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Fluorescent self-assembled molecular monolayer on graphene

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ABSTRACT: We report the first fluorescent molecular self-assembly on graphene. The quenching of the fluorescence of the adsorbed dye by the adjacent graphene is hindered at the molecular scale based on a spacer approach, through a specifically designed dual-functionalized self-assembling building block. This 3D tecton presents two faces, one forming a non-covalent graphene-binding pedestal and the other carrying a dye group linked by a spacer to the pedestal. The spontaneous ordering of the adsorbed layer is investigated by scanning tunneling microscopy whereas the resulting optical properties of the whole graphene-dye hybrid system are characterized by absorption and fluorescence spectroscopies.

Graphene has focused intensive research in the past ten years due to its unusually high electron mobility, atomic thickness, broadband optical absorption and unique flexibility.$^{1,2,3,4}$ CVD-grown graphene being easily transferable onto arbitrary substrates while preserving high electronic mobilities$^5$ it soon appeared as a tantalizing candidate for various applications in photonics$^6$, such as solar cells$^7$, high-speed photodetectors,$^8$ light sources,$^9$ ultrafast lasers$^{10}$ or metamaterials$^{11}$. However, as a zero-bandgap semimetal, pristine graphene has a narrow range of roles, merely limited to transparent electrodes. Advanced applications require graphene to be synergistically combined with nanomaterials providing complementary properties.

Recently, non-covalent functionalization of graphene with organic molecular building blocks has appeared as a promising way to modulate its properties in view of functional applications. Actually, graphene provides an atomic-scale crystallographic lattice acting as a template guiding supramolecular self-assembly.$^{12}$ This bottom-up elaboration process, which implies the physisorption onto graphene of mostly planar molecules (tectons), is now well mastered in view of electronic applications. For example, supramolecular self-assembly on CVD grown graphene was used to dope graphene and back-gated graphene field-effect transistor (G-FET) devices were obtained.$^{13}$

Surprisingly, by comparison with electronics, the non-covalent functionalization of graphene is still in its infancy as concerns applications in photonics. Yet, organic dyes offer a high flexibility in the design of innovative photonic devices. Actually, because of their high oscillator strengths, they can play the role of light harvesters, photon sources, exciton funnels, etc. and as such should provide opportunities to enhance or extend the properties of graphene towards light-based applications.

Very recently, the distinctive optical absorption signature of self-assembled dye-graphene architectures, the structure of which were confirmed by scanning tunneling microscopy, have been analyzed by absorption microspectroscopy$^4$. However, the main challenge remains the emission of light by such systems, since graphene is known as a strong quencher of electronic excited states through ultrafast non-radiative electron-tunneling exchanges (the so-called Dexter excitation transfer) between the dyes and the graphene$.^{5,6,7}$ Hence, the realization of light emitting devices based on dye-graphene assemblies requires to overwhelm Dexter transfers through introduction of an accurately controlled dye-to-graphene electron barrier, for which planar flat-lying molecules are inefficient. Self-assembling building blocks of increasing three-dimensional (3D) complexity emerged recently, in particular following the so-called Janus tecton paradigm$^{8,9}$. These systems have been proposed as a platform to attach functional molecular moieties at a controlled distance from graphene$^{10}$.

Here, we demonstrate the first fluorescent supramolecular self-assembly on graphene with a specifically designed 3D Janus tecton. We investigate its two-dimensional ordering by scanning tunneling microscopy (STM) and its surface optical properties by absorption and fluorescence spectroscopies.
Results and discussion

The 3D Janus tectons are dual-functionalized and present two faces linked by a rigid spacer. One face is designed so as to steer the self-assembly on graphene and, for the present study, the other is a dye unit. More specifically, the dual-functionalized 3D tecton consists of a 3,4,9,10-perylenetetracarboxylic diimide (PTCDI) fluorophore linked to a graphene-binding pedestal by a dithiaparacyclophane pillar and a long leash made of one flexible alkyl chain and a stilbenic unit (scheme 1).

The pedestal unit consists of two molecular clips likely to steer the self-assembly of the whole entity. The clipping motif consists of two C\textsubscript{10}H\textsubscript{21} alkyl chains which, upon physisorption, lay flat on the surface and aligned along the <110> crystallographic direction of graphene. When two such units interdigitate, their lateral self-assembly leads to the formation of lamellae whose crystallographic structure is equivalent to the one observed upon adsorption of linear alkanes. The resulting enhanced 2D crystallization energy is at the origin of the binding nature of the clip. In the following paragraphs, this functional 3D Janus tecton will be designed as PTCDI-JT.

The spectroscopic properties of PTCDI-JT either in solution or self-assembled on graphene will be compared to those of a model system, namely N,N’-ditridecyl-3,4,9,10-perylenedicarboximide (PTCDI-C\textsubscript{13}). PTCDI-C\textsubscript{13} self-assembles on various substrates such as MoS\textsubscript{2}, graphite (Highly Oriented Pyrolytic Graphite, HOPG) or epitaxial graphene.

The self-assembly properties of PTCDI-JT on sp\textsuperscript{2}-hybridized carbon-based substrates were investigated by STM at the liquid/solid interface at room temperature. The roughness imposed on the CVD graphene monolayer by the underlying polyethylene terephthalate PET substrate makes STM observations difficult due to the poor contrast of molecules compared to this roughness. More accurate observations are obtained with an HOPG substrate and will be preferentially used in what follows. STM data recorded on a graphene substrate, including transferred CVD monolayers, show the equivalence with HOPG and are available in the Electronic Supplementary Information.

The image sequence displayed in figure 1 corresponds to observations recorded at the liquid / HOPG interface for three different solutions: (a) 1:4 ratio solution of PTCDI-JT to the PTCDI-JT pedestal, i.e. the naked precursor clipping unit which constitutes the base of the Janus tecton, (b) an equimolar mixture of the PTCDI-JT and PTCDI-JT pedestal, and (c) a solution of pure PTCDI-JT.

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Figure 1. STM images at the liquid/HOPG interface at room temperature, a) 1:4 mixture of Janus tecton and pedestal molecule (60 nm x 60 nm, It = 8.6 pA, Vt = -1320 mV), b) 1:1 mixture of Janus tecton and pedestal molecule (50 nm x 50 nm, It = 9.9 pA, Vt = -1602 mV), c) Janus tecton solution (50 nm x 50 nm, It = 8.6 pA, Vt = -1596 mV).
The pedestal unit is known to self-organize at the graphite surface forming a well-established two-dimensional network with the following parameters: $a = 3.8 \text{ nm}$, $b = 2.1 \text{ nm}$, $\alpha = 64^\circ$.\(^{25}\) Accordingly, STM images obtained with the 1:4 mixture (figure 1a) display large domains of a well-ordered arrangement of the clipping molecules of equivalent surface lattice. Experimentally, thanks to a smaller molecule size which reduces its steric constrains, the clipping molecule is shown to self-assemble more easily on graphite than the 3D Janus tecton.

Upon increase of the PTCDI-JT relative concentration in the applied solution, STM images of the surface display smaller domains of the clipping molecule arrangement together with areas presenting bright spots with rough alignments reminiscent of the underlying ordered regions (figure 1b). Both domains appear in similar proportions to that in the solution. These misaligned bright features are the signature of the physisorbed PTCDI-JT molecules.

Indeed, as far as molecular contrast is concerned, STM favors the signature of aromatic skeletons which present higher electronic surface density of states close to the Fermi level\(^{26}\). In the present case, the long and flexible alkyl spacer permits various locations of the PTCDI upper group relative to its pedestal concealing the periodic signature of the clipped pedestals in the horizontal plane. Still, the spatial distribution of these terminal PTCDI groups is not completely random suggesting that neighboring dye groups do aggregate themselves to form few-dye clusters.

For a pure solution of PTCDI-JT, the STM images show a quasi-random distribution of bright spots (figure 1c) tentatively associated to clusters of aggregated neighboring PTCDI dye moieties and the pedestal network is completely hidden.

To get rough estimates of the dye cluster size and spatial distribution, we built an approximate model based on simple molecular mechanics. Estimating the distance between the end of the fluorescent unit and the dithiaparacyclophane bridge in PTCDI-JT to be ~4 nm (see scheme 1), we computed the possible number of overlapping fluorescent units in every point of the overlaying 3D space based on the known surface lattice parameters of the two-dimensional clipped network (See figure 2 and figure S2 in Electronic Supplementary Information). In this rough model, the aggregation of independent molecules was modelled with all molecules found at the same location, that is the dimensions of the cluster itself are neglected. This simulation shows the possible aggregation of up to nine fluorescent groups belonging to different neighboring PTCDI-JTs.

Looking at the dimensions of the clipped network surface lattice, the parameter $a$ along the interdigitated alkyl chain direction is approximately twice the parameter $b$ along the lamella axis. Consequently, the dye upper groups are expected to interact more importantly with their respective neighbors along the $b$ direction. One can then reasonably predict the existence of stronger interacting subclusters of an even number of molecules. More specifically, the detailed simulation yield a bimodal distribution of the subcluster sizes, whose principal peak corresponds to a cluster size value of 6 ± 1. (See Figure S3 in Electronic Supplementary Information).

![Figure 2. Map of the self-assembled PTCDI-JT tecton indicating the maximum number of aggregated dye-moieties, obtained as the number of intersecting half spheres assuming each dye is located within a 4-nm distance from the center of its pedestal (white crosses).](image)

The most probable location of such 6-dye cluster is computed to be 1.6 nm above the dithiaparacyclophane rigid pillar and 2.8 nm or 4.2 nm away from the bridge in the horizontal plane. The pillar height amounts to about 0.6 nm. The adsorption of the PTCDI-JT pedestal is driven by the physisorption of its aliphatic chains. The aromatic core of the basal unit lies flat on the graphene sheet at a respective distance of 0.33 nm.\(^{27}\) So, adopting a static view of the PTCDI-JT arrangement, the clusters of dye moieties lay at about 0.33 + 0.6 + 1.6 = 2.53 nm above the graphene substrate (See Figure S4 in Electronic Supplementary Information).

Experimental UV-visible absorption spectra of the PTCDI-JT and the PTCDI-C\(_{13}\) molecules in toluene are reported in figure 3.

The light absorption by the clipping motif manifests itself in the UV range of the spectrum ($\lambda < 420 \text{ nm}$) and corresponds to the one recorded for the PTCDI-JT pedestal. The peak at 330 nm and the shoulder at 375 nm are assigned respectively to the four terminal stilbene units and to the central distibene unit. This shoulder in the PTCDI-JT absorption spectrum appears to be two times larger in accordance with the presence of another distibene in the spacer chemical group.

In the visible range 420 nm < $\lambda$ < 550 nm, the PTCDI-JT absorption matches very well that of the PTCDI-C\(_{13}\) dye. This unaltered spectroscopic signature of the core dye moiety demonstrates that the spacer effectively decouples the conjugated electron systems of fluorophore and pedestal units.
The absorption spectra of self-assembled monolayers of PTCDI-JT and PTCDI-C13 on graphite are compared with corresponding toluene solution in figure 4. The relative peak absorption level of PTCDI-JT monolayer (0.4%) is nearly 6-times smaller than that of PTCDI-C13 (2.3%). Considering a similar absorption cross section, a surface density of PTCDI-C13 of 0.5 mol/nm$^2$ and a 2/3 orientational factor since the PTCDI-C13 molecules are lying flat on the graphene surface$^4$ whereas PTCDI-JT dye moieties are randomly oriented, one can estimate a Janus tecton surface density of 0.12 ± 0.03 molecule per nm$^2$. This value is consistent with the surface density extracted from the surface lattice determination by STM of PTCDI-JT self-assembled monolayer on HOPG$^{20}$: 0.14 molecules/nm$^2$. This numerical estimate further supports the conclusion that PTCDI-JT are regularly self-assembled on the graphene surface following the expected binding scheme of the clipping pedestal units.

Both PTCDI-JT and PTCDI-C13 absorption lines drastically change when self-assembled onto graphene as compared with toluene solution (figure 4). Whereas the PTCDI-C13 absorption merely consists in a rigid red shift compared with the solution, the situation is more complex for PTCDI-JT. Despite a reduced signal-to-noise ratio, an increased weight of higher-order vibronic replicas is perceptible and the rising edge appears less abrupt, which can correspond to a spectral broadening. The red-shift observed for PTCDI-C13 is explained by the adsorption of its the aromatic core which lies flat on graphene at a typical π-stacking distance of 0.33 nm.$^{19}$ As inferred from STM analysis in the previous section, for PTCDI-JT, the corresponding distance is much larger. According to the static model discussed above, the distance from the dye moieties to the graphene is typically 7 times larger (~2.5 nm) which prevents such interactions. In contrast with PTCDI-C13, the dye moieties of PTCDI-JT form few-molecule clusters. The increase of higher-order vibronic peaks is typical of a H-aggregate behavior resulting from a π-stacking configuration inside clusters.$^{27, 28}$

Figure 3. Normalized solution absorption spectra of the PTCDI-JT pedestal, PTCDI-JT, and PTCDI-C13 in toluene.

Figure 4. Absorption spectra of PTCDI-JT and PTCDI-C13 self-assembled on graphene. The CVD graphene sheet has been transferred from Cu onto a transparent PET substrate prior to self-assembly. The measurements were done in transmission, referenced to naked graphene. Because of the absorption of PET in the UV, the spectra are limited to $\lambda > 400$ nm. The solution spectra of figure 3 are reproduced for an easier comparison.

The spectral broadening of the vibronic spectral features, already present self-assembled PTCDI-C13 is further increased for PTCDI-JT. This probably reflects the inhomogeneity of the dye-moiety assembly as evidenced by STM imaging of the PTCDI-JT tectons which evidences dye clusters with various sizes.

The main result of this work is presented in figure 5 in which the fluorescence signals of the PTCDI-JT molecule self-assembled on graphene is compared with that of the PTCDI-C13 reference as recorded in solution, in self-assembled monolayer on graphene or in solid state microcrystals as formed by casting the same PTCDI-C13 toluene solution onto naked fused silica instead of graphene. A photoluminescence signal of the PTCDI-JT self-assembly on graphene is readily measurable whereas no such signal can be detected from the PTCDI-C13 ordered domains on the same substrate. Hence, the design of PTCDI-JT based on a spacer between the dye and graphene actually reduces the energy transfer rate, as expected. The measured fluorescence quantum yield is still low, estimated in the range of $2 \times 10^{-5}$ to $2 \times 10^{-6}$ by integrating the fluorescence signal between 560 nm and 745 nm for an excitation at 532 nm. Since the radiative decay rate of PTCDI is of the order of $10^{-9}$ s$^{-1}$, the quenching rate is in the range $10^4$ s$^{-1}$ to $10^5$ s$^{-1}$. Although still high, this value is compatible with a residual Förster resonance energy transfer, the electron-exchange quenching process (Dexter transfer) being slower for the distances between dye and graphene that were estimated above.
Figure 5. Photoluminescence spectrum of self-assembled PTCDI-JT on graphene (blue) compared with PTCDI-C13 reference signals in toluene (green), self-assembled on graphene (black) and in microcrystal solid state (red). The molecules are excited by a CW laser operating at 532 nm wavelength which hides the principal fluorescence peak (0 – 0 line) for all the systems.

The formation of clusters also strongly affects the characteristics of luminescence. Although the above-mentioned H-aggregate coupling is supposed to slower the radiative decay in favor of the quenching, PTCDI-C13 is fluorescent in the solid state, as shown with the fluorescence of microcrystals formed on silica. Non-radiative relaxations are thus hindered by the strong interactions between stacked molecules. The fluorescence spectrum of PTCDI-JT self-assembled onto graphene (figure 5, blue curve) is intermediate between that of the PTCDI-C13 solvated in toluene (green curve) and that of PTCDI-C13 microcrystals on silica (red curve), showing a progressive displacement of the oscillator strength towards higher vibrational replicas with cluster size, consistently with the H-aggregate-like coupling deduced from absorption spectroscopy. A shorter exciton coherence length in self-assembled clusters is consistent with the involvement of a small number of dye groups, spatially limiting the coherent delocalization of the exciton which is of the order of 2 nm for similar crystals. Furthermore, the H-aggregation geometry could be perturbed from a perfect π-stacking by the spacer steric hindrance. Instead of being parallel, aromatic cores can adopt an angle that partially decouple the transition dipole moments. This suggests that the controlled formation of few-molecule dye clusters plays a role in the restoration of luminescence. Note that the Janus tecton surface fluorescence peak is slightly red-shifted from the PTCDI-C13 ones consistently with its red-shifted absorption.

In a forthcoming work, lifetime measurement of the radiative decay states will be carried out. This parameter will give insights into the balance between radiative and non-radiative exciton recombination processes.

In summary, we have achieved the first fluorescent molecular self-assembly on graphene. This milestone has been made possible thanks to the development of a 3D Janus tecton embedding a PTCDI dye uphold away from the substrate plane together with the use of CVD graphene as an optics-friendly 2D substrate. A fluorescent non-covalent functionalization of graphene could help to alter the surface photonic properties of this zero-gap semiconducting monolayer.

Methods

Graphene on PET were bought from graphene-supermarket. The self-assembled monolayer has been produced by dipping graphene on quartz in an approximately $10^{-8}$ mol/L Janus tecton solution with toluene as the solvent. To ensure the fabrication of a single monolayer, the sample has been rinsed abundantly with toluene.

The STM images were recorded under ambient conditions (room temperature $T \sim 300$ K) with a homemade digital system. Imaging proceeds by the immersion of a 250 μm mechanically cut tip of Pt/Ir (90/10, Goodfellow) into a 3 μL droplet of phenyloctane (Aldrich, 98%) containing the molecules at a concentration between $10^{-6}$ and $10^{-5}$ mol/L and deposited on a freshly cleaved 10 mm x 10 mm HOPG surface (Goodfellow). The scanning piezoelectric ceramic was calibrated in XY directions with the help of atomically resolved pictures obtained on HOPG. In order to ensure a correct statistical representation of our measurements concerning the structural organization of the monolayers, several images were recorded at different locations of the samples for each molecule of interest. All the images presented were obtained at a quasiconstant current in the variable height mode (the bias voltage VT and the tunneling current IT are indicated in figure captions) and were corrected for the thermal drift by combining two successive images with downward and upward slow-scan directions.

Absorption spectra were recorded on a Perkin-Elmer Instrument equipped with a tungsten filament white lamp. The sample has been excited with a 532 nm CW diode laser through an Olympus objective (NA = 1.3, x100) and a cooled Andor CCD camera has been used to characterize the fluorescence signals.

Molecular mechanics calculations were performed with a chemical simulation software (Accelrys Discovery Dassault System BioVia).

ASSOCIATED CONTENT

Supporting Information. Synthesis of PTCDI-JT, STM images on graphene and molecular mechanics calculations. This material is available free of charge via the Internet at http://pubs.acs.org."
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ABBREVIATIONS
2D, two dimensional; 3D, three dimensional; CCD, charge coupled device; CVD, chemical vapor deposition; CW, continuous wave; HOPG, highly ordered pyrolytic graphite; PET, polyethylene terephthalate; PTCDI, 3, 4, 9, 10-tetrayanoanthracene; STM, scanning tunneling microscopy

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