable configurations of all the optimized complexes. Efforts were made to study the complexes using the dispersion-corrected DFT (DFT-D) approach, but the optimization process did not converge

constructively as expected, which can be attributed to the large structure of the complexes constructed from the functionalized coronene-based CNP model associated with the metal cations. Furthermore, to avoid the complexity involved in the DFT-D approach and to make the calculations easier and faster, together with reasonably precise outcomes, the B3LYP/6-31(d) level was employed as an alternative. To obtain the equilibrium structures (minimum energy configurations) of the sensing complexes and their constituting units, the Gaussian 09 package (electronic structure calculations) was used.⁴³ Some computational techniques [for instance, the BE/IE parameter, QTAIM tool, hydrogen bond strength based on the interaction coordinate (HBSBIC) approach, NCI-plot tool, compliance constant (CC), and relaxed force constant (RFC) parameters] to quantify the bond strength (bonding/nonbonding interactions) are frequently used to gain deep insights into the type, nature, and strength of various covalent and weak to moderate to strong NCIs.^{44–48}

3. Characterization of *Madhuca longifolia*-derived CNPs

3.1. Morphological characterization

The size and morphology of the CNPs prepared by the hydrothermal treatment of *Madhuca longifolia* flower were studied *via* high transmission electron microscopy (TEM). The synthesized CNPs showed a spherical morphology and were uniformly dispersed in the aqueous medium (Fig. 1a and b). The average diameter of the prepared CNPs was determined by the particle size distribution plot (inset, Fig. 1b), which revealed that they have an average core size of 12 nm.

Scanning transmission electron microscopy (STEM) and the corresponding energy dispersive X-ray spectroscopy (EDS) images were employed to identify the elemental composition and distribution of the carbon, nitrogen, and oxygen elements in the uniformly dispersed CNPs (Fig. 1c). The obtained result indicates that the CNPs are mainly composed of carbon, nitrogen, and oxygen [Fig. S1a in the ESI†]. The EDS mapping showed that there is a uniform distribution of C, O, and N throughout the CNPs.

X-ray photoelectron spectroscopy (XPS) and FTIR analysis were carried out to understand the nature of the functional groups present on the surface of the as-synthesized CNPs. Three peaks were observed at 284.0 eV, 399.5 eV, and 531.5 eV, which are attributed to C 1s, N 1s and O 1s with the atomic percentages



Scheme 1 Schematic representation for the one-step synthesis of the fluorescent CNPs *via* the hydrothermal treatment of *Madhuca longifolia*.



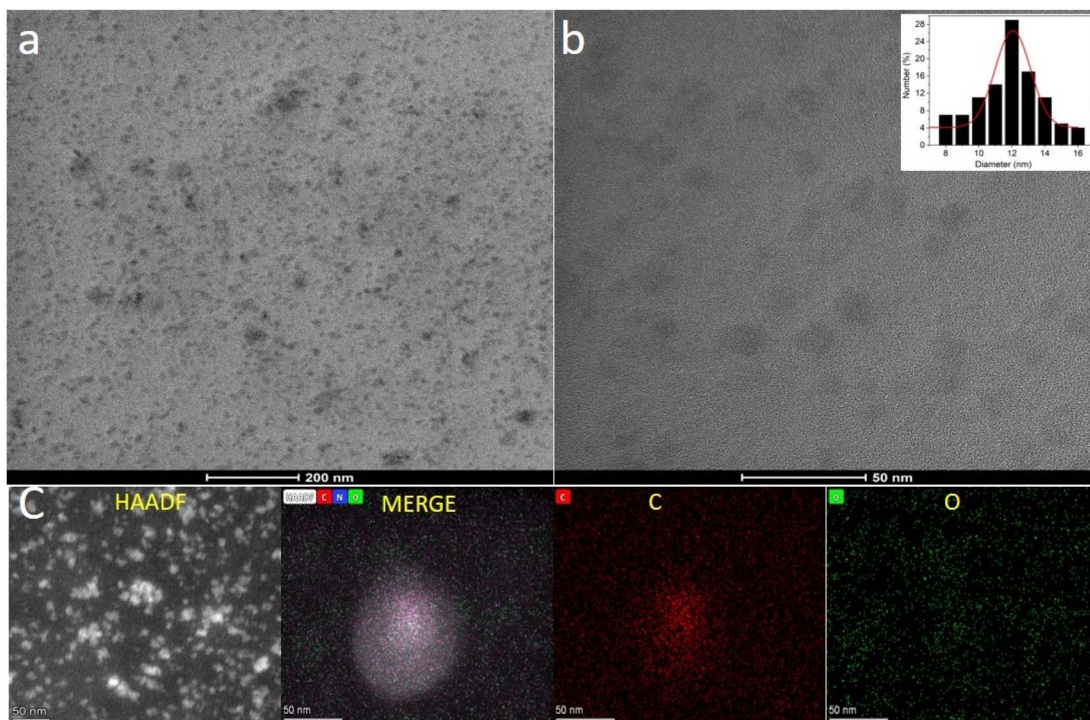


Fig. 1 TEM micrographs and dark-field HAADF-STEM images of CNPs obtained by the hydrothermal treatment of *Madhuca longifolia* flower: (a) lower magnification, (b) higher magnification and inset particle size distribution plot and (c) STEM and EDS mapping.

76.2%, 1.8% and 22.0%, respectively, in the full XPS survey spectrum of CNPs (Fig. 2a).

The full scan (high resolution) spectrum of C 1s and O 1s was obtained to determine the BE values of the various functional groups. The high-resolution scan of the C 1s region shows that carbon is present in three different chemical environments, corresponding to C=C at 283.9 eV, C-OH/C-C at 285.6 eV, and C=O at 287.7 eV (Fig. 2b). The deconvoluted O 1s spectrum shows peaks attributed to C=O at 531.0 eV, C-OH at 532 eV and O-C=O at 532.9 eV (Fig. 2c).

The FTIR analysis facilitated further insight into the surface functional groups present in the CNPs. This study revealed the presence of functional groups, such as a band at 3310 cm^{-1} corresponding to the stretching vibrations of C-OH and 1640 cm^{-1} for the C=O stretching vibration (Fig. 2d). The XPS and FTIR analyses suggest that the CNPs were functionalized with carbonyl, hydroxyl, and carboxylic acid groups.

The zeta potential (ζ) of CNPs was found to be -17.3 mV at neutral pH, which suggests the presence of acid functionalities on the surface of the synthesized CNPs (Fig. S1b, in the ES I[†]). The high negative zeta potential value confirms that the dispersion of CNPs in water is very stable. The presence of the aforementioned functional groups on the surface and the zeta potential of the CNPs confirmed their excellent dispersion and stability in water.

3.2. Photophysical study

The photophysical properties of the as-synthesized CNPs were studied in an aqueous medium at room temperature. Given that

the concentration of the CNPs in terms of molarity could not be calculated accurately, their absorbance was kept in the range of 0.2–0.7 at the excitation wavelength to avoid any inner filter effect. A clear transparent solution of 6.0 mg mL^{-1} CNPs showed absorption at around $250\text{--}550\text{ nm}$ (Fig. 3a, black) with two absorption bands at 271 nm and 330 nm , which can be assigned to the typical absorption of aromatic π systems or the $n\text{-}\pi^*$ and $\pi\text{-}\pi^*$ transition of nitrogen or oxygen-based groups and the trapping of excited-state energy by the surface states, respectively. The water-dispersed CNPs exhibited strong blue fluorescence with an emission maximum at 455 nm (Fig. 3a, blue) at 370 nm excitation. Moreover, the CNP solution displayed strong blue light under UV light exposure (Fig. 3a, inset).

To explore the excitation-dependent emission property of the water-dispersed CNPs, we scanned the samples at excitation wavelengths in the range of 360 to 520 nm (at 10 nm intervals) and collected their emission response from 350 to 650 nm (Fig. 3b). A prominent red-shift in the emission spectrum with the maximum intensity at 370 nm excitation was observed (Fig. 3b). The observed tunability in the normalized emission spectra (Fig. 3c) may be due to the presence of CNPs with different sizes and carboxylic groups on the surface of the CNPs, and structural inhomogeneity originating from the different surface trap states or surface defects during the hydrothermal synthesis of the CNPs from *Madhuca longifolia* flower. The presence of different types of aggregates even at a very low concentration may also contribute to the excitation wavelength dependence of the emission spectrum. The tri-exponential function-fitted fluorescence lifetime decay of CNPs is shown



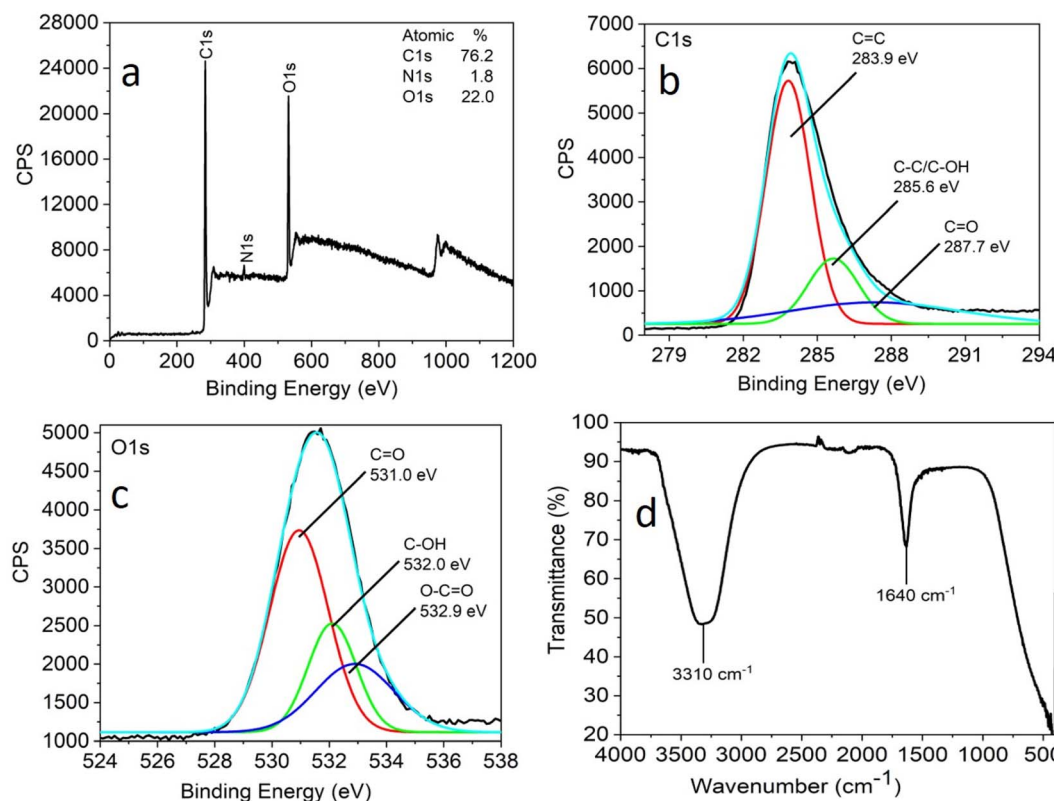


Fig. 2 Full scan XPS spectra (a), high-resolution XPS spectra (b) C 1s region and (c) O 1s region and FTIR spectra (d) of CNPs obtained by the hydrothermal treatment of *Madhuca longifolia* flower.

in Fig. 3d. The average fluorescence lifetime of CNPs was found to be 1.89 ns at a laser excitation of 405 nm.

4. Experimental results and discussion

4.1. Fluorescence sensing study

All the experiments were performed at room temperature in double-distilled water for the detection of Cr^{6+} ions and the concentration of the CNPs was fixed at 2.0 mg mL^{-1} . The fluorescence emission spectra were recorded in the absence and presence of an aqueous solution of Cr^{6+} (5–100 μM). It was observed that the fluorescence intensity of the CNPs was significantly quenched with an increase in the concentration of Cr^{6+} ions (Fig. 4a). The formation of a complex between Cr^{6+} and the CNPs led to quenching of their fluorescence intensity in the aqueous medium. The sensing characteristics or quenching efficiency of the fluorescent CNPs by the Cr^{6+} ion were best studied using the standard Stern–Volmer equation, as follows:

$$F_0/F = 1 + K_{\text{SV}}[Q] \quad (1)$$

where F_0 and F are the fluorescence intensities in the absence (only CNPs) and presence of the quencher (Cr^{6+}), respectively. K_{SV} and $[Q]$ are the Stern–Volmer quenching constant and quencher concentration, respectively. The Stern–Volmer quenching plot for the CNPs was obtained from a fluorescence

quenching titration experiment using Cr^{6+} at 454 nm (Fig. 4b). The plot of F_0/F versus quencher concentration showed good linearity ($R^2 = 0.99$) and the fluorescence intensity decreased with an increase in the concentration of Cr^{6+} . K_{SV} (slope of linear fit) was found to be $3.6 \times 10^3 \text{ M}^{-1}$ based on Fig. 4b (using eqn (1)). Given that the K_{SV} value is large, the quenching may be due to the static quenching (ground state complex formation) phenomenon. If the quenching is dynamic, the Stern–Volmer quenching constant is given by $K_D = k_q \times \tau_0$, where k_q is the bimolecular quenching constant and τ_0 is the lifetime of the CNPs (1.89 ns) in the absence of a quencher. k_q was found to be $1.90 \times 10^{12} \text{ M}^{-1} \text{ s}^{-1}$, which is much larger than the diffusion-controlled quenching value ($1 \times 10^{10} \text{ M}^{-1} \text{ s}^{-1}$), usually indicating some type of binding interaction.

To get a complete scenario of the quenching mechanism, the fluorescence lifetime of CNPs with and without Cr^{6+} was studied because fluorescence lifetime decay is one of the most specific methods to discriminate static and dynamic quenching. In our lifetime experiment, there was almost no change in the average lifetime of the CNPs after the addition of 100 M of Cr^{6+} (Fig. S2 in the ESI†), which confirms a purely ground-state (static quenching) quenching phenomenon.

4.2 Quenching mechanism

Given that Cr^{6+} has a high affinity to form a complex with carboxylate groups,⁴⁹ the formation of a complex between the Cr^{6+} ion and acidic group of the CNPs may lead to a decrease in



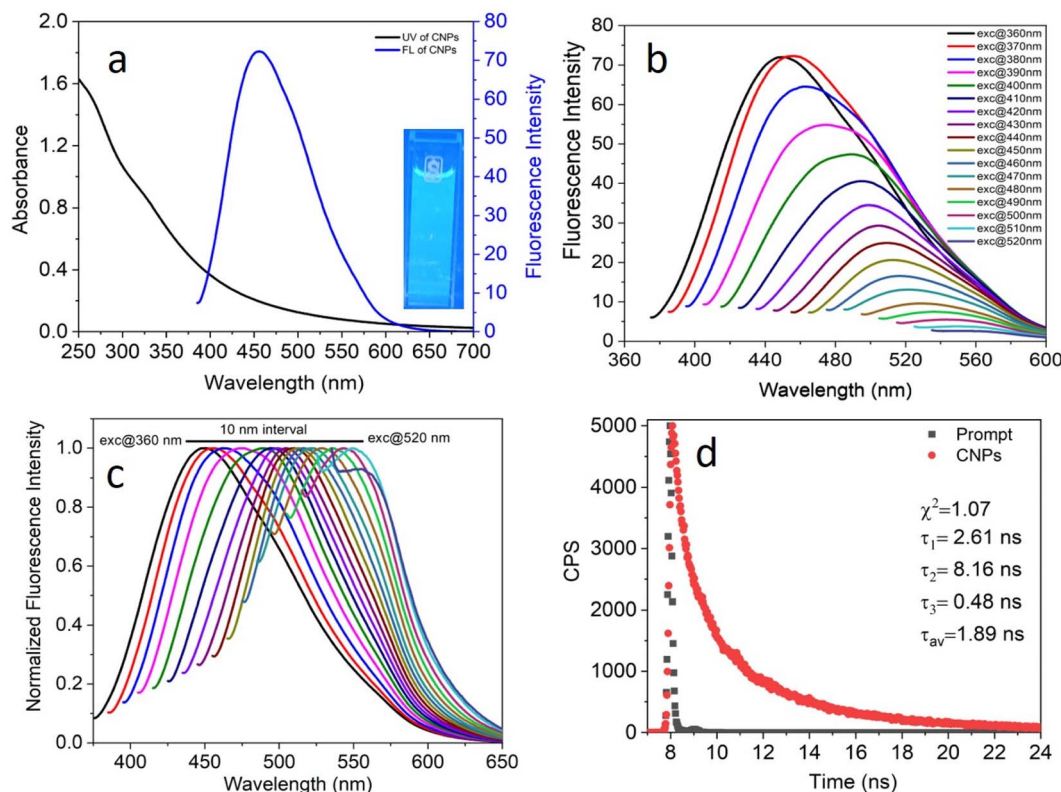


Fig. 3 (a) UV-visible absorption and emission spectra with the inset showing blue fluorescent image ($\lambda_{\text{ex}} = 370 \text{ nm}$ and $\lambda_{\text{em}} = 455 \text{ nm}$), (b) excitation-dependent emission spectra of CNPs, (c) normalized emission spectra of CNPs in an aqueous medium $[\text{CNP}] = 2.0 \text{ mg mL}^{-1}$ and (d) time-resolved fluorescence decay of CNPs ($\lambda_{\text{exc}} = 405 \text{ nm}$ and $\lambda_{\text{emi}} = 490 \text{ nm}$).

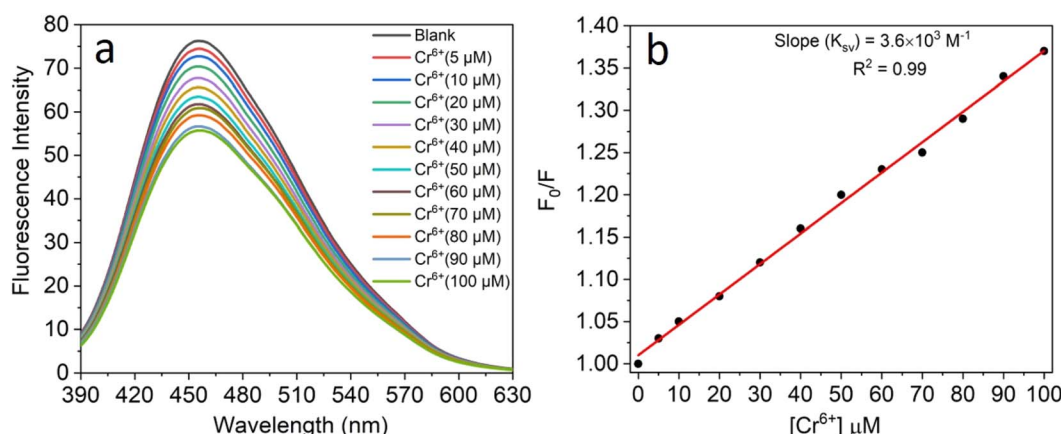


Fig. 4 (a) Fluorescence spectra of CNPs (2.0 mg mL^{-1}) in the presence of different amounts of Cr^{6+} from the top to bottom: 5, 10, 20, 30, 40, 50, 60, 70, 80, 90, and 100 μM in double-distilled water ($\lambda_{\text{exc}} = 370 \text{ nm}$) and (b) plot of F_0/F against the concentration of Cr^{6+} (Stern–Volmer quenching response) at 454 nm.

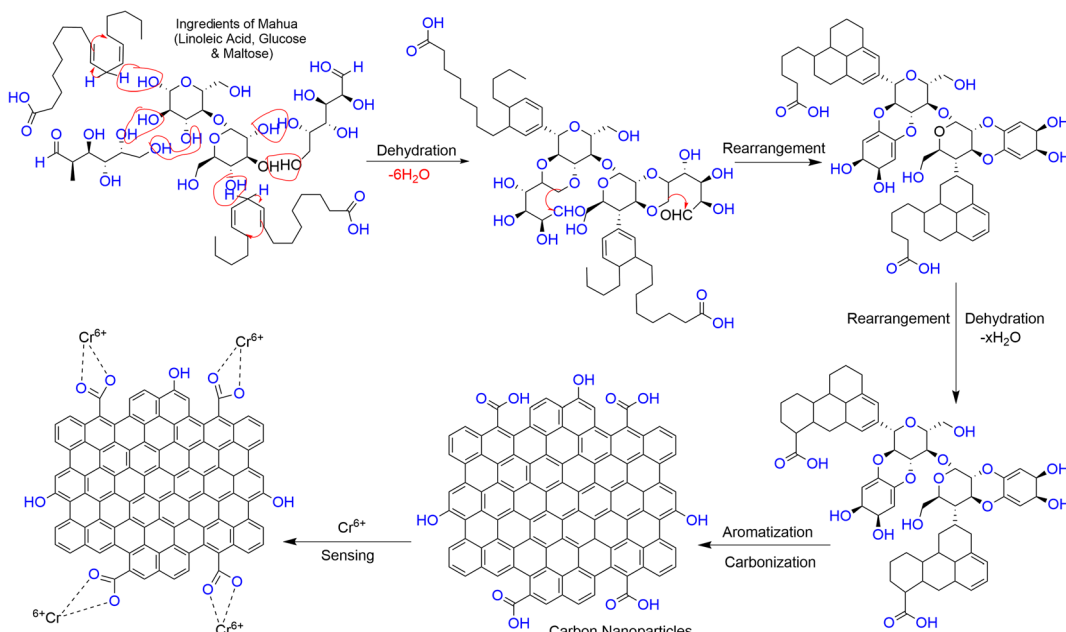
the fluorescence intensity of CNPs through charge transfer (CT) quenching. A schematic diagram for the plausible quenching mechanism is depicted in Scheme 2.

4.3 Selective and sensitive detection of the $\text{Cr}(\text{vi})$ ion

The selectivity towards the detection of the Cr^{6+} ion was verified by carrying out experiments in the presence of other heavy

metal ions (number of samples, $n = 5$). In this study, a variety of environmentally relevant heavy metal ions was chosen in aqueous medium including Co^{2+} , Pb^{2+} , Mn^{2+} , Fe^{3+} , Hg^{2+} , Cr^{6+} , Al^{3+} , Zn^{2+} , Cd^{2+} , Ni^{2+} , Ag^{2+} , and Mg^{2+} . The fluorescence intensity ratio plot for the CNPs (2.0 mg mL^{-1}) in the presence of 100 μM metal ions in double-distilled water was recorded. Fig. 5a shows the selectivity for Cr^{6+} , where about a 29% decrease in the





Scheme 2 Plausible representation of the formation of CNPs via the hydrothermal treatment of the *Madhuca longifolia* flower and their formation of a complex with the Cr^{6+} ion.

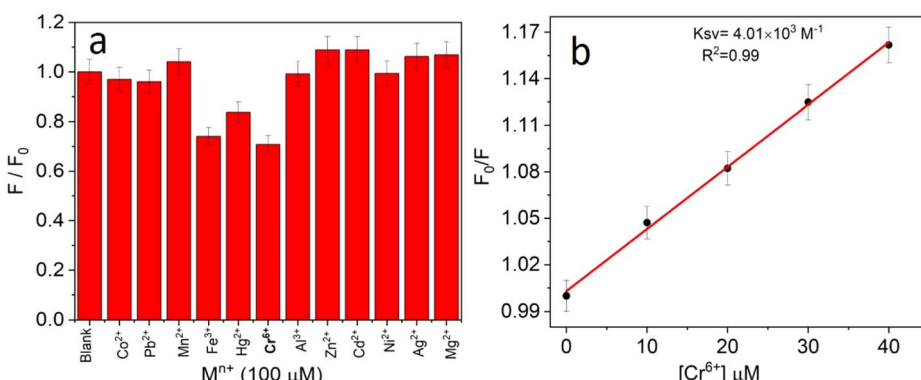


Fig. 5 (a) Fluorescence intensity ratio plot for CNPs in the presence of other heavy metal ions with p -value < 0.05 . (b) Calibration plot, CNPs (2.0 mg mL^{-1}) against Cr^{6+} ($0\text{--}10 \mu\text{M}$) ion concentration for the estimation of LOD in the aqueous solution [standard deviation = ± 0.043 , $[\text{M}^{n+}] = 100 \mu\text{M}$, $[\text{CNPs}] = 2.0 \text{ mg mL}^{-1}$, $\lambda_{\text{exc}} = 370 \text{ nm}$ and $\lambda_{\text{emi}} = 454 \text{ nm}$].

fluorescence intensity was observed in the presence of Cr^{6+} ($100 \mu\text{M}$). However, a slight decrease and increase in the fluorescence intensity ratio were also observed (Fig. 5a) in the presence of other metal ions, mainly Fe^{3+} (25% reduction), Hg^{2+} , Mn^{2+} , Zn^{2+} , and Co^{2+} , respectively. The p -value is less than 0.05, suggesting that the result of this test is statistically significant. The enhancement in the fluorescence intensity after the addition of Mn^{2+} , Zn^{2+} and Co^{2+} may be due to aggregation-induced emission enhancement (AIEE) or coordination-induced aggregation (CIA). When aggregation occurs, the intramolecular motion is restricted, which blocks the nonradiative path and activates radiative decay.

There are numerous studies in the literature where a slight variation in the fluorescence intensity was found at the micromolar level, and calculations were made based on the observed

variation. Employing the change in the fluorescence intensity in the micromolar range, the limit of detection (LOD) from the standard deviation (S.D.) and the slope (m) of the calibration plot (Fig. 5b) were calculated using eqn (2) (number of samples, $n = 5$).

The linear plot of the fluorescence intensity ratio *versus* Cr^{6+} concentration (Fig. 5b) is useful to calculate the LOD knowing the standard deviation (S.D.) and the slope of the plot.

$$\text{LOD} = \frac{3 \times \text{SD}}{m} \quad (2)$$

It was found that the *Madhuca longifolia*-derived fluorescent CNPs have a reasonable LOD (103 ppb), which is almost comparable with the maximum allowed limit of 0.1 mg L^{-1} (100

Table 1 Comparison of the LOD value for Cr^{6+} sensing in the present work with previously reported nanosensors

S. no.	Nanosensor	LOD (ppb) for Cr^{6+}	References
1	Carbon dots	109	Li <i>et al.</i> ⁵⁰
2	Carbon dots	560	Wang <i>et al.</i> ⁵¹
3	Carbon dots	120	Wang <i>et al.</i> ⁵²
4	N-doped carbon dots	216.30	Wang <i>et al.</i> ⁵³
5	Carbon dots	520	Yang <i>et al.</i> ⁵⁴
6	N-doped carbon dots	207	Bardhan <i>et al.</i> ⁵⁵
7	Carbon dots	181	Sakaew <i>et al.</i> ⁵⁶
8	Carbon dots	17 600	Yaun <i>et al.</i> ⁵⁷
9	Titanate and cellulose nanofibers carbon dots	8500	Luo <i>et al.</i> ⁵⁸
10	Lignin-based hydrogel with cellulose nanofibers and carbon dots	11 200	Yaun <i>et al.</i> ⁵⁹
11	N-doped carbon dots	166	Bogireddy <i>et al.</i> ⁶⁰
12	Terylene carbon dots	150	Hu <i>et al.</i> ⁶¹
13	Carbon dots	1700	Anil <i>et al.</i> ⁶²
14	N-doped carbon dots	709.3	Chen <i>et al.</i> ⁶³
15	Carbon dot-based fluorescent hydrogel	147	Feng <i>et al.</i> ⁶⁴
16	Carbon dots	124	Goswami <i>et al.</i> ⁴⁰
17	N-doped carbon quantum dots	109	Wang <i>et al.</i> ³⁸
18	Carbon nanoparticles (CNPs)	103	Our current work

ppb) by the U.S. Environmental Protection Agency (EPA). Furthermore, the LOD for Cr^{6+} sensing in the present study is much better than that in most of the previously reported works in recent years (Table 1).

5. Computational results and discussion

Given that all-inclusive atomistic features (including the atomic/molecular environment) play a dominant role in altering the inter- or intramolecular (metal–nonmetal and nonmetal–nonmetal covalent/ionic bonding, H-bonding, vdW, *etc.*) interactions, the sensing feature of the probed species was inspected in the framework of the DFT approach. Herein, among the twelve experimentally probed metal cations (Co^{2+} , Pb^{2+} , Mn^{2+} , Fe^{3+} , Hg^{2+} , Cr^{6+} , Al^{3+} , Zn^{2+} , Cd^{2+} , Ni^{2+} , Ag^{2+} , and Mg^{2+}), the three most selective and high sensitivity feature environmentally relevant heavy metal cations (Cr^{6+} , Fe^{3+} , and Hg^{2+}) were chosen for the quantum chemical calculations. A thorough and organized structural, stability/energetic (*i.e.*, diagnostic of the sensitivity feature using the BE parameter), and electronic feature analysis was performed employing the widely used DFT-based quantum mechanical calculation approach. Interestingly, and importantly, a comprehensive theory-based study on the above-mentioned interactions can be viewed in a few previous reports, portraying the structural, energetic/stability, and electronic features of the vastly used polyacrylamide (PAM) and hydroxyapatite (HAP)-based bio-composite models, where the experimental outcomes were supported by the employed *in silico* approach.^{65,66}

5.1 Characterization of structural, stability/energetic, and electronic features

A comprehensive study on the characterization and validation of the structural (optimized parameters such as bond length

and bond angle), (HOMO, LUMO, and HOMO–LUMO gap), sensing and selectivity features (in terms of the BEs and visualization of metal–nonmetal and other probable (nonbonding) interactions using the NCI-RDG plot *via* 3D-isosurface and 2D-scatter plots), and electronic properties (HOMO–LUMO 3D-isosurface maps, MESP, and QTAIM topological picture and parameters) of a free CNP moiety and its associated complexes with the metal ions (Cr, Fe, and Hg cations) is discussed in the following section.

5.2 Free functionalized CNP constituent

5.2.1 Structural and electronic feature analyses. We begin the discussion by choosing the parent framework as the CNP designed by using the molecular modeling approach as a theoretical model, and then the electronic structure calculations were executed using the DFT approach. Initially, to inspect the binding interaction (sensing feature) between the metal ion and the CNP, a total of six COOH and two OH groups was chosen during modelling the isolated parent CNP structure. The optimized (minimum energy/equilibrium) structure of the free functionalized CNP can be seen in Fig. 6a with proper atomic labeling. The equilibrium (optimized) structure of the CNP shows that the C–C bond distance of its central ring (*i.e.*, benzene ring) ranges from 1.429 Å to 1.437 Å. Substitution of the COOH and OH group is supposed to slightly distort the optimized parameters of the outer six benzene rings. Moreover, the bond distance of the C–H bonds of some of the outer benzene rings of the CNP ranges from 1.077 Å to 1.080 Å. The two OH bonds constructing H-bonds have a bond length of 1 Å. The C=O bond lengths of all six COOH functional groups range from 1.209 Å to 1.239 Å. Importantly, two O–H bonds and four O–H bonds of the COOH group have the bond lengths of 0.976 Å and 0.977 Å, respectively. Because of the presence of multiple COOH groups, many intramolecular H-bonding NCIs were probed using the QTAIM and NCI-plot tools, which show that



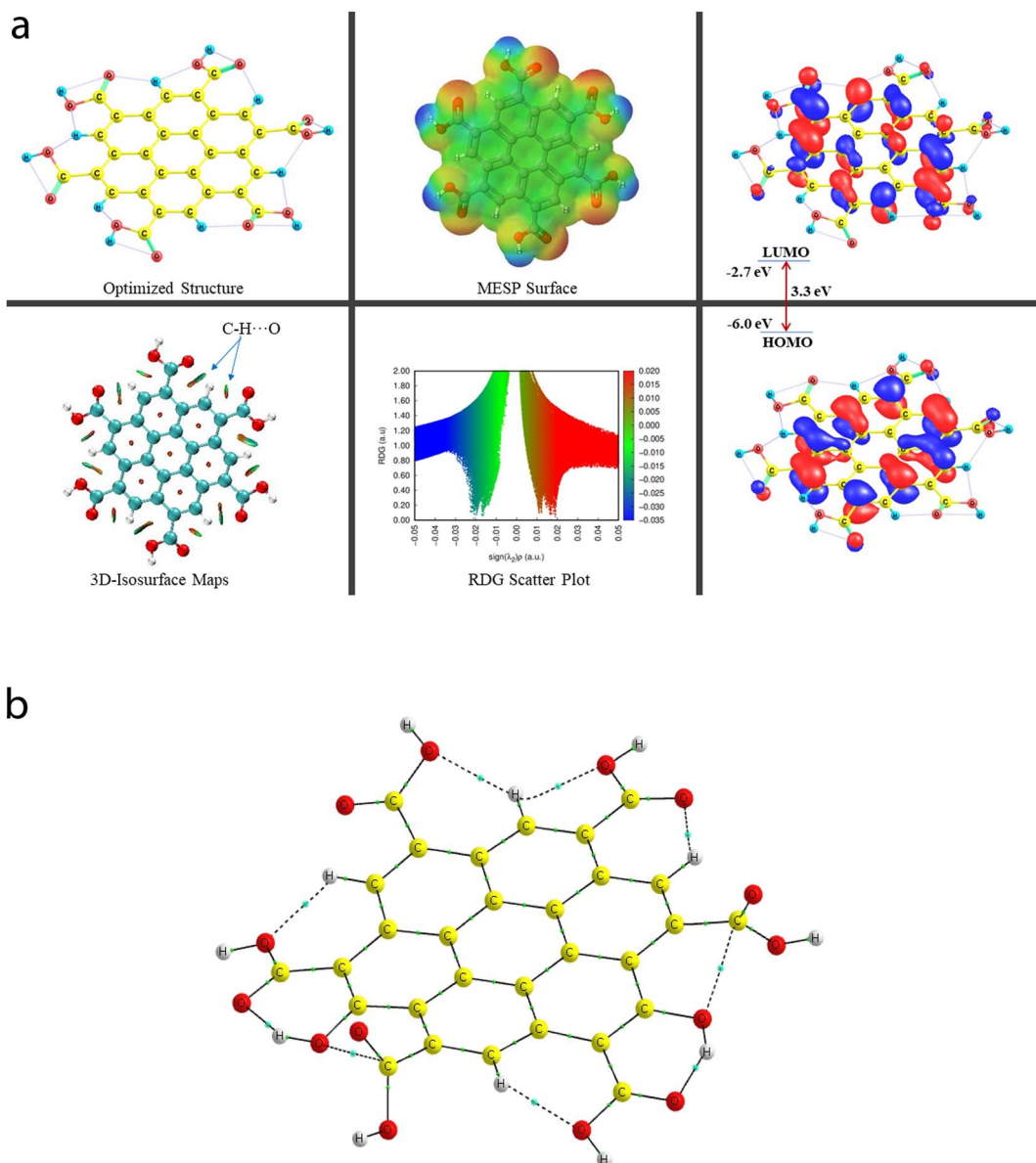


Fig. 6 (a) Optimized structure, MESP surface, HOMO–LUMO 3D maps, 3D-isosurface maps, and RDG scatter plot of the free functionalized CNP constituent model. (b) QTAIM molecular graph of the free functionalized CNP constituent model.

these (a total of two moderate O–H···O and five weak C–H···O) interactions take place, and also illustrate that two non-conventional bond paths (NBPs) (C···O) stabilize the CNP, as shown in Fig. 6b. The O–H···O H-bond distances were calculated to be 1.599 Å and 1.597 Å, respectively. The corresponding bond angles were found to be 147.3° and 147.6°, respectively. The distance between two atoms for one NBP (C29···O55) is 2.597 Å, whereas it was computed to be 2.573 Å for the second NBP (C45···O53). In seven types of NCIs (excluding both C···O NBPs), the O atom acts as a proton acceptor, whereas two O atoms and five C atoms act as proton donors in all the probed NCIs.

The energy difference between the two frontier molecular orbitals (FMOs), *i.e.*, highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO), is

denoted as the HOMO–LUMO energy gap (E_{gap}). The strength, stability, and chemical behavior of any chemical/molecular species are described by the E_{gap} value. For example, a system having a small E_{gap} value generally shows higher reactivity (*i.e.*, less stability), and thus shows good optical features (*i.e.*, easy photochemical excitation possible) because of the lower energy gap (bandgap) between the valence and conduction bands. Moreover, the electron-accepting and electron-donating capabilities of a particular species are linked to its E_{HOMO} and E_{LUMO} values, respectively. The E_{HOMO} and E_{LUMO} values for the free functionalized CNP species were calculated to be –6.0 eV and –2.7 eV, respectively, and the associated E_{gap} value is 3.3 eV, which are presented in Table 2. Interestingly, Fig. 6a presents the HOMO–LUMO 3D isosurface maps of the free functionalized CNP constituent, where the HOMOs (in the plane) are

Table 2 The DFT (B3LYP/6-31G(d)) calculated binding energies (in kcal mol⁻¹) of TM@CNPs (TM: Cr⁶⁺, Fe³⁺, and Hg²⁺)

Species	BE (kcal mol ⁻¹)	BSSE complexation energy (kcal mol ⁻¹)	HOMO (eV)	LUMO (eV)	<i>E</i> _{gap} (eV)
Free functionalized CNP	—	—	-6.0	-2.7	3.3
Cr ⁶⁺ @CNPc	-4395.1	-4759.8	-23.0	-21.5	1.5
Cr ⁶⁺ @CNPt	-4476.7	-4688.9	-23.2	-22.0	1.2
Fe ³⁺ @CNPc	-696.5	-753.1	-15.1	-13.8	1.3
Fe ³⁺ @CNPt	-794.4	-785.6	-15.0	-13.8	1.2
Hg ²⁺ @CNPt	-183.9	-226.2	-11.6	-11.2	0.4

mainly positioned over almost all the six-membered rings of the free functionalized CNP moiety and the LUMOs follow a similar pattern, but the orbital phases are out-of-plane. This indicates the occurrence of a π to π^* transition in the free functionalized CNP framework.

The molecular size, shape, and charge distribution are concurrently described by the molecular electrostatic potential surface (MESP) map, which is widely used to understand the NCI(s), ways for the recognition of one molecule by another, relative polarity, and electrophilic and nucleophilic attack, reactive nature, *etc*. The MESP surface is displayed in Fig. 6a, where the reactive sites such as electron-poor region (high probability of nucleophilic attack) are shown by the absolute positive charge distribution indicated in a blue colour scheme. The neutral regions are exemplified by a green colour scheme, while the strong negative region (shown in the red colour scheme) can be viewed by the absolute negative charge distribution, indicating the electrophilic attack. The positive regions (favorable for nucleophilic attack) are positioned over the H-atoms of the COOH groups, and the negative regions (preferable for electrophilic attack) are located over the O atoms of the COOH groups of the free functionalized CNP constituent.

5.2.2 NCI-RDG isosurface plots and QTAIM analyses. The NCI-reduced density gradient (NCI-RDG) tool allows an all-encompassing portrayal of the H-bonding, extremely weak vdW interactions, and steric repulsion engaged in the probed complexes. Here, in this work, the NCI-RDG approach was capable of producing a graphical view of the NCI regions considering the nature and strength of the NCIs. Generally, strong attractive interaction(s) such as metal–nonmetal bonding and strong H-bonding are described by localized blue lentils. However, moderate to weak H-bonding is shown by a bluish-green to green colour scheme. The extremely weak vdW interactions are illustrated by thin and delocalized green regions, whereas the steric clashes are exposed by the red isosurfaces. Complete information about all three colour schemes can be realized employing an example of the functionalized CNP model (see Fig. 6a).^{65,66} The horizontal axis on the scale of 0.05 in the range of -0.05 to +0.05 shows the RDG isosurface, whereas the scale ($\mathcal{Q}(r)$ values) on the vertical axis is in the range of -0.035 atomic unit (au) to +0.02 au, where the colored isosurfaces of the species can be viewed on the blue-green-red scale. The higher density values ($\mathcal{Q}(r) < 0$) indicate stronger attractive interactions, whereas the repulsive interactions are shown by very low-density values ($\mathcal{Q}(r) > 0$).

For instance, the mixed blue-green (light bluish) colour peak (values ranging from -0.02 au to -0.025 au in the 2D scatter plot) shows the H-bonding interactions, which can also be confirmed by the light-bluish (mixed blue-green) disc-shaped 3D-isosurface maps shown in the seven NCIs (two O–H···O and five C–H···O H-bonding interactions) (see Fig. 6a). These attractive interactions give a vibrant sign of the existence of intramolecular HBs, which indeed play a prime role in stabilizing the functionalized CNP itself. The extremely weak vdW interactions are validated by the 2D scatter plot (green colour peaks lying between -0.015 au and -0.02 au) and the 3D isosurface maps (green colour lentils between some C and O atoms). The low-gradient broad peaks exposed in the red colour scheme and located at the positive side (values ranging from +0.01 au to +0.02 au in the 2D scatter plot) show the existence of a steric effect as an indication of repulsive interaction, which was also confirmed by the 3D isosurface maps. This effect represents the electron density depletion, which is due to the presence of electrostatic repulsion.

To confirm that the NCIs play an important role in the sensing features, the QTAIM tool was employed for all the sensing model complexes together with the free COOH-functionalized CNP constituents. Several parameters such as ρ (electron density), $\nabla^2(\rho)$ (Laplacian of electron density), V (local potential energy density), and delocalization index (DI) were used to quantify the strength of the NCIs and the NBPs. The bond path lengths (BPLs) acquired from the QTAIM tool appeared to be consistent with the bond length (BL) obtained from the optimized parameters using the Gaussian package, which can be verified in Table 3.

A total of nine [seven NCIs (two O–H···O and five C–H···O HBs) and two NBPs (C···O)] can be visualized in Fig. 6b, which are represented by the bond paths (BPs) shown in dotted/solid lines (black color), where small green color spheres are displayed between two interacting atoms (O and H atoms for the HBs and C and O atoms for the NBPs). A clear QTAIM molecular graph (with proper atomic labeling and numbering) of the free functionalized CNP constituent can be seen in Fig. S3 in the ESI.† As the BLs of the H-bonding interactions increase (ranging from 1.597 Å to 2.234 Å), the corresponding ρ values decrease (range of 0.0175 au to 0.0606 au) as expected. The strongest H-bonding interaction (ρ : 0.0606 au, $\nabla^2(\rho)$: +0.1930 au, V : -0.0565 au, and DI: 0.131) was detected as the O50···H54 bond, whereas the weakest NCI is the O43···H22 HB (ρ : 0.175 au, V : -0.0140 au, and DI: 0.04) having the $\nabla^2(\rho)$ value +0.0811 au, which is greater



Table 3 Some useful QTAIM parameters of the NCIs (H-bonding) and nonconventional bond paths (NBPs) for the free functionalized CNP constituent model

BP	BL (Å)	BPL (Å)	ρ (au)	$\nabla^2(\rho)$ (au)	V (au)	DI (A, B)
Noncovalent interaction (NCI) (intramolecular)						
O50···H54	1.597	1.622	0.0606	+0.1930	-0.0565	0.131
O34···H56	1.599	1.624	0.0604	+0.1922	-0.0563	0.131
O35···H17	2.057	2.087	0.0230	+0.0878	-0.0190	0.064
O51···H18	2.064	2.088	0.0227	+0.0869	-0.0188	0.066
O42···H26	2.077	2.099	0.0233	+0.0810	-0.0191	0.072
O39···H22	2.132	2.158	0.0201	+0.0743	-0.0163	0.058
O43···H22	2.234	2.411	0.0175	+0.0811	-0.0140	0.04
Nonconventional bond path (NBP) (intramolecular)						
C45···O53	2.573	2.583	0.0178	+0.0687	-0.0139	0.044
C29···O55	2.597	2.609	0.0172	+0.0664	-0.0133	0.041

than that of the second weakest O39–H22 HB. The QTAIM parameters (ρ , $\nabla^2(\rho)$, V , DI) for two NBPs, C45···O53 and C29···O55, were computed to be (0.0178, +0.0687, −0.139, 0.044) and (0.172, +0.0664, −0.0133, 0.041) (in au), respectively, which also stabilize the CNP constituent model. These strength estimating parameters correspond to the analyzed BLs and BPLs, as expected.

5.3 Cr⁶⁺@CNP, Fe³⁺@CNP, and Hg²⁺@CNP complexes

5.3.1 Structural, sensing, and electronic feature analyses. To acquire the maximum sensing feature in terms of the maximum number of metal–nonmetal interactions (MNIs) between the metal cation (Cr⁶⁺/Fe³⁺/Hg²⁺) and the functionalized CNP moiety, careful fine-tuning was executed during the optimization and frequency calculations, showing all minima in the potential energy surface (PES), followed by the molecular modelling approach. Considering the experimental facets and to gain deep insights into the sensing features, the molecular modelling approach was employed in two ways. A small unit of a modified CNP (here, functionalized coronene) was chosen as the model, which appears to show two binding sites. Firstly, the functionalized CNP can potentially complex with the TM (Cr⁶⁺/Fe³⁺/Hg²⁺) lying at the terminal of the functionalized CNP and forms the TM@CNPt complex, followed by facilitating minima during the optimization process, where ‘t’ refers to the terminal position. Secondly, the TM, Cr⁶⁺/Fe³⁺, is positioned at the top of the central six-membered ring of the CNP framework during the formation of the TM@CNPc complex, where ‘c’ refers to the top of the centre. The optimized structures of the two corresponding Cr⁶⁺@CNP supramolecular complexes are displayed in Fig. 7a (Cr⁶⁺ at the top of the center) and Fig. 8a (Cr⁶⁺ at the terminal) together with their MESP surfaces, HOMO–LUMO isosurfaces, NCI-2D scattered plot, and 3D isosurfaces. A few chosen optimized parameters (such as BLs and BPLs) and electronic feature-based parameters of metal–nonmetal interactions (MNIs), NCIs, and NBPs for different complexation modes (TM@CNP) between the CNP constituent unit and the metal cation (Cr⁶⁺) are summarized in Tables 4 and 5 for the Cr⁶⁺@CNPc and Cr⁶⁺@CNPt complexes, respectively. The

QTAIM molecular graphs (showing the MNIs, NCIs, and NBPs) of Cr⁶⁺@CNPc and Cr⁶⁺@CNPt can be viewed in Fig. 7b and 8b, respectively. Importantly, clear pictures of the QTAIM molecular graphs of Cr⁶⁺@CNPc and Cr⁶⁺@CNPt (with proper atomic labeling and numbering together with their BPs) can be seen in Fig. S4 and S5 of the ESI,[†] respectively.

The QTAIM analyses (*vide infra*) revealed that the Cr metal appears to associate with two C atoms (Cr53–C4 and Cr52–C9) of the central benzene ring of the CNP in the case of the Cr metal ion located at the top of the center of the CNP (Cr⁶⁺@CNPc complex) (see Fig. 8b) and the bond distances were detected to be 2.215 Å and 2.298 Å, respectively. Structural changes in the CNP can be seen in the Cr⁶⁺@CNPc complex, which is due to incorporation of the Cr⁶⁺ ion possessing the highest oxidation state with a smaller size and higher positive charge, thus altering the structural, sensing and selectivity feature (enhanced) and electronic features. A increase in the C–H bond distances in the range of 1.087 Å to 1.09 Å was observed when the Cr metal cation is associated with the CNP (in free functionalized CNP: C–H bond range from 1.077 Å to 1.080 Å). However, interestingly and importantly, another configuration of the Cr metal cation with the CNP (Cr⁶⁺@CNPt complex) provides more favorable outcomes because its larger BE than the Cr⁶⁺@CNPc complex (*vide infra*). A total of three MNIs (two Cr–O and one Cr–H) can be discerned in Fig. 7b. The bond distances of the two MNIs, Cr–O were analysed to be 1.979 Å (Cr51–O37) and 1.988 Å (Cr51–O41) and the bond length of the Cr–H bond was computed to be 2.029 Å. It is noteworthy to mention that the higher oxidation state (with highly positive charge) and smaller size of the Cr⁶⁺@CNP complex (both cases, Cr⁶⁺@CNPc and Cr⁶⁺@CNPt) are responsible for the larger degree of deformation of the CNPs.

In the case of the Fe³⁺@CNPc complex, one bond (MNI) between the Fe metal ion and one of the C-atoms of the central benzene ring of the CNP was detected using the QTAIM tool (*vide infra*), where the bond length of the Fe57–C3 bond was analysed to be 2.336 Å. Moreover, the equilibrium (optimized) structures of the Fe³⁺@CNPc complex can be seen in Fig. S6[†] (structural/electronic feature in Fig. S6a[†] and QTAIM molecular graph in Fig. S6b[†] for Fe³⁺@CNPc) and Fig. S7[†] (geometrical/electronic features in Fig. S7a[†] and QTAIM molecular graph in Fig. S7b[†] for Fe³⁺@CNPt) in the ESI.[†] Some useful and selected structural [BL and bond path length (BPL)] and QTAIM-based parameters (such as ρ , $\nabla^2(\rho)$, V , and DI indices) for Fe³⁺@CNPc and Fe³⁺@CNPt can be viewed in Tables S1 and S2 of the ESI,[†] respectively.

However, surprisingly, only one configuration of the Hg²⁺-related complex, Hg²⁺@CNPt (Hg²⁺ at the terminal), can be viewed in the ESI (see Fig. S8a[†]). Also, the electronic feature-based picture and QTAIM molecular graph of the Hg²⁺@CNPt complex are displayed in Fig. S8a and b,[†] respectively. The structural (BL and BPL) and QTAIM-based (ρ , $\nabla^2(\rho)$, V , and DI) parameters of the Hg²⁺ ion (probed at the terminal position only) can be seen in Table S3 of the ESI.[†] The bond distances of two probed MNIs of Hg–O were computed to be 2.943 Å (Hg53–O43) and 3.153 (Hg53–O39), which are much larger than that of the Fe–C bond (2.336 Å) of the Fe³⁺@CNPc complex.



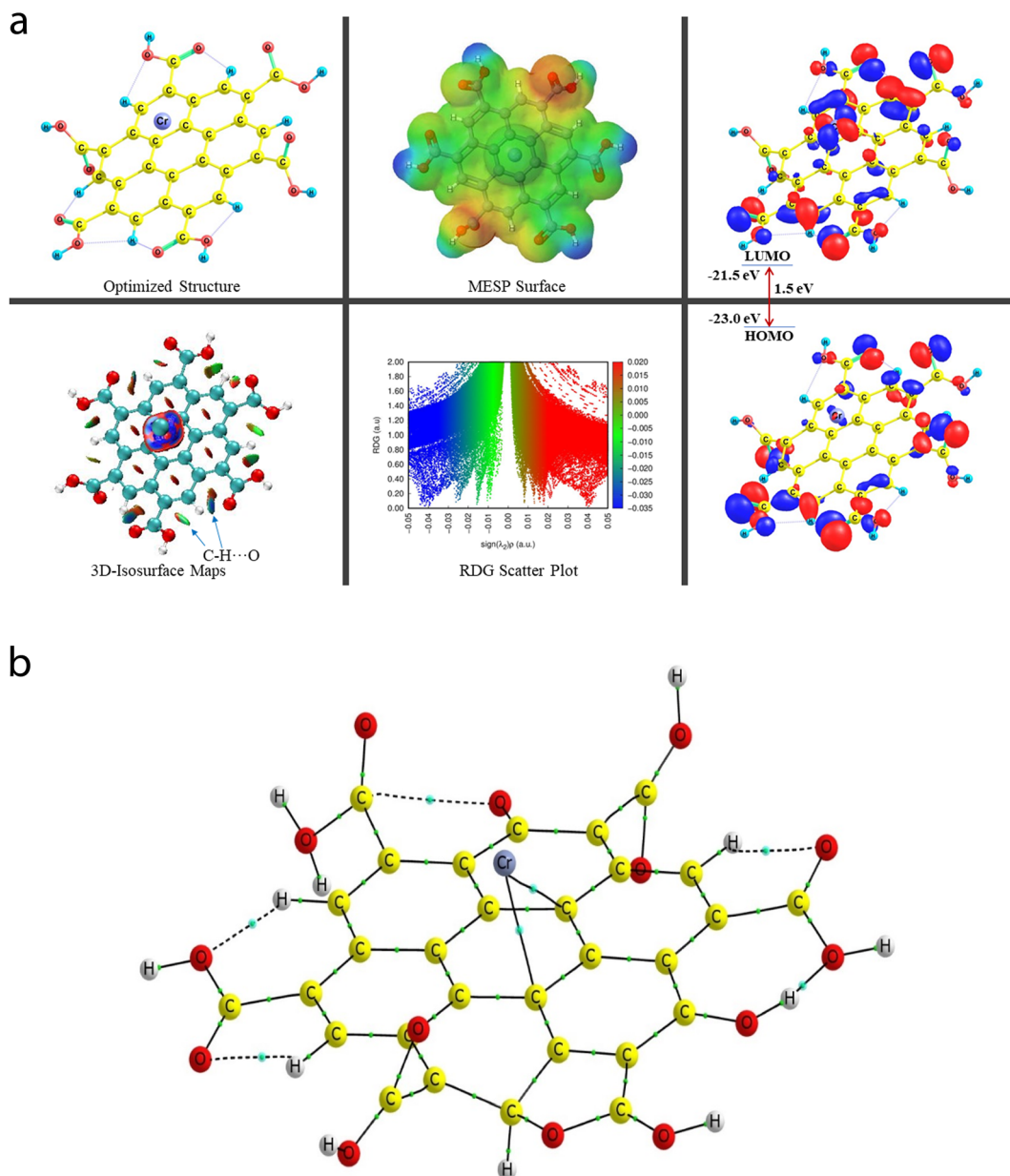


Fig. 7 (a) Optimized geometric structure, MESP surface, HOMO–LUMO 3D maps, 3D-isosurface maps, and RDG scatter plot of the Cr⁶⁺ ion interacting with the functionalized CNP to form the Cr⁶⁺@CNPc complex in the center of the CNP moiety. (b) QTAIM molecular graph of the Cr⁶⁺ ion interacting with the functionalized CNP to form the Cr⁶⁺@CNPc complex in the center of the CNP moiety.

The calculated BE values for the TM@CNPt complexes associated with the terminal TMs, *i.e.*, Cr⁶⁺, Fe³⁺, and Hg²⁺ ions, were calculated to be $-4476.7\text{ kcal mol}^{-1}$, $-794.4\text{ kcal mol}^{-1}$, and $-183.9\text{ kcal mol}^{-1}$, whereas the analyzed BEs of the CNP complex with the central Cr⁶⁺ and Fe³⁺ metal ions forming the Cr⁶⁺@CNPc and Fe³⁺@CNPc complexes are $-4395.1\text{ kcal mol}^{-1}$ and $-696.5\text{ kcal mol}^{-1}$, respectively. A high BE difference can be realized among all three cations, which appear to be due to the charge and size of the TM cations. Given that the Cr⁶⁺ ion consists of the highest charge and has the lowest size among the three cations, it is expected to show the highest BE with a large difference, revealing that it is a more selective and the most

sensitive complex. A comprehensive DFT study on the interaction between different valence metal chromium ions (Cr⁶⁺, Cr³⁺, and Cr²⁺) and reduced glutathione (GSH) was reported by Jianhua *et al.* in 2013, where they showed that the BEs for GSH associated with the Cr²⁺, Cr³⁺, and Cr⁶⁺ metal ions are $-834.8\text{ kJ mol}^{-1}$ ($199.5\text{ kcal mol}^{-1}$), -1162 kJ mol^{-1} ($277.7\text{ kcal mol}^{-1}$), and $-17568.6\text{ kJ mol}^{-1}$ ($-4199\text{ kcal mol}^{-1}$), respectively.⁶⁷ Here, the emphasis is to compare the selective and sensitivity feature (in terms of the BE value of $-4199\text{ kcal mol}^{-1}$) of the Cr⁶⁺ ion interacting with the GSH skeleton (Cr⁶⁺@GSH complex), which is $277.7\text{ kcal mol}^{-1}$ and $196.1\text{ kcal mol}^{-1}$ smaller than that of the probed Cr⁶⁺@CNPt



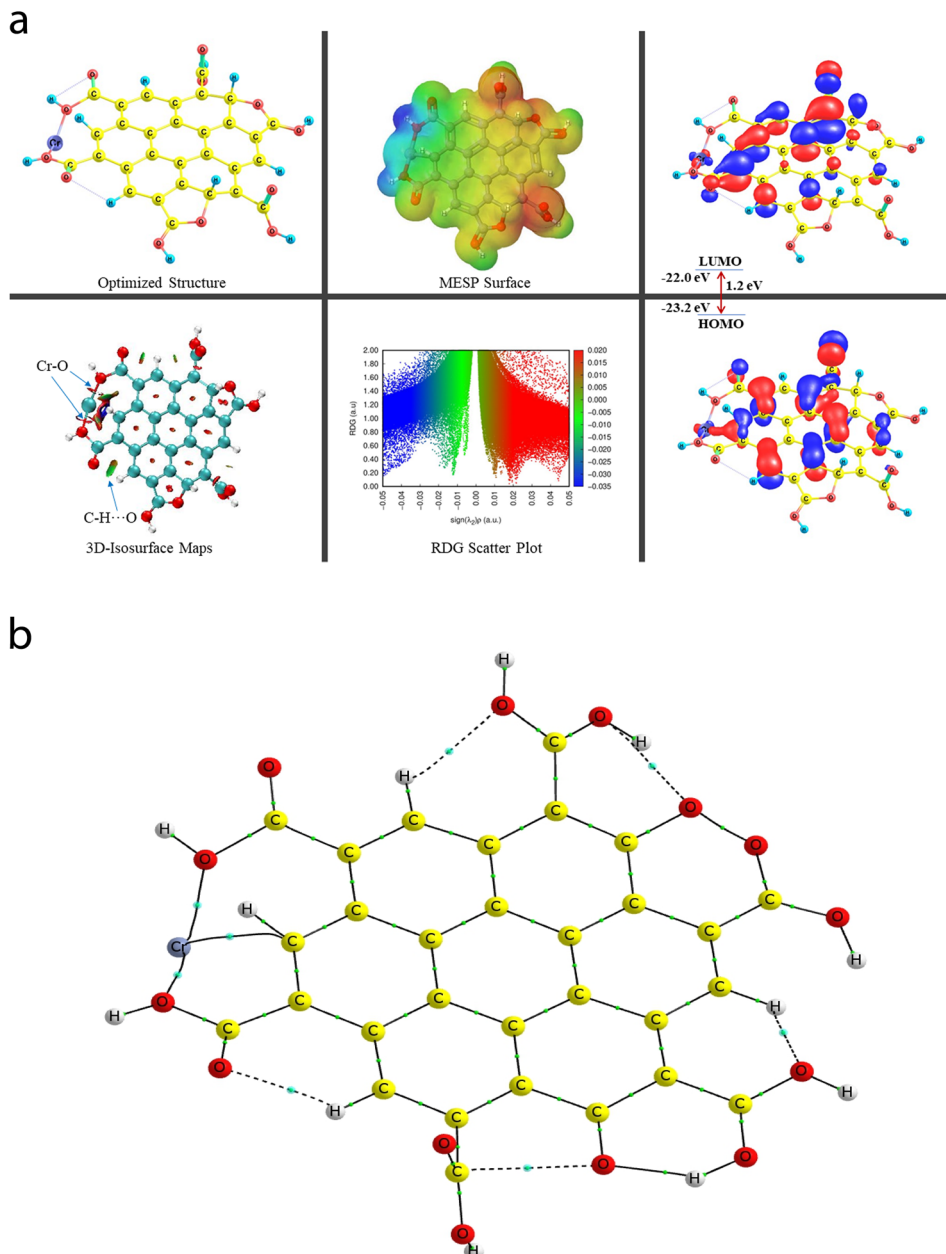


Fig. 8 (a) Optimized structure, MESP surface, HOMO–LUMO 3D maps, 3D-isosurface maps, and RDG scatter plot of Cr^{6+} interacting with the functionalized CNP to form the $\text{Cr}^{6+}@\text{CNPt}$ complex at the terminal of the CNP moiety. (b) QTAIM molecular graph of the Cr^{6+} ion interacting with the functionalized CNP to form the $\text{Cr}^{6+}@\text{CNPt}$ complex at the terminal of the CNP moiety.

and $\text{Cr}^{6+}@\text{CNPc}$ complexes, respectively. These findings suggest that the probed $\text{Cr}^{6+}@\text{CNPt}$ and $\text{Cr}^{6+}@\text{CNPc}$ complexes reported in this work are much more stable (more selective and higher sensitivity) than the $\text{Cr}^{6+}@\text{GSH}$ complex.

Interestingly, the following scenario was seen for the Cr^{6+} - and Fe^{3+} -associated CNP complexes (see Table 2), where it was observed that the calculated BE/IE values suggest that the complexation with both metal ions (Cr^{6+} and Fe^{3+}) located at the terminal position of the CNP unit is more favorable (*i.e.*, more selective and highly sensitive) ($\text{Cr}^{6+}@\text{CNPt}$: $-4476.7 \text{ kcal mol}^{-1}$ > $\text{Cr}^{6+}@\text{CNPc}$: $-4395.1 \text{ kcal mol}^{-1}$) compared to the complexation of the CNP moiety with the TM cations positioned in the

center. This can be attributed to the ion-dipole interaction between the metal cation and the COOH-functionalized CNP framework. A similar pattern can be viewed in the case of both $\text{Fe}^{3+}@\text{CNP}$ complexes ($\text{Fe}^{3+}@\text{CNPt}$: $-794.4 \text{ kcal mol}^{-1}$ > $\text{Fe}^{3+}@\text{CNPc}$: $-696.5 \text{ kcal mol}^{-1}$). According to Table 2, the BE results suggest that the TMs located at the terminal of the CNP moiety appear to show much more intense sensing features compared to the TMs located in the center of the CNP framework.

The E_{HOMO} and E_{LUMO} values of the $\text{Cr}^{6+}@\text{CNPc}$ complex were calculated to be -23 eV and -21.5 eV , respectively, whereas the E_{HOMO} (-23.2 eV) decreased and E_{LUMO} (-22 eV)

Table 4 The QTAIM parameters of the Cr^{6+} ion (in the center) interacting with the functionalized CNP, forming the $\text{Cr}^{6+}@\text{CNPc}$ complex

BP	BL (Å)	BPL (Å)	ρ (au)	$\nabla^2(\rho)$ (au)	V (au)	DI (A, B)
MNI (intermolecular)						
Cr53-C4	2.215	2.253	0.0572	+0.2194	-0.0739	0.374
Cr53-C9	2.298	2.337	0.0483	+0.1942	-0.0570	0.311
NCI (intramolecular)						
O56-H57···O31	1.467	1.490	0.0777	+0.2019	-0.0766	0.159
C23-H26···O43	2.077	2.101	0.0213	+0.0770	-0.0171	0.061
C12-H18···O30	2.161	2.332	0.0215	+0.0966	-0.0172	0.051
C19-H22···O42	2.195	2.402	0.0204	+0.0907	-0.0162	0.049
NBP (intramolecular)						
C45···O54	2.689	2.704	0.0154	+0.0573	-0.0109	0.042
C45···O54	2.872	2.704	0.0154	+0.0573	-0.0109	0.042

Table 5 The QTAIM parameters of the Cr^{6+} ion (at the terminal) interacting with the functionalized CNP, forming the $\text{Cr}^{6+}@\text{CNPt}$ complex

BP	BL (Å)	BPL (Å)	ρ (au)	$\nabla^2(\rho)$ (au)	V (au)	DI (A, B)
MNI						
Cr51-O37	1.979	2.020	0.0768	+0.4741	-0.1244	0.546
Cr51-O41	1.988	2.030	0.0775	+0.4408	-0.1225	0.513
Cr51-C18	2.268	2.400	0.0528	+0.2113	-0.0643	0.318
NCI						
O48-H57···O56	1.652	1.677	0.0515	+0.1676	-0.0448	0.106
C12-H53···O49	2.090	2.116	0.0207	+0.0789	-0.0168	0.058
C22-H52···O40	2.221	2.259	0.0156	+0.0587	-0.0116	0.045
C6-H17···O33	2.279	2.350	0.0149	+0.0596	-0.0110	0.037
NBP						
C43···O56	2.668	2.687	0.0156	+0.0587	-0.0114	0.040
O32···O54	2.743	2.853	0.0141	+0.0586	-0.0114	0.055
O28···O54	1.451	2.746454	0.275065	-0.006080	-0.417653	1.135357

increased for the $\text{Cr}^{6+}@\text{CNPt}$ complex. The E_{gap} value of the $\text{Cr}^{6+}@\text{CNPc}$ complex (1.5 eV) was found to be slightly larger than that of $\text{Cr}^{6+}@\text{CNPt}$ (1.2 eV), which demonstrates that the latter is more sensitive compared to the former, which is also confirmed by the BE values. The HOMO-LUMO 3D isosurface maps of the $\text{Cr}^{6+}@\text{CNPc}$ and $\text{Cr}^{6+}@\text{CNPt}$ complexes can be seen in Fig. 7a and 8a, respectively. In the case of the $\text{Cr}^{6+}@\text{CNPc}$ complex, its HOMOs appear to be primarily spread over the O atoms of the carbonyl group of the COOH functional groups, whereas the LUMOs seem to be distributed around the O atoms (out-of-plane), which depict an n to π^* transition (see the HOMO-LUMO plot in Fig. 7a). In the case of the $\text{Cr}^{6+}@\text{CNPt}$ complex (see Fig. 8a), the HOMOs are mainly located over the CC bond (in-plane) of the six-membered ring of the CNP framework, while the LUMOs are around the CC bond but in the out-of-plane phase, which describes the π to π^* transition. As shown by the 3D MESP surfaces of both chromium-related complexes, it can be seen that the positive regions are positioned over the H atoms of the COOH functional groups, whereas the negative regions are located over the -COO segments of the COOH functional groups in both complexes ($\text{Cr}^{6+}@\text{CNPc}$ in Fig. 7a and $\text{Cr}^{6+}@\text{CNPt}$ in Fig. 8a).

5.3.2 NCI-RDG isosurface plots and QTAIM analyses. The NCI-plot based 2D scatter plot and 3D isosurfaces of the $\text{Cr}^{6+}@\text{CNPc}$ complex are displayed in Fig. 7a. The 3D isosurface map of the $\text{Cr}^{6+}@\text{CNPc}$ complex validates the presence of two MNIs (see blue colour discs) between the chromium metal cation and a few C atoms of the central six-membered ring of the CNP skeleton, which is also confirmed by the QTAIM analysis (see two MNIs of Cr53-C4 and Cr53-C9), as shown in Fig. 7b. The MNIs were also confirmed by the 2D scatter plots, where the blue colour spikes range from -0.04 au to -0.045 au and indicate the existence of MNIs. The QTAIM parameters for two MNIs, i.e., Cr53-C4 (ρ : 0.0572 au, $\nabla^2(\rho)$: +0.2194 au, V : -0.0739 au, and DI: 0.374) and Cr53-C9 (ρ : 0.0483 au, $\nabla^2(\rho)$: +0.1942 au, V : -0.0570 au, and DI: 0.311), can be seen in Table 4, where the former is stronger than the latter. A total of six interactions [four NCIs (one O-H···O and three C-H···O) and two NBPs (C···O)] are involved in the $\text{Cr}^{6+}@\text{CNPc}$ complex. The bond distances for all four HBs range from 1.467 Å to 2.195 Å, where the respective BPLs are consistent, as expected. The associated strength parameters are almost reciprocal for the corresponding BLs and BPLs. The QTAIM parameters for the NCIs and NBPs are shown in Table 4.



The 2D scatter plot and 3D isosurface maps of the Cr^{6+} @CNPt complex are shown in Fig. 8a. It can be seen that three strong MNIs [two Cr–O MNIs: Cr51–O37 and Cr51–O41 and one Cr–C MNI: Cr51–C18] exist in the Cr^{6+} @CNPt complex, which play a major role in stabilizing it. The QTAIM molecular graph of the Cr^{6+} @CNPt complex (see Fig. 8b) and the associated parameters confirmed the presence of these three MNIs. It showed that two Cr–O MNIs (ρ : 0.0768 au and 0.0775; $\nabla^2(\rho)$: +0.4741 au and +0.4408 au, V : −0.1244 au and −0.1225 au, and DI: 0.546 and 0.513) in the Cr^{6+} @CNPt complex are stronger in comparison to the two Cr–C MNIs present in the Cr^{6+} @CNPt complex, while the Cr–C MNI of the former is weaker than that of the latter and *vice versa* for the other Cr–C MNI involved in the former. Additionally, the Cr^{6+} @CNPt complex consists of a total of seven intramolecular interactions (one O–H···O and three C–H···O HBs, as well as one C···O and two O···O NBPs), which are shown in the 3D isosurface maps and 2D scatter plot. The QTAIM topological parameters for all the NCIs and the NBPs can be seen in Table 5, where the BLs of the NCIs range from 1.652 Å to 2.279 Å. The opposite trend can be observed in the case of the QTAIM-based strength quantifying parameters against their corresponding BLs and BPLs.

The associated QTAIM topological parameters (ρ , $\nabla^2(\rho)$, and V) were calculated (0.0423 au, +0.196 au, and −0.0454 au) for the Fe57–C3 bond of the Fe^{3+} @CNPc complex, respectively. The QTAIM-based topological features for the various intramolecular NCIs such as six C–H···O and two NBPs (C···O) involved in the case of the Fe^{3+} @CNPc complex can also be viewed in Table S1 of the ESI.† Alternatively, the Fe^{3+} @CNPt complex demonstrates three MNIs (two Fe–O and one Fe–C), which are slightly stronger than that of the Fe^{3+} @CNPc complex. The (ρ , $\nabla^2(\rho)$, and V) values for two observed Fe–O intermolecular MNIs, Fe53–O43 and Fe53–O39, of the Fe^{3+} @CNPt complex were determined to be (0.0568 au, +0.3931 au, −0.088 au) and (0.0545 au, +0.399 au, −0.0892 au), while they are (0.0537 au, +0.1918 au, −0.0058 au) for the Fe53–C19 bond, respectively. Alternatively, the five intramolecular NCIs (C–H···O) and two NBPs (O···O) acquired from the QTAIM technique for the Fe^{3+} @CNPt complex are shown in Table S2 of the ESI.†

The QTAIM topological parameters (ρ , $\nabla^2(\rho)$, and V) for the Hg53–O43 bond were found to be (0.0163, +0.0491, and −0.0116) (in au), whereas for the Hg53–O39 bond, they were detected to be (0.0108, +0.0305, −0.0067) (in au), respectively, showing that the former is much stronger (approximate one and half times) than the latter. Hence, these MNIs are much weaker than the MNIs (one Fe–C in the Fe^{3+} @CNPc and Fe^{3+} @CNPt complexes, whereas two Fe–O MNIs in only the Fe^{3+} @CNPt complex). The QTAIM parameters for several other intramolecular NCIs (seven C–H···O) and NBPs (two O···O) taking part in the Hg^{2+} @CNP complex can be seen in Table S3 of the ESI.†

Conclusively considering the sensitivity and selectivity features in all five inspected complexes here in this study, it is noteworthy to mention that the BEs for the COOH-functionalized CNP associated with the TM cations located at the terminal of the framework follow the order of

(Cr^{6+} @CNPt: −4476.7 > Fe^{3+} @CNPt: −794.4 > Hg^{2+} @CNPt: −183.9) (kcal mol^{−1}). Importantly and interestingly, this trend is also consistent with the BSSE complexation energy (Cr^{6+} @CNPt: −4688.9 > Fe^{3+} @CNPt: −785.6 > Hg^{2+} @CNPt: −226.2) (kcal mol^{−1}). Table 2 demonstrates that Cr^{6+} @CNPt is the most stable complex (consisting of the largest BE value, *i.e.*, is the most selective and sensitive) among the five computationally examined TM-associated CNP complexes, which assists in portraying the excellent sensitivity and selectivity of the as-prepared CNP for Cr^{6+} . By observing the five analyzed BE findings, the Cr^{6+} @CNPt species is highly preferred in terms of highest selectivity and most sensitive behavior.

6. Conclusions

Herein, a novel, convenient, cost-effective, and green synthetic route for the preparation of CNPs from *Madhuca longifolia* flower using a hydrothermal approach was proposed in this study. The average diameter of CNPs was determined by TEM and found to be 12.0 nm. The as-synthesized CNPs displayed strong blue fluorescence with an emission maximum of 454 nm at 370 nm excitation, and the prepared CNPs also exhibited excitation-dependent emission properties. The XPS and FTIR studies suggested that the CNPs were functionalized carbonyl, hydroxy and carboxylic acid groups, and the interaction between the carboxylic acid with Cr^{6+} is responsible for the quenching of fluorescence intensity *via* the static quenching phenomenon. The fluorescence lifetime decay study also validated the static quenching mechanism given that no change was observed in the fluorescence decay profiles. The LOD was found to be 102 ppb, which is much better in comparison with most of the reported nanosensors. The sensing mechanism for the detection of Cr^{6+} was well supported by the theoretical study.

The experimentally acquired outcomes were strongly supported by the DFT approach. The theoretical findings suggested that the interaction between the functionalized CNP framework and the Cr^{6+} (located at the terminal) is the strongest among the five studied complexes and demonstrated the highest sensitivity. It should be noted that the bonding features were mainly directed by the MNIs (Cr–C, Fe–C, Fe–O, and Hg–O) in all the investigated complexes, where the Cr^{6+} -associated bonding feature played a dominant role in terms of sensitivity. Moreover, many other NCIs (O–H···O and C–H···O) and NBPs (C···O and O···O) taking part in all complexes stabilized the CNP skeleton itself. Given that the quantum chemical calculations provide a good understanding of the structural, stability, and electronic features of the association of three TM cations with the functionalized CNP, these model complexes can facilitate new insights into understanding the sensing features analysed in the field of environmental chemistry.

Author contributions

Methodology, experimental performing, and writing original manuscript (Tuhin Mandal); steady-state fluorescence study



(Ashish Kumar Ghosh); reviewing and editing (Shiv Rag Mishra); designing, performing, and analysis of the theoretical study along with the conceptualization and computational methodology (Sarvesh Kumar Pandey); conceptualization, methodology, supervision, analysis of experimental results, writing/editing original manuscript and funding acquisition (Vikram Singh).

Conflicts of interest

The authors declare that they have no conflict of interest.

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