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Abstract

Energy transfer in non-covalently bound porphyrin / carbon nanotube compounds is investigated at the single nano-compound scale. Excitation spectroscopy of the luminescence of the nanotube shows two resonances arising from intrinsic excitation of the nanotube and from energy transfer from the porphyrin. Polarization diagrams show that both resonances are highly anisotropic with a preferred direction along the tube axis. The energy transfer is thus strongly anisotropic despite the almost isotropic absorption of porphyrins. We account for this

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unexpected result by local field effects induced by the large optical polarizability of nanotubes. We show that the local field correction extends over several nanometers outside the nanotubes and drives the overall optical response of the functionalized nanotubes.
Single-Wall Carbon Nanotubes (SWNTs) have attracted much attention in the last decades due to promising properties for future applications. However, it appeared rapidly that their performances can be highly enhanced by means of functionalization. This latter allows to combine the exceptional properties of nanotubes with those of various molecules. In particular, in view of light harvesting or opto-electronics applications, the combination of a photo-active molecule (chromophore) with nanotubes is particularly attractive. Energy transfer, charge transfer, luminescence quenching are some of the possible effects that can result from the combination of chromophores with nanotubes and could possibly lead to new opportunities in organic photovoltaics or bio labeling. In all cases, the goal is to exploit the large optical absorption of the chromophore to funnel the light energy into the device. Therefore, it is essential to know how the optical response of the chromophore is affected by the presence of the nanotube and how efficient is the coupling between the chromophore and the nanotube.

Here, we focus on non covalently bound porphyrin / carbon nanotube compounds that display efficient energy transfer: absorption in the porphyrin molecules results in enhanced near infrared (NIR) emission from the nanotube. We performed an all-optical investigation of the coupling between the chromophore and the nanotube at the single nano-compound scale by means of micro photoluminescence excitation spectroscopy and polarization spectroscopy. Each (of a few tens of specimens) nano-compound shows two resonances arising from intrinsic excitation of the nanotube and from energy transfer from the porphyrin. The weak temperature dependence of the transfer efficiency (down to 10 K) is compatible with a purely electronic process. Moreover, polarization spectroscopy allowed us to reveal original antenna effects that lead to a peculiar coupling between light and the nano-compounds. We show that the local field effects due to the large optical polarizability of the nanotubes extend over several nanometers outside of the nanotubes and lead to a strongly anisotropic energy transfer resonance despite the almost isotropic absorption of the chromophore. A simple model based on the electromagnetic response of a dielectric cylinder allows to account quantitatively for the excitation diagrams and to gain more insight into the local field map surrounding the nanotube.
Free base tetraphenyl porphyrin molecules (H$_2$TPP, see ??(b) for the molecule structure) are non-covalently attached (Π-stacking) to the wall of carbon nanotubes (CoMoCat produced by Southwest Nanotechnologies SG65) by means of the micelle swelling method that we developed to achieve high yield functionalization in water. In solution, the average coverage of the nanotube wall by porphyrin molecules is of the order of 1 (one layer of porphyrins stacked on the nanotube wall with an almost 100% surface coverage). In such systems, we showed by absorption and luminescence spectroscopies that an efficient energy transfer occurs, that leads to near infrared (NIR) luminescence of the nanotubes upon excitation of the Soret band of the porphyrin (440 nm). Such an energy transfer has characteristic signatures in photoluminescence excitation (PLE) measurements. In this paper, we extended this technique to single nano-compound measurements, which allowed us to reveal new anisotropic effects in the energy transfer.

Figure 1: (color online) Scheme of (a) a Single Wall Carbon Nanotube showing its axis along which preferential light absorption and emission occur, (b) the H$_2$TPP molecule with its x and y axis along which the optical oscillator strengths are almost identical. In the visible, the absorption is negligible for an incident polarization perpendicular to the plane of the molecule.

(a) shows the photoluminescence (PL) map of a 1.5 x 1.5 $\mu m^2$ area of the sample where the emission of a single nano-compound is clearly visible. In ??(b) we show typical PL spectra of single nano-compounds that shrink into a single narrow line (3 meV) at low temperature, clearly showing the presence of single nanotubes. The excitation spectrum of such nano-compounds (see next section) allows to assign their chiral family following the scheme introduced in Ref. We need however to take into account the global redshift of the transitions induced by the interaction with the H$_2$TPP molecules, as discussed in Ref.
Figure 2: (color online) (a) Spatial map of the photoluminescence signal of a single compound based on a (6,4) nanotube at room temperature. Detection wavelength: 912 nm, excitation wavelength 633 nm, incident power: 100 W/cm². Color scale is in counts/s. (b) Normalized PL spectra of single (6,4) nanotubes at room temperature and low temperature. The broad (32 meV) PL at 300 K shrinks into one single narrow (4 meV) line at 10 K, which shows that the signal arises from a single nanotube.

**Energy transfer**

In order to investigate a possible energy transfer in individualized compounds, we performed excitation photoluminescence spectroscopy (PLE) on single nanocompounds by recording the NIR PL intensity of the nanotubes while tuning the excitation wavelength. The NIR PL spectrum of each nano-compound consists in one narrow line whatever the excitation wavelength (see ??b-e). The PLE spectrum is the integrated PL line intensity as a function of the excitation wavelength. Two typical results are shown in ??b-e. The PLE spectra show two resonances for each compound. One of them occurs at a wavelength that depends on the nanotube under investigation, whereas the other always appears at 440 nm. We assign the former to a direct excitation of the nanotube $S_{22}$ excitonic transition, whereas the latter corresponds to a resonant excitation of the $\text{H}_2\text{TPP}$ molecules Soret band followed by energy transfer to the nanotube and NIR luminescence at the nanotube $S_{11}$ excitonic resonance.

We first discuss the new insight given by such PLE measurements at the single nanocompound scale on the intrinsic properties of the nanotubes. The $S_{22}$ resonances widths are of the order of 60 meV, even at low temperature, which is very similar to the width obtained in ensemble measurements. This is in strong contrast with the lower $S_{11}$ transition that shows a linewidth of the order of 45 meV in ensembles but shrinks down to a few meV at the single nanotube scale at low temperature. This means that inhomogeneous broadening is dominant for the lower transition but almost negligible for upper transitions. This is consistent with the reduced lifetime and dephasing
time reported for these upper levels. We note that the functionalization of the nanotubes by the H$_2$TPP molecules does not induce any sizable broadening of the $S_{11}$ transition which remains of the order of a few meV at 10 K (see (b, c)). This confirms that non-covalent functionalization hardly perturbs the electronic system of the nanotubes and preserves the phase coherence which is known to be very sensitive to environment effects.

![Figure 3](PLE-54-65_v8-eps-converted-to.pdf)

Figure 3: (color online) (a) Integrated photoluminescence intensity as a function of the excitation wavelength for two single compounds based on a (5,4) nanotube (blue line, detection at 864 nm) and a (6,5) nanotube (green line, detection at 1002 nm). The gray line shows the spectrum of a bare (6,5) nanotube. (c) and (e) PL spectra of these nanocompounds excited on their $S_{22}$ transition. (b) and (d) PL spectra of the same nanocompounds excited on the energy transfer resonance at 440 nm. All spectra were divided by the excitation power and recorded at 10 K.

The second resonance showing up at 440 nm in all individual compounds (and absent in bare nanotubes (see ??)) corresponds to the observation of energy transfer at the single compound scale. In (a) we show that a clear energy transfer resonance occurs for both (5,4) and (6,5) nanotubes. Other results (not shown here) suggest that this energy transfer occurs whatever the chiral species, at least for the small diameter nanotubes (up to a diameter of 1 nm) investigated in this study. An important point here is that the energy transfer yield, that is closely related to the ratio between the intrinsic resonance and the energy transfer resonance, turns out to be constant (within a 20% error bar) when decreasing the temperature down to 10 K. This is consistent with previous time-resolved studies that showed an ultrafast process (of the order of 100 fs) in agreement with a pure electronic process (Dexter type mechanism) involving the overlap of the $\pi$ orbitals of the chromophore and the nanotube and leading to a very efficient energy transfer yield.
Polarization spectroscopy

SWNTs are well known for their highly anisotropic optical properties with light absorption and emission occurring essentially with a polarization parallel to the tube axis. In contrast, H$_2$TPP molecules absorb and emit light with an almost identical probability for polarizations along their x and y axis? (??(b)), whereas no transition is expected in the visible range for light polarized along the out-of-plane (z) axis. It is therefore interesting to check whether this in-plane isotropic absorption of H$_2$TPP can be used to modify the effective absorption properties of functionalized nanotubes and enhance their cross-polarized absorption through energy transfer.

Polarization spectroscopy is the perfect tool to address this issue. Although such a spectroscopy can be performed in ensembles of randomly oriented objects,? it still suffers from ad-hoc instrumental corrections. In contrast, single object spectroscopy allows us to perform a thorough analysis of polarization effects in the optical properties of functionalized nanotubes and to investigate the role of the polarization of light in energy transfer processes where the donor has an isotropic absorption and the acceptor is highly anisotropic.

We first checked the intrinsic properties of the nanotubes by analyzing the polarization diagram of the emitted light ($S_{11}$ transition) and the absorption polarization diagram of the $S_{22}$ transition by means of PLE measurements (??). We fit the data according to :

$$I(\theta) = A \cos^2(\theta - \theta_0) + B \sin^2(\theta - \theta_0)$$

where $\theta_0$ stands for the nanotube axis direction in the sample plane and $\theta$ is the direction of the polarization of light (see inset of ??). The results shown in ?? are consistent with the literature: the dipole of the nanotubes is essentially along the tube axis.?? We note that the extinction ratio, defined as $B/A$ is stronger in emission ($\sim 0.05$) than in absorption ($\sim 0.15$). This may be related to residual absorption from cross-polarized states or to the fact that the nanotube is not straight. In this latter case, the absorption has a perpendicular component, whereas emission can remain highly polarized since it arises from localized excitons.?
Figure 4: (color online) Polarization diagram of a single NT/H$_2$TPP nano-compound based on a (6,4) nanotube. (a) Excitation diagram at 633 nm on the S$_{22}$ transition. (b) Emission diagram (S$_{11}$ transition at 900 nm). The blue solid line is a fit to the data according to eq. (??). The axis of both diagrams are identical within the error bar.

The polarization diagram of the energy transfer resonance is shown in ?? (a) for a (6,5) based compound. Even if the absorption of light is expected to be quite isotropic for the H$_2$TPP molecules, the energy transfer resonance is clearly anisotropic with a preferential axis identical to the one of the S$_{22}$ resonance (?? (b)), that is to the tube axis. Furthermore, the extinction ratio is of the order of 0.12 which is comparable to the one observed for the S$_{22}$ transition.

Figure 5: (color online) Polarization diagram of a single compound based on a (6,5) nanotube (same as in ??). (a) Excitation diagram on the Soret transition at 440 nm (energy transfer). The red solid line is a fit to the data according to the local field model (see text) with $\varepsilon = 7$ and $\varepsilon_m = 2$. (b) Excitation diagram on the S$_{22}$ transition at 580 nm. The blue solid line is a fit to the data according to eq. (??).

It is well known that the so-called crossed optical transitions (S$_{12}$ for instance) that are allowed for an incident electric field perpendicular to the tube axis, are strongly suppressed in bare carbon nanotubes because of screening effects. This can be understood as a consequence of the depolarizing field induced in the nanotube in response to an incident field, that causes a strong cancellation of the latter inside the nanotube. Similar local field effects occur at the surface of the nanotube and in its close vicinity. Therefore, we base our interpretation of the induced anisotropy in the energy transfer resonance on local field effects due to the large polarizability of the nanotube. This effect can be quantitatively accounted for by modeling the response of a dielectric cylinder.
permittivity $\varepsilon$ of diameter $d$ in a dielectric environment (dielectric permittivity $\varepsilon_m$) to an incoming field $\vec{E}_0$ and writing down the boundary conditions at the nanotube interface in the electrostatic approximation (valid for $d \ll \lambda$)\(^{(??)}\).

Figure 6: Response of a dielectric cylinder to an external electric field. Inset: Incoming field orientation with respect to the tube axis. (a) Parallel component. (b) Perpendicular component.

For an incoming field $\vec{E}_0$ in the direction $\theta$ with respect to the tube axis ($Oz$) and in the limit of an infinite cylinder, the effective electric field at a point of cylindrical coordinates $M(r, \Phi, z)$ in the outer vicinity of the nanotube reads\(^{(??)}\):

$$E^\parallel_z = E_0 \cos \theta$$

for the field component parallel to the tube axis, and:

$$E^\perp_r = E_0 \sin \theta \left[ 1 - \frac{d^2 \varepsilon_m - \varepsilon}{4r^2 \varepsilon_m + \varepsilon} \right] \cos \Phi$$
$$E^\perp_\Phi = -E_0 \sin \theta \left[ 1 + \frac{d^2 \varepsilon_m - \varepsilon}{4r^2 \varepsilon_m + \varepsilon} \right] \sin \Phi$$

for the field components perpendicular to the tube axis.

An accurate value of $\varepsilon$ of the nanotubes in the optical range is not available, but realistic values can be taken from the literature and turn out to be of the order of 5 to 30 for non resonant and resonant conditions respectively\(^{(??)}\). Therefore, the electric field can be strongly suppressed in a perpendicular geometry, whereas neither screening nor amplification is expected in the longitudinal geometry. The results of this simple model accounting for local field effects are summarized in \(?\). In the close vicinity of the tube ($r - d \ll d$), the field remains almost radial except for $\Phi \simeq \pm \pi/2$ where it becomes tangential with an amplitude reduced by a factor $(\varepsilon + \varepsilon_m)/2\varepsilon_m$. Bearing in mind that porphyrin absorption is forbidden for an electric field perpendicular to its plane, we can readily
conclude that the absorption on the Soret band will be strongly suppressed in the perpendicular geometry (?? (b)). In fact, although the local field amplitude can reach up to twice the amplitude of the incoming field, its direction remains essentially radial, that is perpendicular to the average plane of the porphyrin molecule. On the other hand, for \( \Phi \sim \pm \pi/2 \), the field becomes parallel to the porphyrin plane, but its amplitude is strongly reduced. In total, the absorption of the porphyrin is strongly suppressed whatever its location on the tube surface. In contrast, its absorption is optimal for an incoming field parallel to the tube axis (??(a) and ??(a)). As a consequence, the energy transfer resonance shows a strong anisotropy along the tube axis, whereas each individual chromophore has no preferential axis.

![fieldmap-vect-eps-converted-to.pdf](fieldmap-vect-eps-converted-to.pdf)

Figure 7: (color online) Local electric field amplitude in the vicinity of a nanotube for an incoming field parallel to the tube axis (a) or perpendicular to the tube axis (b) calculated from the model presented in the body of the paper. In the latter case, the field direction at the interface is essentially radial except near the poles where it becomes orthoradial while its amplitudes decreases. The distances are in \( \text{nm} \) and the field amplitude is in units of the incident field. The porphyrin molecules are figured at a realistic scale on the surface of the tube.

The first piece of information that arises from these experimental data, is a clear confirmation of the local structural properties of the compound. Actually, this polarization effect can only occur if the \( \text{H}_2\text{TPP} \) molecules are properly attached with their plane stuck onto the nanotube wall. This would not be the case if the molecules were attached in an amorphous way or as small aggregates. Therefore, these polarization measurements first give an important insight into the molecular arrangement in the compounds, that was only predicted from energetic arguments before.

This anisotropy effect does not depend on the particular location of the porphyrin molecules on the nanotube wall (??). Actually, for a 500 nm long nanotube, several hundreds of porphyrin molecules are randomly attached on the wall leading to a strong averaging even at the scale of a single nanotube. Therefore, the right parameter for estimating quantitatively the anisotropy of the system is the average value \( \langle |E_\perp / E_0|^2 \rangle_\Phi \) over all orientations \( E_\perp \) gives no contribution to
the absorption since the absorption of H$_2$TPP is negligible for a field perpendicular to its plane). Actually, $<|E_{\Phi}/E_0|^2>_{\Phi}$ is directly comparable to the anisotropy coefficient $B/A$ of the phenomenological fit (Eq. (21)). The best fit to the data according to Eq. (21) is obtained for $\epsilon = 7$, with $\epsilon_m = 2$ (see (23)). These values seem reasonable: the former corresponds to the value predicted in the case of a non resonant frequency in carbon nanotubes$^7$ and the latter corresponds to a typical value for a transparent material such as the environment of the nanotubes.

It is enlightening to use the ratio $B/A$ to single out the role of the local field in contrast to a pure geometrical effect. In the latter case, $B/A = <\sin^2\Phi>$ turns out to be 1/2, whereas the full local field model leads to $B/A \sim 0.1$ in much better agreement with the experimental value (of the order of 0.1 as well). This shows that local field effects due to the large polarizability of the nanotube is actually the main reason for the overall optical anisotropy of the compounds.

This study shows that local field corrections (or antenna effects) are essential in understanding the optical properties of nano composites. In particular, in the case of nanotubes, the large polarizability at optical frequencies allows to reshape drastically the absorption of the chromophore. The compounds inherit the spectral absorption features of the chromophore and the polarization absorption features of the nanotube. Although no significant local field amplification is observed in the particular case of semi-conducting nanotubes (positive dielectric susceptibility), the effective absorption cross section of carbon nanotubes is clearly enhanced (by a factor of 5) due to the large number of attached chromophores and their large intrinsic absorption cross section. This opens the way to labeling applications where all semi-conducting species could be excited efficiently with one single wavelength. In addition, new effects are expected for metallic nanotubes for which the negative dielectric permittivity can lead to strong local amplification of the electric field through surface plasmon resonances. In that latter case, the effective absorption (or scattering) of attached chromophores could be strongly enlarged, which is another attractive possibility in labeling or medical applications.$^7$
Conclusion

We have investigated energy transfer in non-covalently bound porphyrin / carbon nanotube compounds at the single nano-compound level. Photoluminescence excitation spectroscopy reveals an efficient energy transfer from the porphyrin to the nanotube down to cryogenic temperatures in line with a purely electronic process. Emission and excitation polarization diagrams allow to address the question of local field effects in the overall optical response of the nano-compound. We show that the compound inherits the spectral absorption features of the porphyrin together with the anisotropy of the nanotube, resulting in a strongly anisotropic energy transfer resonance. This result is quantitatively modeled by the electromagnetic response of a dielectric cylinder, which strongly modifies the field structure at the tube interface. This study opens the way to applications where nanotubes could serve as optical antenna especially in the case of metallic nanotubes that could benefit from an enhancement of the local field due to surface plasmon resonances.

Methods

The preparation of the micellar suspension of functionalized nanotubes is described in Ref.?

Micro-PL setup

In order to investigate the energy transfer mechanism at the single nanocompound scale, we deposited a drop of the suspension on the back side of a zirconium oxide hemispherical lens (solid immersion lens). Atomic Force Microscopy (AFM) imaging shows that we reach a nanotube density of the order of 10 nanotubes per micrometer square with an average length of the nanotubes of the order of 500 nm. However, most of these nanotubes are not detected in our setup because we can only detect the semi-conducting ones with a band gap in the detecting range of our CCD camera. Therefore, only the (5,4), (6,4), (8,3), (9,1), (6,5) and (7,5) chiral families are observed. This leads to an effective detectable tube density of the order of $10^{-1} \mu m^{-2}$.

The solid immersion lens (SIL) configuration allows a strong enhancement of the signal due to an increase of the
effective numerical aperture and due to the reshaping of the emission diagram of the emitter in the collecting direction. In contrast to regular oil immersion objective, the SIL configuration is further compatible with low temperature (10 K) measurements. The optical setup is based on a home made confocal microscope. The light source is a high brightness cw Xe lamp filtered by a monochromator and injected in a small core optical fiber, or a He-Ne laser (632.8 nm). The excitation power ranges from 10 to 1000 W/cm². The injected and collected beams are polarized and analyzed by a rotating achromatic half-wave plate associated to a polarizer with a fixed direction. The collected light is dispersed in a 32 cm spectrometer and detected by a nitrogen cooled Si CCD camera.

**PLE measurements**

PLE spectra are obtained by recording an full PL spectrum for each excitation wavelength. Then, for each PL spectrum, the PL line of the nanotubes is extracted, integrated over its width and divided by the incident excitation power. This normalized PL intensity is reported as the PLE intensity for each excitation wavelength.

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