A sensing mechanism for the detection of carbon nanotubes using selective photoluminescent probes based on ionic complexes with organic dyes

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The multifunctional properties of carbon nanotubes (CNTs) make them a powerful platform for unprecedented innovations in a variety of practical applications. As a result of the surging growth of nanotechnology, nanotubes present a potential problem as an environmental pollutant, and as such, an efficient method for their rapid detection must be established. Here, we propose a novel type of ionic sensor complex for detecting CNTs – an organic dye that responds sensitively and selectively to CNTs with a photoluminescent signal. The complexes are formed through Coulomb attractions between dye molecules with uncompensated charges and CNTs covered with an ionic surfactant in water. We demonstrate that the photoluminescent excitation of the dye can be transferred to the nanotubes, resulting in selective and strong amplification (up to a factor of 6) of the light emission from the excitonic levels of CNTs in the near-infrared spectral range, as experimentally observed via excitation-emission photoluminescence (PL) mapping. The chirality of the nanotubes and the type of ionic surfactant used to disperse the nanotubes both strongly affect the amplification; thus, the complexation provides sensing selectivity towards specific CNTs. Additionally, neither similar uncharged dyes nor CNTs covered with neutral surfactant form such complexes. As model organic molecules, we use a family of polymethine dyes with an easily tailorable molecular structure and, consequently, tunable absorbance and PL characteristics. This provides us with a versatile tool for the controllable photonic and electronic engineering of an efficient probe for CNT detection.

Keywords: carbon nanotubes; ionic surfactant; organic dye; photoluminescence sensor

INTRODUCTION

In addition to being one of the best and most readily available one-dimensional (1D) model systems, carbon nanotubes (CNTs) demonstrate remarkable potential for multifunctional applications in electronics, reinforced composite materials, chemical and biological sensing, scanning probe microscopy, and many other areas1–6. Chemically functionalized CNTs could be designed for various advanced applications, including light-harvesting devices, sensors, drug delivery, and energy storage systems4–10. Whereas some of the mentioned applications remain theoretical, others, such as composite frames and coatings and rechargeable batteries, are being realized in practice10. However, with the strong, continuous growth of nanotechnology and a total mass production of CNTs exceeding thousands of tons per year10, CNTs could represent a significant source of industrial pollution in the near future. This concern is heightened by the fragmentary knowledge available regarding their toxicity8,9,11,12. CNTs can escape into the environment during production, application, and recycling; thus, the ambient risk posed by CNTs must be carefully controlled. To this end, we must be well equipped for the rapid detection of CNTs to prevent toxic exposure to the human body and in case of technological catastrophes.

Among the known techniques for chemical sensing, one of the most promising is photoluminescence (PL) by virtue of its high sensitivity, efficiency, and simplicity13,14. There are several PL-sensing approaches, categorized on the basis of their responses in terms of absolute intensity, intensity ratio, or redistribution of the PL emission spectrum, change in lifetime, and alteration in anisotropic features. Furthermore, some of these approaches can be beneficially combined, thereby improving sensitivity, selectivity, dynamic range, or other sensor parameters.

The most efficient PL sensing can be achieved through enhancement of the emission in the presence of the compound to be detected13,14. The quantum yield of the PL emission of CNTs is low1–6, and the PL is quenched by both the bundling of the nanotubes15,16 and interactions...
with \(\pi\)-conjugated compounds and biomaterials\(^{17-20}\), for example, rhodamine, fullerenes, or DNA. The “brightening” of CNT PL has been pursued for various photonics applications through the encapsulation of organic materials inside the nanotubes\(^{21,22}\), the covalent incorporation of sp\(^3\) defects\(^{23}\), the embedding of luminescent local states\(^{24}\), admixture with an electrolyte\(^{25}\), and the non-covalent attachment of \(\pi\)-conjugated organic molecules\(^{26-29}\). The merits of non-covalent attachment of organic molecules include preserving the unique intrinsic features of the nanotubes while simultaneously endowing them with novel advantages associated with the organic conjugates\(^{30,31}\).

Among the various organic conjugated compounds available, polymethine dyes (PDs), which are organic molecules that contain polymethine chains of variable lengths and terminal functional groups, can be easily modified and tuned for particular opto-electronic tasks\(^{30}\). Specifically, the most important distinguishing feature of PDs and their donor-acceptor conjugates is their intense and relatively narrow absorption and fluorescence bands, which can be deliberately tuned in the visible and NIR range through modification of the polymethine chain length or through the attachment of specific terminal groups with donor-acceptor conjugates. Among the most important distinguishing features of PDs and their donor-acceptor conjugates is their intense and relatively narrow absorption and fluorescence bands, which can be deliberately tuned in the visible and NIR range through modification of the polymethine chain length or through the attachment of specific terminal groups with their own large \(\pi\)-electron systems. Furthermore, PDs behave as advanced photonic materials known as spectral sensitizers\(^{31}\), light harvesters\(^{32}\), and "laser" dyes with large optical nonlinearity\(^{33-35}\).

One of our objectives in this study was to pioneer an efficient method for the detection of CNTs through considerable amplification of CNT PL via their interaction with tailorable PDs. In water, CNTs become encompassed by ionic surfactant to form micelles\(^{36}\), thus interfering with the interactions between most studied PL probes and the CNTs themselves. Thus, we need to identify an efficient mechanism for sensing CNTs in aqueous media and appropriate probe materials that will help us to overcome the presence of surfactant around the CNTs and thus significantly enhance the sensitivity of the PL sensor.

**MATERIALS AND METHODS**

**Sample preparation**

Purified single-wall CNTs (CoMoCAT) were purchased from SWent, Inc. (SWent CG100, Lot # 000-0012) and used as a source material to prepare the CNT dispersions. In a typical experimental procedure, 1.2 mg of CNTs was dispersed in 20 mL of deionized water in the presence of a surfactant. The ionogenic surfactants sodium dodecylbenzene sulfonate (SDBS), sodium dodecylsulfate (SDS), sodium deoxycholate (SDOC), and sodium taurodeoxycholate (STDOC) and the neutral surfactants polyoxyethylene octyl phenyl ether (Triton X-100) and polyvinylpyrrolidone (PVP) were used to disperse the CNTs. We used the following amounts of surfactant in the above dispersions: 6.5 mg of SDBS, SDS, and SDOC; 5.0 mg of STDOC; 15.2 mg of Triton X-100; and 19.2 mg of PVP. The use of higher concentrations of the last two surfactants, Triton X-100 and PVP, was motivated by the need to increase the low concentrations of debundled CNTs in the supernatant dispersions. First, the above dispersions were subjected to ultrasonication using a NanoRuptor (Diagenode) processor for 1 h at 21 kHz and 250 W. Second, the dispersions were subjected to ultracentrifugation for 2 h 30 min at 17°C at 47 000 rpm for SDBS and 20 000 rpm for the other surfactants (Beckman Coulter Optima Max-XP, MLS 50 rotor) to remove the aggregate phase and obtain debundled supernatant solutions of CNTs.

A series of PDs was supplied by the Institute of Organic Chemistry (NASU, Ukraine). The following dye concentrations were studied: 0.0005, 0.001, 0.002, 0.004, and 0.008 mg mL\(^{-1}\). Mixtures with higher concentrations of PDs (>0.008 mg mL\(^{-1}\)) were not considered in the discussion because of the effect of PL emission re-absorbance. To prepare mixtures of CNTs and PDs, we used approximately 20% of the initial solutions of CNTs dispersed in SDBS and STDOC, approximately 33% of the initial solutions of CNTs dispersed in STDOC and Triton X-100, and approximately 80% of the initial solutions of CNTs dispersed in PVP and SDS. The different percentages of the initial CNT solutions were chosen to ensure that the PL signal was compatible with the sensitivity limits of the PL detector. The concentrations of specific CNTs in all mixtures and neat CNTs for the reference media used in the experiments were the same (20%, 33%, and 80%, respectively, for the abovementioned surfactants).

To study the dependence of the PL on the CNT concentration dispersed in SDBS, we tested a range of diluted concentrations (from 3% to 20% of the initial CNT concentration).

**Experimental setup**

The absorption spectra in the visible and NIR ranges were measured using a Lambda 1050 UV/VIS/NIR (Perkin Elmer) spectrometer. The fluorescence emission spectra at various excitation wavelengths were recorded using a Horiba NanoLog excitation–emission spectrophotometer equipped with a nitrogen-cooled InGaAs array detector to generate PL excitation-emission maps (PLE maps), with the X-axis representing the wavelength of the PL emission, \(\lambda_{EX}\), and the Y-axis representing the excitation wavelength, \(\lambda_{EM}\). Entrance/exit slits of 14 nm in width were used for both the excitation and emission monochromators in the NIR PL measurements. The PL experiments in the visible range were performed using a silicon detector and 2 nm entrance/exit slits for the monochromators. The time-resolved PL decays were measured using a LifeSpec II spectrophotometer (Edinburgh Instruments Ltd., UK) to perform time-correlated single-photon counting (TCSPC) under excitation by a picosecond pulsed diode laser at 405 nm.

Raman spectra were recorded using a Horiba–Jobin–Yvon LabRAM Raman microscope with a laser wavelength of 660 nm, operating at a power of approximately 14.5 mW and with a grating of 600 lines mm\(^{-1}\). A slit of 50 \(\mu\)m in width was used to improve the spatial resolution. The detector was a Synapse CCD detector. The spectra were collected by averaging 4 acquisitions, each of 30 s in duration. The Raman shift was calibrated using the Raleigh peak and the 520.7 cm\(^{-1}\) silicon line from an Si(100) reference sample. Samples were dropcast from solution onto Si(100) wafers and dried at room temperature prior to analysis. A minimum of three spectra were recorded from different areas of each sample to ensure statistical relevance.

**RESULTS AND DISCUSSION**

By testing a series of newly synthesized and well-known PDs for the rapid recognition of CNTs covered with ionic and non-ionic surfactants in water, we demonstrated enhancement of the PL from the studied semiconductor tubes functionalized by astraphloxin dye (Figure 1). The astraphloxin molecule consists of cyanine terminal groups connected to polymethylene chains, and this molecule has an uncompensated positive charge on one of its nitrogens (Figure 2a). We selected SDBS as a basic surfactant in our experiments because of its high efficiency in dispersing CNTs and thus yielding stable debundled nanotubes in water\(^{36}\).

In the mixtures of CNTs with dozens of the tested PDs (some of them are described in the Supplementary file), we observed slight quenching of the PL with no influence on the NIR emission of the CNTs. However, astraphloxin resulted in significant amplification of the PL emission from the CNTs, which we attribute to the formation of complexes consisting of CNTs, surfactant, and dye. The behaviour
observed in the absorption spectra of the studied complexes helped us to understand the nature of this complexation. The maximum of the absorption spectrum for a PD molecule ($\lambda_M$) is predominantly determined by the length of the $\pi$-conjugated polymethine chain. In our study, the absorption spectrum of an astraphloxin solution exhibited a maximum at $\lambda_M = 540$ nm (Figure 3a, curve 1) regardless of its concentration (Supplementary Figure S1). Because of the complexation of astraphloxin with CNTs, the absorption spectra for the mixtures of astraphloxin with CNTs contained red-shifted dye absorption peaks. The extent of the red-shift depended on the dye concentration (Figure 3a, Supplementary Figure S2). Essentially, the absorption peak of the astraphloxin monomers ($\lambda_M$) vanished in the CNT mixtures at low dye concentrations (Figure 3a, curve 2). The red-shifted absorption bands in the range of 560–600 nm were referenced to J-like aggregates of astraphloxin of various lengths. J-aggregation is a supramolecular self-organization of particular dyes in solution to form “staircase”-type units (“head-to-tail” aggregation). Note that neat solutions of the dye remained stable for weeks with no aggregate formation, whereas mixtures of astraphloxin with CNTs that were aged overnight (or longer) contained small, shiny crystals/particles (Figure 2b) as well as dye deposited on the walls of the sampling cells. An increase in the dye concentration resulted in an increase in the intensity ($\lambda_M$) of the peak associated with the dye monomers (Figure 3a, curves 3 and 4). Mixtures with high concentrations of astraphloxin yielded absorption spectra that could be fitted with a superposition of three components: one associated with the dye monomers, one associated with the interactions between the dye and the CNTs, and one associated with the CNTs themselves (Figure 3a, Supplementary Figure S2). Additionally, the PL excitation spectra (specifically, the dependence of the PL emission at a specific wavelength ($\lambda_{EM}$) on the excitation wavelength ($\lambda_{EX}$)) displayed a consistent red-shift of the peak (from 549 to 557 nm, curves 2–4 in Figure 3b) with increasing dye concentration in the mixtures. Thus, the explicit red-shift of the absorption and PL excitation peaks can be explained by the interaction of the PD molecules with the CNT micelles and the formation of J-like aggregates of astraphloxin in the vicinity of the nanotubes. As shown in Figure 3a, in the studied mixtures, we observed a threshold in the appearance of the dye monomers ($\lambda_M$) at concentrations in the range of 0.002–0.004 mg mL$^{-1}$, indicating saturation in the attachment of the dye molecules to the nanotubes. From this observation, we infer that astraphloxin may cover the surface of a CNT only in a single layer. A similar phenomenon of single-layer formation has been demonstrated in a system of CNTs non-covalently covered with porphyrin molecules.

The interaction of astraphloxin with CNTs is evidenced by the absorbance data in the range of transparency of the dyes (Supplementary Figure S3), where we analysed the excitonic peaks $E_{11}$ (900–1250 nm) and $E_{22}$ (550–750 nm) in the CNT absorption spectra for the above mixtures. The high-energy $E_{22}$ peak of the CNTs, with its maximum at 655 nm, does not change in position, whereas the low-energy $E_{11}$ peaks at 994 and 1024 nm are red-shifted to 999 and 1029 nm, respectively.

In the PL studies of the mixtures of PDs with CNTs in the range of intrinsic dye emission, we observed significant quenching (by a factor of 4) and red-shifting of the maxima (at approximately 11 nm) of the emission of the astraphloxin as a result of its admixture with CNTs.
for neat CNTs and mixtures of CNTs with various concentrations from the picosecond timescale to the nanosecond timescale. This fixation hinders the rapid oscillations of the astraphloxin molecule and thus delays the relaxation process of PL excitation-emission.

Therefore, the second lifetime component appears to be attributable to the fixation of the dye on the surfaces of the CNT–surfactant micelles. By fitting the time-resolved PL kinetics in the dye emission range, we obtained a PL lifetime of 30–40 ps for neat astraphloxin, whereas in the solutions with CNT complexes, this lifetime was supplemented by a second component amounting to 1.5 ns regardless of the astraphloxin concentration. A lifetime in the range of 1 ns has been observed for astraphloxin deposited on a substrate as a thin layer 37.

(Figure 3c and d). The observed PL quenching can be partially explained by the non-radiative transfer of the excitation energy from the dye to the CNTs, followed by further emission from the CNT excitonic levels. The magnitude of the PL quenching is much lower than that for CNTs covered with porphyrin in the absence of surfactant molecules (this quenching is by three orders of magnitude) 28. This finding indicates that in the studied complexes, the surfactant remains in the CNT and the dye, limiting the approach distance between them. By fitting the time-resolved PL kinetics in the dye emission range, we obtained a PL lifetime of 30–40 ps for neat astraphloxin, whereas in the solutions with CNT complexes, this lifetime was supplemented by a second component amounting to 1.5 ns regardless of the astraphloxin concentration. A lifetime in the range of 1 ns has been observed for astraphloxin deposited on a substrate as a thin layer 37.

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In the NIR range, we measured PL excitation-emission (PLE) maps for neat CNTs and mixtures of CNTs with various concentrations of astraphloxin (Figure 4, and see Supplementary Figure S4 for more PLE maps). The PLE maps for the neat CNTs (Figure 4a) with λ_{EX} at E_{22}, E_{33}, and E_{44} and λ_{EM} at E_{11} clearly reveal the presence of CNTs with chiralities of (6,5), (7,5), and (8,4) in the studied dispersions. The PLE maps of the mixtures indicate an increase in the PL intensity from tubes of all chiralities (E_{11}) in the dye excitation range, with a maximum at λ_{EM} = 555 nm (Figure 4b). The maximum of λ_{EX} does not coincide with λ_{EM}, the absorption maximum of the monomers. However, it does coincide with the absorption maximum for the interaction of the dye with the CNTs (554 nm at 0.0005 mg mL^{-1} in Figure 3a) and corresponds to E_{Dx} in Figure 4b. The resonant energy transfer (RET) from the astraphloxin–surfactant (donor) complex attached to the CNT (acceptor) is the most obvious cause of this effect and can be explained in terms of dipole–dipole interactions between the molecules. Moreover, the dye monomers present in the mixtures at high concentration do not transfer energy to the CNTs, as seen from the absence of PL peaks at λ_{EX} = λ_{M} = 540 nm (Figure 4b). This means that short-range energy transfer of the Dexter type is more likely than a Förster mechanism for RET, in which energy can be transferred over distances of up to 90 Å 39.

To more explicitly demonstrate the amplification of CNT emission, we extracted the PL emission spectra at λ_{EX} = 555 nm from the PLE...
maps for presentation in Figure 5a. The maximum PL amplification was achieved at an astraphloxin concentration of 0.004 mg mL$^{-1}$. Moreover, the PL spectra indicate the different effects of the dye on CNTs of each chirality, revealing selectivity in the sensing of CNTs of various diameters (Figure 5b). In Figure 5b, we selected the data from Figure 5a corresponding to a dye concentration of 0.004 mg mL$^{-1}$ and to neat CNTs and plotted a histogram of the PL amplification ranges for CNTs of various chiralities — (6,5), (7,5), and (8,4). The highest (up to a factor of 6) PL amplification was achieved for the (7,5) chirality. The varying selectivity for CNT chirality can be explained in terms of factors related to electronic structure and geometry. The latter are related to the diameter of the nanotubes or, more precisely, the surface area of the CNTs to which the dye molecules can become attached. Considering this approach only, a larger surface area implies that a greater number of dye molecules can complex with the CNTs. However, the CNTs of the (8,4) chirality, those with the largest diameter, did not exhibit the highest PL amplification. This finding indicates that the electronic structure parameters of the CNTs and the organic dye must simultaneously influence the PL amplification. Therefore, the structural matching of the π-electron systems of specific chiral CNTs and organic molecules is of considerable importance in determining the selectivity of non-covalent interactions.

The different dependences of the PL in the visible range (emission from the dye) and the NIR range (emission from the CNTs) on the astraphloxin concentration in the mixture are depicted in Figure 5c. The NIR emission tends to saturate, whereas the PL in the visible range grows until the phenomenon of re-absorbance of the emission begins to hinder this effect. The saturation of the NIR PL suggests that the number of astraphloxin molecules that are able to attach to a CNT is limited. An increase in the dye concentration results in the presence of free dye molecules that do not interact with the CNTs and emit light at the maximum of the monomers (Figure 3d).

We note that the intensity of the intrinsic PL of the CNTs (peak at $\lambda_{EX} = 645$ nm and $\lambda_{EM} = 1028$ nm in Figures 3b and 4) changes very little and can be regarded as constant with the variation in the dye concentration. Thus, the interactions of the CNTs with the astraphloxin...
sustain their intrinsic PL properties while endowing them with novel advantages. Furthermore, the preservation of the intrinsic features of the CNTs observed in our study is a common phenomenon in the non-covalent functionalization of CNTs by organic molecules.

To explore the possible application of the studied complexes in sensors, we analysed the dependence of the PL amplification on the concentrations of nanotubes of the three basic chiralities (Figure 5d, and see Supplementary Figure S5 for a description of the estimation of the CNT concentrations). Regarding this concentration dependence, in addition to establishing the shape of the calibration curves for CNT sensing, we evaluated the detection limit, which was found to lie in the range of (1–4) ng mL$^{-1}$ for chiralities of (6,5), (7,5) and (8,4) (see the Supplementary file for details). This detection limit is comparable to that previously obtained by another research group.

Modification of the molecular structure of astraphloxin can help us to understand the specific nature of the CNT–PD interaction and, potentially, to improve the PL amplification. A version of the astraphloxin molecule modified with a long alkyl chain, Cy3, exhibited behaviour similar to that of astraphloxin, but the PL amplification effect was lower, yielding a maximum amplification of a factor of 4 at a Cy3 concentration of 0.002 mg mL$^{-1}$ (Supplementary Figures S4, S6, and S7). This means that more astraphloxin molecules could contribute to the RET because of the possibility of better regularity in their attachment to the CNTs.

Another investigated approach was to test the PL sensing properties of astraphloxin molecules substituted with different anionic groups. The specific purpose of the test was to exploit PDs with the same molecular backbone modified by an anionic-linking group and thus to neutralize the positive charge of the astraphloxin (i.e., the formation of betaine systems). In the absorption and PL spectra, the mixtures of the betaine dyes, namely neutral forms of modified astraphloxin or Cy3, with CNTs resulted in the simple superposition of the two compounds without any new features or PL amplification (see the molecular structures of some of these dyes in Supplementary Figure S8). Thus, we demonstrated that astraphloxin analogues neutralized with anionic groups do not transfer energy to CNTs dispersed with SDBS and that
there is little or no interaction between such a neutral dye and a charged CNT micelle.

The next logical step was to consider the effects of various surfactants used to disperse the CNTs on the complexation and, perhaps, PL amplification. We tested CNT micelles dispersed with the non-ionic surfactants Triton X-100 and PVP and compared their interactions with astraphloxin and its analogues. In the case of the mixtures of the nanotubes with astraphloxin, a superposition of the spectra of the mixed components was observed and the PL signal of the CNTs decreased in the presence of the dye (Supplementary Figure S9, Supplementary Tables S1–S3). Furthermore, no RET to the nanotubes was observed.

It should be emphasized that the astraphloxin analogues neutralized with anionic groups did not transfer energy to the CNTs regardless of the anionic or neutral nature of the surfactants used. These findings help us to understand the nature of the interaction between the molecules and reveal the mechanism of the complexation between astraphloxin and CNTs. In this manner, the Coulomb attraction of positively charged PDs to anionic groups of the surfactant that forms the CNT micelles is established as a key requirement for the formation of energy transfer complexes and the aggregation of the dyes on CNT micelles. Thus, the utilization of the interactions between charged PDs and surfactant-CNT micelles can be regarded as a novel approach to creating efficient PL sensors for CNTs in aqueous environments.

A series of CNT dispersions with various ionic surfactants (SDBS, SDS, SDOC, and STDOC) were tested for complexation with astraphloxin. A comparison of the absorption spectrum of a neat solution of astraphloxin (with constant $\lambda_{\text{EX}}$) with those of its mixtures with different CNT micelles revealed the emergence of new absorption features of the mixtures in the range of 550–600 nm (Figure 6a). A new explicit peak at 576 nm appeared in the absorption spectra of the mixtures with CNTs dispersed with SDOC and STDOC, whereas the spectra of the mixtures with CNTs dispersed with SDBS and SDS exhibited a shoulder in that range. The peak at 576 nm is attributed to the longer shift (up to 36 nm) in absorption that occurs when the dye interacts with CNTs covered with deoxycholate surfactants (SDOC and STDOC). As a result, the absorption peaks of the astraphloxin monomers and astraphloxin-CNT complexes do not overlap in the case of deoxycholate-treated CNTs. This longer shift can be explained in terms of a better arrangement of the astraphloxin molecules in the J-like aggregation because of the influence of the π-electron systems in the SDOC and STDOC surfactants. For excitation in the range of the new absorption peak (550–600 nm), we observed enhancement of the PL from the CNT levels ($E_{\text{FL}}$ in Figure 4b) for all mixtures containing anionic surfactants (Supplementary Figure S9, Supplementary Table S3). The excitation maximum for the PL amplification varied from $\lambda_{\text{EX}} = 555$ nm for SDBS to $\lambda_{\text{EX}} = 565$ nm for SDS and to $\lambda_{\text{EX}} = 570$ nm for SDOC and STDOC. CNTs with the (7,5) chirality displayed the maximum PL amplification for all anionic surfactants. In the histogram presented in Figure 6b, we summarize the data for the PL enhancement of (7,5) CNTs, including the reduction of the PL intensity for CNTs dispersed with neutral surfactants due to the bundling process mentioned above. The maximum PL enhancement was observed for mixtures of astraphloxin with CNTs dispersed with SDBS (see Supplementary Tables S1–S3 for more data). Thus, this study demonstrated the different effects of the dyes on CNTs of each chirality, revealing selectivity in the sensing of CNTs of various diameters. The effect is similar to the selectivity observed when sorting CNTs using SDOC and STDOC$^{43}$. Additionally, we observed a striking selectivity in PL amplification among CNT-dispersing surfactants.

To confirm that the changes observed in the above systems were caused by the interaction of the PDs with the CNTs, we studied a two-component system of mixtures consisting only of the dye (astraphloxin or its analogues) and the surfactant. No PL features appeared in the NIR range of the spectra. Thus, the fundamental contribution to the amplification in the NIR range stems from the interaction of the surfactant-covered CNTs with the dye molecules.

In addition, we performed an alternative characterization of the studied systems via Raman spectroscopy to gain a better understanding of the PL amplification mechanism and the role of ionic and non-ionic surfactants in the astraphloxin–CNT interaction. The most important feature of the acquired Raman spectra was an enhancement in the dye bands (relative to the vibrational modes of the CNTs themselves) at 1122, 1238, 1272, 1373, 1405, and 1473 cm$^{-1}$, which was assigned to the cyanine chromophore and polymethine chain modes$^{44}$, in the mixture of CNTs with ionic SDBS compared with the dye-containing mixture of CNTs with non-ionic Triton X-100 (Figure 7, and see Supplementary Figure S10 for an extension of the

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**Figure 6** Absorption spectra of astraphloxin with CNTs dispersed with various surfactants. (a) Absorption spectra for water solutions of neat astraphloxin (1) and mixtures of astraphloxin with CNTs dispersed with various surfactants: (2) SDBS, (3) SDS, (4) SDOC, and (5) STDOC. All spectra are normalized to the maximum at approximately 540 nm. (b) Histogram of the PL amplification ($\lambda_{\text{EX}} = 555–570$ nm) in the NIR range for astraphloxin with (7,5) CNTs dispersed with various surfactants – SDBS, SDS, SDOC, STDOC, Triton X-100, and PVP. The astraphloxin concentration in all solutions was 0.004 mg mL$^{-1}$. 

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the lowest electron transitions in CNTs (E_11) and in CNT–PD complexes. Strictly speaking, the features of the frontier levels directly affect the enhancement of the Raman bands to aggregation of the dye molecules, as demonstrated by Akins et al. (for example^{27}) and as discussed above for ionic complexes.

RET from the dyes aggregated with the CNTs is the most obvious explanation for the PL amplification effect. Thus far, because CNTs and PDs are \( \pi \)-electron systems, the most feasible form of the interaction between them is an aromatic stacking interaction, as previously discussed in the scientific literature^{27,28,46,47}. By means of preliminary quantum-chemical modelling (using the ZINDO/S method), we showed that the interaction between a CNT and a PD molecule separated by a surfactant molecule should be decreased but still present. Strictly speaking, the features of the frontier levels directly affect the lowest electron transitions in CNTs (E_11) and in CNT–PD complexes. Our model shows that as a result of the complexation of a CNT with astraphloxin, the highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO) are located on the CNT and the astraphloxin, respectively. Thus, in the CNT–PD complex, a new frontier LUMO emerges that is not present in the pure CNT. In this manner, the features of the frontier orbitals determine the transition between the LUMO on the astraphloxin and the HOMO on the CNT (E_11 in Figure 4b) in the studied complexes could be responsible for the strong PL amplification observed.

CONCLUSIONS

We describe a new photonic nanosystem consisting of PDs attached to CNT-surfactant micelles in water, which results in significant amplification of the CNT PL emission. The Coulomb attraction between the dye and surfactant molecules covering the CNTs is key to the emergence of novel photonic complexes. We observe that the creation of such complexes occurs through the Coulomb coupling of the positively charged astraphloxin molecules with the anionic surfactants covering the CNTs. The astraphloxin attached to a micelle interacts with the CNT via weak \( \pi - \pi \) complexation, resulting in quenching of the dye emission and enhancement of the CNT PL. PL excitation-emission mapping in the spectral ranges of the dye and CNTs allowed us to experimentally demonstrate that only PD molecules associated with nanotube-surfactant micelles contribute to the PL amplification of the CNTs. Moreover, we observed firm evidence of the J-like aggregation of astraphloxin on anionic CNT micelles, supported by absorbance, PL, and Raman spectroscopy measurements. Reference systems with neutral surfactants or modified astraphloxins with negatively charged sulfonated groups were unable to form such complexes with CNTs. We also report the physicochemical model of the complexation process, which is fully supported by a complementary study of the reference systems.

In this paper, we propose an efficient molecular-engineering approach to the detection of CNTs and establish a mechanism for the selective PL sensing of CNTs by means of ionic complexes. It should be emphasized that the limit of CNT detection estimated based on the dependence of the PL amplification on the concentrations of CNTs of various chiralities is in the range of ng mL^{-1}. The most prominent feature of PDs is their vast variety of functional versatility, including polymethine chain elongation and the possibility of applying various anions/cations to neutralize the charged dyes. We can exploit the easy tailoring of the molecular structures of PDs, which offers excellent prospects for the optimization of the chemical and physical properties of complexes of CNTs and organic dyes with tunable optical properties and thus demonstrates great potential for application in chemical sensing and other photonic applications.

The developed macromolecular CNT–PD complexes provide an innovative means of functionalizing carbon nanomaterials, particularly with the purpose of strong enhancement of PL emission, thereby extending the multifunctional applications of these materials in electronics, ultra-fast photonics, reinforced composite materials, chemical and biological sensing, scanning probe microscopy, drug delivery, energy storage, and many other areas.

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