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Photoinduced energy and electron transfer at graphene quantum dot/azobenzene interfaces[†]

Magdalena Kaźmierczak, ^a Samuele Giannini ^{‡b} and Silvio Osella ^{*a}

To respond to the constant miniaturization of modern devices, it is imperative to develop novel classes of hybrid materials with outstanding functionalities. One of the most promising developments in this regard is photoswitchable nanoelectronics, which offer non-invasive structural changes *via* irradiation. Azobenzenes, in particular, represent a widely investigated group of compounds due to their reversible photoisomerization and flexibility to introduce modifications by functionalizing the aryl ring. Here, an interface consisting of graphene quantum dots (GQDs) and a pyrene functionalized azobenzene (AZO) is built and investigated to unravel the energy and charge transfer dynamics at play. The influence of the conformation, isomerization and thermal effects on the photophysics is described using a multiscale computational approach coupled to electronic structure calculations. By computing the photoinduced energy and charge transfer rates we found that the small exciton reorganization energy and favorable alignment of the energy levels at the interface favor excitation energy transfer. A significant increase of photoinduced hole transfer from AZO to GQDs is observed when thermal effects are considered. Moreover, these photoinduced processes for the azobenzene molecules in their *trans* configuration are always faster than those related to the *cis* isomer. This is due to a favorable overlap and a stronger interaction of the *trans* isomer, considered the active state, with the GQDs. On the other hand, the *cis* configuration, featuring slower photoinduced energy and charge transfer processes, can be considered as the inactive species. Nevertheless, its contribution to the overall photophysics remains non-negligible.

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Introduction

Graphene quantum dots (GQDs) are unique, zero-dimensional materials derived from graphene with properties depending on the size of the particles.¹ Unlike graphene, GQDs are semiconductors and exhibit a tunable non-zero band gap due to their constricted size^{2,3} as well as their edge geometry⁴ and functionalization.^{5,6} Quantum confinement and edge effects are also responsible for improving GQDs' optical properties compared to other carbon allotropes, not only by opening their band gap but also by enhancing their dispersibility. The highly adjustable optical properties such as photoluminescence,

electrochemiluminescence and photostability, together with the simplicity of functionalization, make GQDs highly versatile materials.⁷ The main drawback – low photoluminescence quantum yield – can be significantly improved by further changes in their surface composition or doping.^{8,9} The formation of an interface between GQDs and small organic molecules can lead to functional materials, whose optoelectronic properties can be easily tuned by incorporating changes within the molecules.^{10–13} GQDs are applied to a number of applications such as solar cells,^{14–17} light-emitting diodes^{18–21} and supercapacitors.^{22–24} Moreover, their low toxicity and biocompatibility enable the use of these materials in medical bioimaging²⁵ and drug delivery applications.²⁶

The ability to control a device based on a GQD-small molecule interface with an external stimulus, for example through irradiation, is a further step towards enhancing the complexity of the device. Azobenzene can be considered as an archetype in this field since full control over geometrical changes is enabled by photoisomerization *via* absorption of a photon. Since *cis/trans* isomerization is a fully reversible process,²⁷ the different sets of optoelectronic properties of two isomers can be accessed and swapped.^{28,29} Additionally, the photoswitching performance at different excitation wavelengths and/or intensities, which is a result of those fundamental

^a Chemical and Biological Systems Simulation Lab, Centre of New Technologies, University of Warsaw, Banacha 2C, 02-097 Warsaw, Poland.

E-mail: s.osella@cent.uw.edu.pl

^b Laboratory for Chemistry of Novel Materials, University of Mons, 7000 Mons, Belgium

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[‡] Present address: Institute of Chemistry of Organometallic Compounds, National Research Council (ICCOM-CNR), I-56124 Pisa, Italy.



properties, can be readily modified by adding electron donating or withdrawing groups in the *para* position of the benzene ring forming an azobenzene molecule. As a consequence, tunable on/off states in which one isomer facilitates charge or energy transfer whereas the other inhibits this process, allow using these materials as switches in biosensors,³⁰ energy storage devices^{31–33} and molecular machines.³⁴

Upon excitation of a photoactivated interface three distinct processes can occur: photoinduced electron transfer (PET), photoinduced hole transfer (PHT) and excitation energy transfer (EET). When considering PET and PHT, an excitation of a generic donor/acceptor interface leads to excitation of an electron from the ground state to the excited state of either the donor or acceptor, respectively. The excitation process is followed by the transfer of an excited electron from the donor to the acceptor in PET, while a hole is transferred from the acceptor to the donor in the PHT process. On the other hand, EET involves a simultaneous transition of electrons from an excited state to the ground state for the donor and *vice versa* for the acceptor (Fig. 1(a)).

In this study, the optoelectronic properties of a push-pull 4-[(4-aminophenyl)diazenyl]phenyl 4-pyrenebutanoate (hereafter labelled as AZO, Fig. 1(b)) molecule physisorbed on GQDs are investigated through multiscale simulations combining molecular dynamics and electronic structure. The AZO molecule considered in this study has already been synthesized and reported in the literature.³⁵ Webb and coworkers reported the synthesis and absorption properties of a series

of AZO-pyrene molecules connected with an ester linker, in which substituents (either NH₂ or OH) were added in the *para* position of the azobenzene moiety. This allowed the modulation of the absorption spectra, and for the amine substitution the authors observed that the strong electron-donating effect was responsible for a red shift in the spectrum with respect to pyrene. In this work, we focus on the amine-substituted system and compute the UV-Vis spectra. Along with the examination of the optical properties of the AZO-pyrene molecules, we examine here the excited state characters and describe possible charge or energy transfer processes in the single molecule and at the interface of the AZO-pyrene and the GQDs.

Specifically, by calculating the interaction between different states (*i.e.*, electronic couplings), the reorganization energies of the various photoinduced processes, driving force and finally the Marcus rates between electron donor and acceptor sites, we aim to describe the photophysics of the GQD–AZO interfaces. We first consider the isolated AZO molecule to assess the possibility of such a system undergoing intramolecular excited state processes upon irradiation (Fig. 1(b)). Marcus rates were computed to rationalize various competitive pathways. Next, we discuss how the formation of an interface between the AZO molecule and a GQD (Fig. 1(c)) alters the optical and electronic properties and the photophysics at the interface. Finally, we take into account thermal effects in molecular dynamics simulations to assess the effects of structural disorder and temperature on the transfer rates.

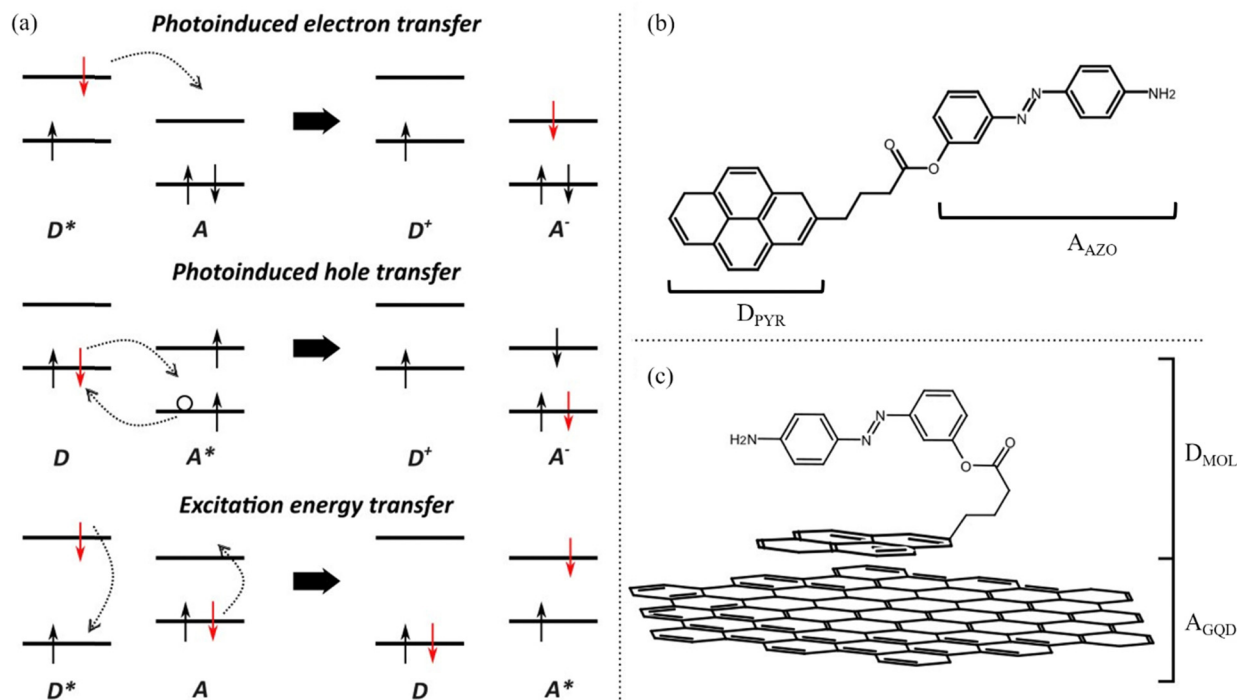


Fig. 1 (a) Schematic representation of the photoinduced processes, (b) structure of the AZO molecules and (c) of the GQD–AZO interface considered in this study. Note that in the isolated molecule the pyrene moiety is the electron donor (D_{PYR}) while the azobenzene one is the electron acceptor (A_{AZO}), and for the GQD–AZO interface the electron donor is the AZO molecule which we refer to as (D_{MOL}), while the electron acceptor is the GQD (A_{GQD}).



Methodology

Azobenzenes can be converted from the *trans* to the *cis* isomer at a specific wavelength of UV light, which relates to the $\pi \rightarrow \pi^*$ excitation, whereas the opposite isomerization appears at longer wavelengths (in the visible range) and corresponds to the $n \rightarrow \pi^*$ transition. The characteristic trait of the investigated AZO molecule is that upon irradiation it undergoes photoisomerization as the parent azobenzene.³⁵ In order to systematically evaluate the optoelectronic properties' changes occurring when AZO molecules are interacting with a GQD, isolated AZO molecules in different conformations as well as GQD-AZO interfaces are optimized and analyzed.

First, isolated AZO molecules as well as GQD-AZO interfaces are optimized at the DFT level using the CAM-B3LYP functional³⁶ together with GD3 van der Waals dispersion³⁷ with the 6-31G(d) basis set³⁸ as implemented in the Gaussian 16 software.³⁹ The PET, PHT and EET processes at the interface are evaluated by combining TD-DFT calculations performed with the same functional and basis set used for the geometry optimization and evaluation of the transfer rates (as explained below).

The GDQ-AZO interface is built by considering 42 AZO molecules (either *trans* or *cis*) physisorbed on graphene and then performing molecular dynamics (MD) simulations to assess the stability and magnitude of the interactions taking place at the interface. The simulations are performed using the GROMACS package⁴⁰⁻⁴⁵ with the CHARMM36 force field.⁴⁶ To correctly describe the AZO *trans* and *cis* isomers, two different sets of parameters for the N=N bond, angles and dihedrals are taken from the literature.²⁷ Dihedral angles for the ester bond are reparametrized at the MP2 level of theory using Gaussian 16 software, whereas the ESP charges for the AZO are taken from the optimized structure at the DFT level. Both interfaces are solvated with water molecules using the TIP3P description for the solvent. The calculations are performed at room temperature using the *NVT* ensemble and the Nose-Hoover thermostat (300 K and 0.5 ps). Periodic boundary conditions are considered while building the cell, which has dimensions equal to $a = 8.08$ nm, $b = 8.11$ nm and $c = 5.0$ nm to prevent spurious interaction with repeating images along the c direction. 60 ns MD simulations with a time step of 2 fs are run to equilibrate and minimize the energy of the system (the estimated error of total energy and Lennard-Jones (LJ) interactions between the GQDs and AZO fell below 20 kJ mol⁻¹, which is less than 0.01% for average total energy and less than 0.5% for LJ interactions). A 30 ns production simulation is sampled, and from it, 50 frames, consisting of AZO and GQDs (114 carbon atoms in a hexagonal shape with all armchair edges), are extracted along the trajectory. The selected frames are subsequently considered as input structures for TD-DFT calculations, to assess the thermal effects on the optical properties (such as the absorption spectrum) and electronic properties (such as electronic couplings and reorganization energies).

To characterize the photophysics of the systems and the characteristic rates of the different EET, PET and PHT

processes, we adopt here the well-established Marcus rate equation. In the weak coupling and the high-temperature classical limits, the rate constant (k_{if}) for nonadiabatic transfer between two diabatic states (referred to as the initial and final states, $|i$ and $|f$) can be written as:

$$k_{if} = \frac{2\pi}{\hbar} \cdot V_{if}^2 \cdot \frac{1}{\sqrt{4\pi\lambda k_B T}} \cdot \exp\left(-\frac{(\lambda + \Delta G)^2}{4\lambda k_B T}\right) \quad (1)$$

where V_{if} is the coupling between the diabatic electronic wave functions which is computed as described below using a powerful diabatization scheme called multi-state fragment charge transfer-fragment exciton transfer (MS-FED-FCD).⁴⁷⁻⁴⁹ In eqn (1), λ is the total reorganization energy of the process which is defined as the sum of the internal and external contributions $\lambda = \lambda_i + \lambda_0$, where the external reorganization energy λ_0 is set to 0.05 eV.⁵⁰ Throughout the study, we will consider the total reorganization energy.

The internal reorganization energy (λ_i) instead is the energy cost to bring the nuclear configuration of the initial state into that of the final state while keeping the electronic configuration fixed at the initial state and was computed *via* the so-called four-point scheme:⁵¹

$$\lambda_{PET} = E^-(Q_N) - E^-(Q^-) + E_N(Q^-) - E_N(Q_N) \quad (2)$$

$$\lambda_{PHT} = E^+(Q_N) - E^+(Q^+) + E_N(Q^+) - E_N(Q_N) \quad (3)$$

where $E_N(Q_N)$ and $E^{+/-}(Q^{+/-})$ are the total energy of the optimized neutral and charged structures, while $E^{+/-}(Q_N)$ and $E_N(Q^{+/-})$ are the total energy after the vertical ionizations.

When considering the interface consisting of donor and acceptor fragments, one can separately calculate the internal reorganization energy as an average of the backward (λ_{BW}) and forward transitions (λ_{FW} , see the ESI† for more details).^{47,52,53} Finally in eqn (1), ΔG is the free energy difference between the initial and final states, and is approximated in this work as the energy difference between the states that are coupled.

The electronic couplings are computed considering both the intramolecular couplings between the electron donor pyrene (D_{PYR}) and electron acceptor azobenzene (A_{AZO}) moieties in the isolated AZO molecules, and intermolecular coupling between the *trans/cis* pyrene-azo molecule (D_{MOL}) and the electron acceptor GQDs (A_{GQD}) in the interfaces considered (see Fig. 1(b) and (c)). As mentioned above, to quantify these interactions, we used a multi-state diabatization procedure (MS-FED-FCD).⁴⁷⁻⁴⁹ This method involves the partition of a given donor-acceptor pair into two fragments and by using appropriate additional operators, the adiabatic Hamiltonian of the dimer (formed by two or more adiabatic states of the system) is transformed into a fragment-related diabatic basis, to subsequently extract the couplings between states of different characters from the Hamiltonian matrix written in such a localized basis. In practice, first, the adiabatic excited states of the donor-acceptor pair are calculated followed by diabatization to localized, (maximally) diabatic excitonic states. Then, the couplings can be found as the off-diagonal elements of this matrix. Importantly, MS-FED-FCD allows us to include several



adiabatic excited states of the donor–acceptor system in the diabatization procedure (in this work, we used up to 40 excited states) to ensure a complete de-mixing between excitations of different nature and an optimal reconstruction of localized excitonic and charge transfer states. This is particularly useful for the investigated system where there is a high density of states and a given adiabatic state is the combination of many diabatic states of both D and A moieties. We refer to ref. 47–49 for a more detailed description of the method. When degenerated states are present in the system, the procedure is modified to consider the effective coupling (V_{eff}) computed as the square root of the sum of the squared couplings related to the number of degenerate states found in the calculation. For example, for two degenerate initial states i_1, i_2 and two degenerate final states f_1, f_2 , the effective coupling will be:

$$V_{\text{eff}} = \sqrt{V_{i_1 f_1}^2 + V_{i_1 f_2}^2 + V_{i_2 f_1}^2 + V_{i_2 f_2}^2} \quad (4)$$

Results and discussion

Isolated AZO molecules

Before considering the optoelectronic properties of the interfaces, we assess the properties of the isolated AZO molecule for both its *trans* (*t*AZO) and *cis* (*c*AZO) isomers, and the possible intramolecular photoinduced processes which can occur from the different isomerization. As reported in the literature,³⁵ two different conformations can be present, in which the AZO part of the molecule is either perpendicular to the pyrene moiety

(labelled as ‘open’) or parallel to it (labelled as ‘fold’) (see Fig. S1, S2, ESI[†] and Fig. 2).

As the *t*AZO fold conformation is more stable than the open one by 0.4 eV (in agreement with the 0.3 eV energy difference reported in the literature),³⁵ we will consider only the fold conformers for further analyses (details of the open conformers can be found in the ESI[†]). As expected, and in agreement with the parent molecule, *t*AZO is found to be the more stable state compared to *c*AZO, with a stabilization energy of 0.7 eV. The modifications of the AZO structure compared to the parent molecule are also responsible for the dipole moment of 1.26 Debye for *t*AZO and 6.4 Debye for *c*AZO, due to the presence of both the amino group and the pyrene moieties. Moreover, for both isomers, the HOMO energies have similar values of −6.3 eV, while the LUMO has an energy of −0.5 eV for *t*AZO and it is slightly lower for the *c*AZO molecule (−0.3 eV). This leads to a similar wide energy gap of about 6.0 eV for both isomers. When considering the localization of the frontier molecular orbitals (FMO), we observe that for the *trans* isomer, the HOMO is localized on pyrene, whereas the LUMO is localized on the AZO moiety, suggesting the formation of an intramolecular charge transfer state upon excitation (Fig. 2(a)). On the other hand, for *c*AZO, the HOMO is delocalized over the whole molecule, while the LUMO is localized over the pyrene moiety, suggesting a different behavior in intramolecular charge transfer ability for this isomer (Fig. 2(b)).

The absorption spectra of the two isomers are reported in Fig. 2(c). Four main peaks can be found for *t*AZO in the absorption spectrum, with the fold conformation bathochromically

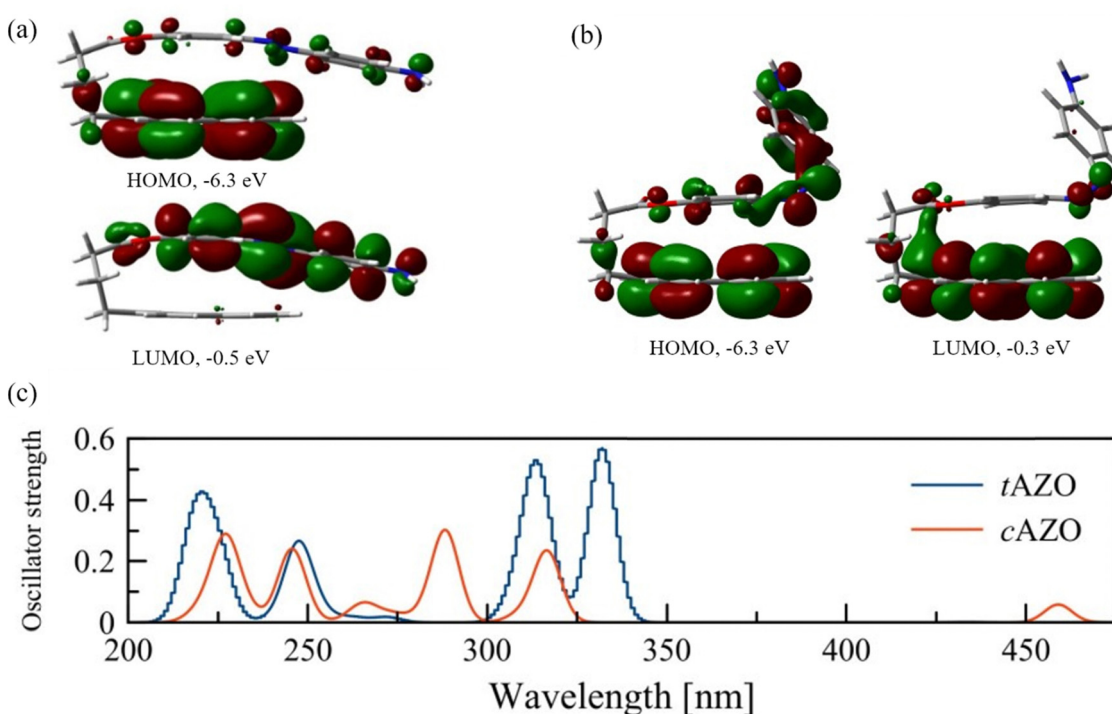


Fig. 2 Frontier orbitals of the isolated AZO molecules: (a) *trans* isomer, (b) *cis* isomer. (c) Computed UV-Vis spectra for the isolated AZO molecules in *trans* (*t*AZO) and *cis* (*c*AZO) configurations.



shifted by approximately 5 nm with respect to the open one (details for the open conformation are reported in the ESI†). The first excited state is a forbidden $n \rightarrow \pi^*$ transition (HOMO-3 to LUMO, see Table S1 (ESI†) for a detailed description of all transitions). The first bright peak at 332 nm relates to the second excited state (S_2) and corresponds to the HOMO-1 to LUMO transition ($\pi \rightarrow \pi^*$). The next peak at 314 nm corresponds to the HOMO to LUMO+1 transition. For the *c*AZO isomer, similar absorption peaks are obtained compared to *t*AZO, albeit blue-shifted by 20 nm. In addition, the first excited state is now allowed, and it shows a weak peak at 459 nm, which corresponds to the $n \rightarrow \pi^*$ (HOMO-1 to LUMO+1) transition. This observation confirms that AZO undergoes photoisomerization because a similar transition is responsible for this process in the unsubstituted *cis*-azobenzene.⁵⁴ The next peak at 317 nm corresponds to the $\pi \rightarrow \pi^*$ HOMO to LUMO transition. Interestingly, our results well agree with the reported absorption in the literature,³⁵ although some expected discrepancies are present due to the absence of solvent contributions in our calculations.

To gain deeper insight into the charge and energy transfer rates of the AZO isomers, we compute the natural transition orbitals (NTOs) and intramolecular coupling for the investigated AZO isomers. This analysis suggests that the electron donor site is the pyrene moiety (D_{PYR}), whereas AZO acts as the acceptor (A_{AZO}) (see Fig. S3, ESI†). The energy transfer process taking place due to pyrene local excitation at 314 nm to AZO local excitation at 332 nm features a coupling value of 46.4 meV. A nearly identical coupling value is found for the PET process of the same isomer, which occurs from the excited D_{PYR}^* at 314 nm, transferring an electron to the AZO moiety. A considerably higher coupling value of 64.3 meV is found for PHT, which involves AZO excitation at 332 nm and AZO to pyrene charge transfer state. Similar trends can be observed for *c*AZO for which both EET and PET couplings have rather low values of 9.0 meV and 8.9 meV, respectively, compared to 105.5 meV for the PHT coupling, which occurs between the AZO local excitation at 288 nm and the AZO to pyrene charge transfer state at 246 nm (see Table 1). On the whole, we can conclude that the better overlapping states of the *trans* configuration tend to favor greater interactions with respect to the *cis* form.

To quantify the transfer rates, we calculate next the reorganization energies for the different processes considering the four point model (eqn (2) and (3)) described in the Method section. For the PET process (Fig. 1(a)), we compute a λ_{PET} of

Table 1 Total reorganization energy, electronic coupling, driving force and Marcus rates for the three processes considered in this study, for the isolated AZO molecule

	Process	λ [eV]	Coupling [eV]	ΔG [eV]	k [s^{-1}]
<i>t</i> AZO	EET	0.19	0.046	-0.220	8.0×10^{13}
	PET	0.80	0.045	-0.048	3.9×10^{10}
	PHT	1.28	0.064	0.172	6.9×10^6
<i>c</i> AZO	EET	0.21	0.009	0.335	3.2×10^6
	PET	1.38	0.009	1.083	2.9×10^{-7}
	PHT	1.14	0.106	0.748	1.0×10^1

0.80 eV for *t*AZO, and a nearly double value for *c*AZO of 1.38 eV (a similar situation holds true for the open conformers, see the ESI† for more details). The difference in reorganization energies for holes between the two isomers for the PHT process is much less pronounced compared to the PET process, with values of 1.28 eV and 1.14 eV for *t*AZO and *c*AZO, respectively. As a result, the geometrical changes related to *c*AZO require less energy for the hole transfer, whereas *t*AZO should favor the PET process. The high reorganization energy values are due to the strong geometrical distortions arising from the presence of an additional electron/hole in the structure. For PHT, we observe planarization of the NH_2 group and changes in the CNNC dihedral angle for both isomers; for PET, there is a bending of the *t*AZO structure towards the pyrene moiety, while for *c*AZO, we observe an opening of the CNNC dihedral angle from 9 to 42 degrees (see Fig. S4, ESI†). Much lower values of 0.19 and 0.21 eV have been obtained for the EET process for *t*AZO and *c*AZO, respectively, suggesting that this process might be favored from a geometrical point of view due to much less severe molecular distortions (Fig. S4, ESI†). These results are confirmed by the analysis of the driving force, for which low values for EET are observed for both *t*AZO and *c*AZO isomers (Table 1).

With all the computed parameters, we can calculate the Marcus rates for the three processes, reported in Table 1. First of all, we can observe that the rates for the three processes – EET, PHT, PET – are always higher for *t*AZO compared to *c*AZO, suggesting that the former can be considered as an active state for intramolecular photoinduced charge/energy transfer. Going into the details of the specific mechanism, we observe that for *t*AZO, the EET is favored, with a k_{EET} of $8 \times 10^{13} \text{ s}^{-1}$, compared to the photoinduced charge transfer processes, following the $k_{EET} > k_{PET} > k_{PHT}$ trend. Although the coupling value for PHT is the highest among the three different processes, its reorganization energy and driving force also have the highest values, resulting in a lower rate for hole transport. On the other hand, for *c*AZO, the $k_{EET} > k_{PHT} \gg k_{PET}$ trend is found. EET is once more the favored process, with a k_{EET} of $3.2 \times 10^6 \text{ s}^{-1}$, while PET is now completely inhibited (due to very low coupling and high values of both reorganization energy and driving force). PHT shows very slow rates, despite the high coupling value, because of the strong geometrical deformation translating into a high reorganization energy value of 1.14 eV. As a result, the intramolecular EET is the main process occurring upon irradiation, with a higher rate for the *trans* isomer compared to the *cis* isomer, making *t*AZO the active state for this photophysical process. As the parent azobenzene molecule, the system investigated here can be used not only as an optical probe (due to the *cis*–*trans* isomerization) but also as an efficient on/off switch in which the *trans* isomer is the active state.

GQD–AZO interface

Next, we built an interface between the AZO molecule and a GQD to study its intermolecular photophysical properties. For this system, we consider the GQDs as an electron donor and the whole pyrene–AZO molecule as a single accepting unit



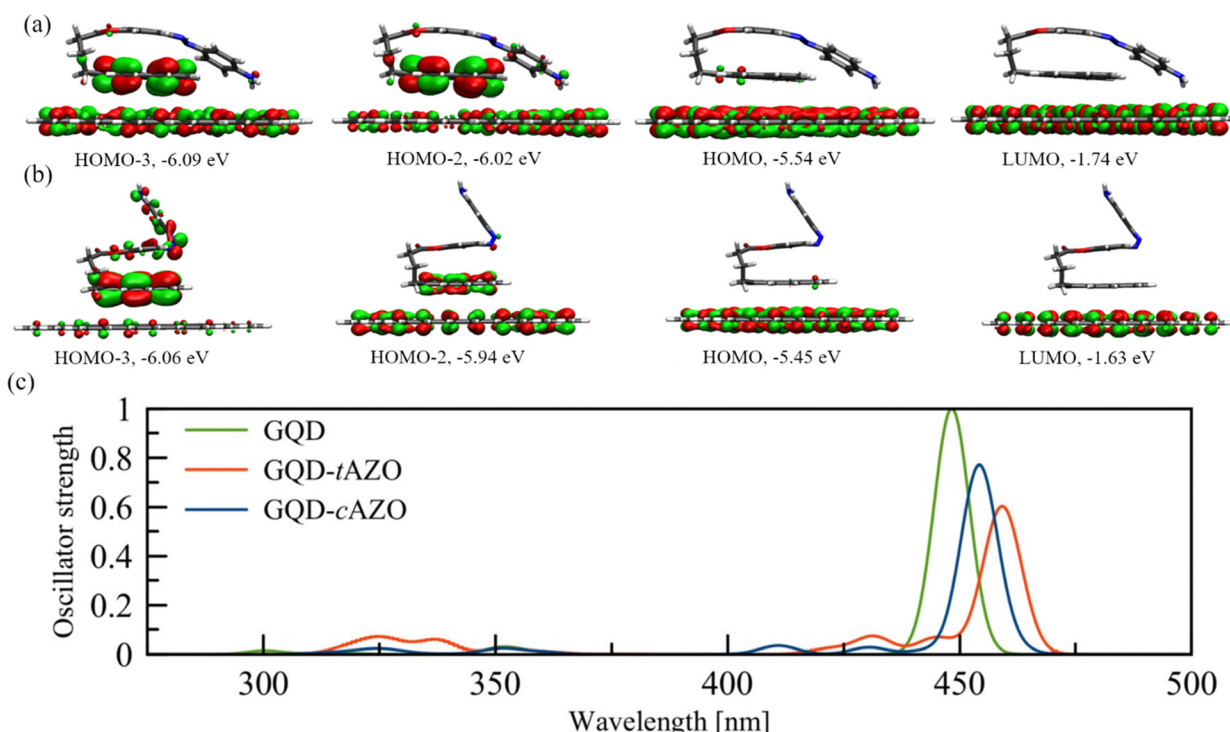


Fig. 3 Frontier orbitals of (a) GQD-tAZO and (b) GQD-cAZO. (c) UV-Vis spectra of GQD-AZO interfaces (red and blue lines) and GQDs (green line).

(see Fig. 1(c)). For the sake of brevity, we indicate in the following the whole pyrene-AZO molecule as AZO. The GQD-AZO interfaces are optimized and analyzed only for the fold conformation, since it is more stable than the open one (Fig. 3). For both interfaces, the distance between graphene and pyrene nearly ideally corresponds to a π - π stacking distance of 0.33 nm. The formation of the interface significantly changes the dipole moments, increasing its absolute value from 0.89 Debye for *t*AZO to -4.03 Debye for GQD-*t*AZO, and from -0.8 Debye to 1.7 Debye for *c*AZO alone and at the interface, respectively. For both isomers, the occupied (HOMO-1 and HOMO) and virtual (LUMO and LUMO+1) frontier orbitals are degenerated in energy, lying at -5.5 eV and -1.7 eV for the *trans* isomer and -5.5 eV and -1.6 eV for the *cis* isomer, respectively. Therefore, the presence of the GQDs is responsible for the decrease of the energy gap from about 6 eV to 3.8 eV, making the interface a semiconductor (see Fig. S5–S7, ESI†). The localization of the frontier orbitals over the GQDs indicates no contribution arising from the AZO molecules, with the nearest orbital taking part in PET and PHT processes being HOMO-2 (delocalized over graphene and pyrene for both isomers).

The absorption spectrum of the GQD-*t*AZO interface is redshifted with respect to the spectra of *t*AZO in the fold conformation by 100 nm, due to the presence of GQDs. Degenerate electronic transitions are observed. The first two excited states are dark and correspond to the π (GQD) \rightarrow π^* (GQD) transitions, and the first bright peak consists of two strongly allowed transitions at 461 and 458 nm, delocalized over the AZO moiety

and GQDs, respectively (Fig. S8, ESI†). The peak at 445 nm corresponds to the $\pi \rightarrow \pi^*$ transition localized over the GQD fragment. The same shift towards longer wavelengths and degenerate electronic transitions are observed for the GQD-*c*AZO interface. Once again, the first two transitions are dark and localized over the GQD fragment, while the first bright peak at 454 nm consists of two strong transitions, involving both AZO and GQD fragments. As for the GQD-*t*AZO interface, the peak at 443 nm corresponds to the two localized transitions on the GQDs. The presence of the AZO molecule at the interface is also responsible for the redshift of the absorption spectra compared to the bare GQDs, of 10 and 15 nm for *c*AZO and *t*AZO, respectively, also decreasing the shift in absorption between the two isomers compared to the isolated case. In fact, now the GQD-*t*AZO interface exhibits a redshift of 10 nm compared to the GQD-*c*AZO interface, while for the isolated molecule, the *c*AZO isomer exhibits a redshift of more than 100 nm compared to the *trans* one. In addition, the presence of the graphene dots at the GQD-*c*AZO interface is responsible for the suppression of the weak, long wavelength peaks related to the S_1 $n \rightarrow \pi^*$ transition observed for the isolated molecule.

The NTO analysis of the orbitals involved in the photophysics of the two configurations indicates that for these interfaces, AZO acts as the donor (D_{MOL}), whereas the GQDs is the acceptor (A_{GQD}). It is important to notice that since both interfaces exhibit degenerate orbitals, an effective coupling has been calculated as discussed in the Method section. Depending on the number of degenerate states involved in the transition, we considered a combination of multiple coupling values



(as in eqn (4)). For GQD-*t*AZO, the EET process is facilitated by a local AZO excitation at 436 nm, which is followed by an energy transfer to two degenerated GQD local excitations at 461 and 458 nm, respectively. Although the EET takes place between two GQDs' transitions with high oscillator strengths, the unfavorable overlap between the transition densities of these localized states causes a relatively low coupling value of 5.3 meV. For GQD-*c*AZO, the states involved in the EET are local AZO excitation at 462 nm, from which energy is transferred to the two GQDs' local excitations at 455 and 453 nm, leading to a coupling value of 10.4 meV. On the other hand, the PET process is hindered for both interfaces, with the local excitation of the AZO at 436 (462) nm for the *trans* (*cis*) isomer for the electron transfer from AZO to GQDs (visible as two almost degenerated final states at 423/420 nm and 411/409 nm for *trans* and *cis*, respectively), leading to a very weak coupling with a value lower than 4 meV (Table 2). This is a consequence of the weaker overlap between the diabatic wavefunctions of the initial and final states, which in the PET case are, in a first approximation, the LUMOs of AZO and GQD moieties. As one can already see in Fig. 3, where the frontier orbitals of the whole donor:acceptor system are represented for both interfaces, the LUMO is localized only on the GQDs. Ultimately, the dominant process for both interfaces is the PHT, with the local excitations of the GQDs (two degenerated A_{GQD}^* at 433/430 nm), allowing a transfer of a hole from the GQDs to AZO (visible as two almost degenerated final states at 423/422 nm and 411/409 nm for *trans* and *cis*, respectively), leading to holes' coupling values up to 51.2 and 70.4 meV, for the *trans* and *cis* isomers, respectively. The higher value of PHT coupling compared to PET coupling is a consequence of the larger delocalization of the HOMO orbitals involved in the former process compared to the LUMOs (as visible in Fig. 3). Moreover, the different delocalization and overlap between the wavefunctions of the initial and final states are slightly more favorable for GQD-*c*AZO compared to the *trans* isomer, although the difference is too small to be visually appreciated.

The PHT reorganization energy for GQD-*t*AZO computed using the forward and backward reorganization energy scheme (eqn (S1)–(S3), ESI[†]), where the AZO molecule is considered as a donor and GQDs as an acceptor, has values of 0.71 and 0.64 eV, respectively (Table 2). For the EET process, the values are similar: 0.19 eV and 0.20 eV for *t*AZO and *c*AZO isomers, respectively. A different scenario is observed for the PET process, for which the presence of the GQDs decreased the

Table 2 Total reorganization energy, electronic coupling, driving force and Marcus rates for the three processes considered in this study, for the GQD–AZO interfaces

	Process	λ [eV]	Coupling [eV]	ΔG [eV]	k [s^{-1}]
GQD- <i>t</i> AZO	EET	0.19	0.005	-0.146	1.0×10^{12}
	PET	0.33	0.001	0.095	3.6×10^7
	PHT	0.71	0.051	0.068	1.3×10^{10}
GQD- <i>c</i> AZO	EET	0.20	0.010	0.043	2.3×10^{11}
	PET	0.11	0.004	0.337	2.8×10^4
	PHT	0.64	0.070	0.142	9.4×10^9

λ to 0.33 eV for GQD-*t*AZO and to 0.11 eV for GQD-*c*AZO. Overall, the presence of GQDs has a strong influence on the reorganization energies in the GQD–AZO interfaces, significantly decreasing their magnitude to similar values. This can be explained considering the nature of the GQDs, since they can easily accommodate one additional electron/hole, with little or negligible distortions in the interfacial structure. Thus, since it is not obvious which photoinduced process will dominate due to the lack of a clear trend in the different components (reorganization energy, electronic coupling and driving force), it is crucial to compute the Marcus rates, to quantify and disentangle the different contributions.

From the analysis of the Marcus rates, we observe that the trend characterizing the rates for the different processes follows: $k_{\text{EET}} > k_{\text{PHT}} > k_{\text{PET}}$ for the GQD-*t*AZO interface, in which the fastest process after photoexcitation is EET, with a k_{EET} value of $1 \times 10^{12} \text{ s}^{-1}$. PHT might also take place, although with a smaller rate of $1.3 \times 10^{10} \text{ s}^{-1}$. On the other hand, PET is rather slow (Table 2). Opposite to GQD-*t*AZO, the GQD-*c*AZO interface presents an overall lower rate for all the three processes, following the $k_{\text{EET}} > k_{\text{PHT}} \gg k_{\text{PET}}$ trend. PHT could become favorable also for the GQD-*c*AZO interface, as its rate of $9.4 \times 10^9 \text{ s}^{-1}$ is high, albeit smaller compared to the EET rate ($2.3 \times 10^{11} \text{ s}^{-1}$). It is also worth noting the effect of the isomer on the rate values. In fact, as already discussed for the intramolecular processes for the isolated molecules, also when intermolecular photoinduced transfer at the interface is considered, the rates for the *trans* isomer are always higher in magnitude compared to the *cis* isomer. At first glance, this can appear counterintuitive, because the couplings are smaller for GQD-*t*AZO compared to GQD-*c*AZO, and the reorganization energies have rather similar values. We can thus infer that this effect is mainly due to the driving force, which presents large variations for the different processes for both isomers. This confirms that *t*AZO is the active state for the interface, presenting higher rates for all the processes, while *c*AZO is the inactive state.

GQD–AZO – MD simulations

In order to assess the effects of structural disorder and temperature on the transfer rates, we consider here the analysis of an assembly of AZO molecules, in which we explicitly consider the thermal effects using MD simulations.

The first observation from MD simulations is that the majority of the AZO molecules prefer the fold conformation (88% for *trans* and 83% for *cis*) over the open one and they tend to aggregate in one place of the surface (Fig. S9–S11, ESI[†]). The presence of thermal energy results in the distortion of the AZO molecule geometry, which in turn tends to decrease the degeneracy of the occupied and virtual frontier molecular orbitals. The GQD-*t*AZO interface shows a double distribution for all four frontier orbitals, leading to an energy gap of 3.8 eV (Fig. S12 and S13, ESI[†]).

The convoluted absorption spectra for all the extracted frames from the MD simulations are reported in Fig. 4. The first absorption peak for the GQD-*t*AZO interface is slightly



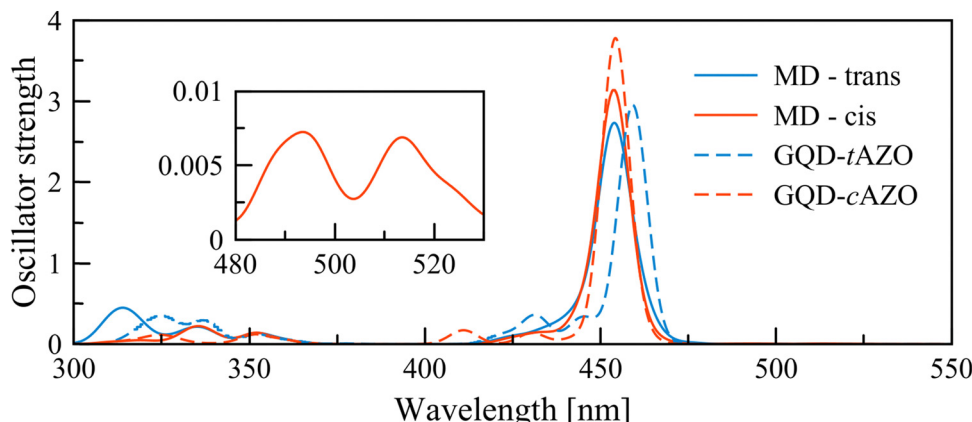


Fig. 4 Convoluted UV-Vis spectrum recorded using 50 frames extracted from the MD simulations and the absorption spectra of the optimized interfaces (replotted from Fig. 3). The inset shows a magnification of the long wavelength peaks for the cAZO interface.

blue shifted by 8 nm compared to the optimized interface. This peak at 459 nm corresponds to the $\pi(\text{GQD}) \rightarrow \pi^*(\text{GQD})$ transition. It is worth noting that the first four excited states are now dark. The shoulder at 427 nm consists of series of three $\pi(\text{GQD} + \text{AZO}) \rightarrow \pi^*(\text{GQD})$ transitions. Opposite to *trans*, the absorption spectrum for the GQD-cAZO interface is not shifted compared to the model interface, however two additional weakly allowed peaks at 515 and 494 nm appear due to the molecular distortion induced by the thermal energy. The peak at longer wavelength corresponds to the first excited state and the second relates to the second excited state and both have a $\pi(\text{GQD}) \rightarrow \pi^*(\text{GQD})$ character. The high-intensity peak at 454 nm is related to a transition localized over the GQD fragment of the interface. A mixture of contributions from different excited states (S_8 (25%), S_9 (37%), S_{10} (16%) and S_{11} (22%)) is found for the peak at 430 nm and correspond to $\pi(\text{GQD} + \text{AZO}) \rightarrow \pi^*(\text{GQD})$ transitions. Overall, the distortion of the molecular geometry of the interface due to thermal energy marginally affects the GQD-tAZO interface while it has a much stronger effect on the GQD-cAZO interface, resulting in two weakly allowed transitions at longer wavelengths, which might result in an enhanced the photoinduced transfer rate.

To quantify and assess how the thermal effects affect EET, PET and PHT processes at the GQD-AZO interfaces, the couplings are evaluated for all the frames extracted from the MD simulations. Once again the GQD is found to be an acceptor (A_{GQD}) and AZO acts as a donor site (D_{MOL}). For all the transfer processes, we observed that the same states already found in the optimized interfaces take part in the transitions. Surprisingly, the EET coupling between the first allowed degenerated GQD (A_{GQD}^*) local excitations and AZO (D_{MOL}^*) local excitation for the GQD-tAZO interface increases four times in value compared to the optimized interface, with a final value of 18 meV. Similarly, the EET coupling between A_{GQD}^* and D_{MOL}^* for the GQD-cAZO interface exhibits a value of 57 meV, which corresponds to an almost six time increase in value compared to the optimized interface (Table S3, ESI[†]). We believe that this is due to the fact that thermal effects and related structural

Table 3 Total reorganization energy, electronic coupling, driving force and Marcus rates for the three processes considered in this study, for the GQD-AZO interface, obtained using MD simulations. Standard deviations of the different quantities are reported in brackets

Process	λ [eV]	Coupling [eV]	ΔG [eV]	k [s ⁻¹]
GQD-tAZO	EET	0.05 (0.01)	0.018 (0.02)	-0.152 (0.06) 2.8×10^{12}
	PET	0.08 (0.03)	0.002 (0.002)	0.119 (0.02) 1.3×10^9
	PHT	0.19 (0.79)	0.038 (0.03)	0.070 (0.03) 1.7×10^{12}
GQD-cAZO	EET	0.06 (0.03)	0.057 (0.03)	0.297 (0.12) 1.4×10^5
	PET	0.10 (0.04)	0.004 (0.003)	0.480 (0.12) 5.6×10^{-3}
	PHT	0.23 (0.94)	0.024 (0.01)	0.055 (0.02) 6.5×10^{11}

fluctuations increase the conformational motion allowing for a broader sampling of the potential energy surface for the selected process, resulting in the discovery of different local minima which can enhance the electronic coupling. A similar effect has been reported in the literature for biosystems,⁵⁵ as well as for other types of couplings in organic crystals and semiconducting polymers depending on the transport regime.^{56,57} On the other hand, for the PET process, the coupling values have not changed compared to the optimized interfaces (Table 3). A strong decrease in coupling is observed for the PHT process, with 35% lower value for the GQD-tAZO interface compared to the optimized interface, leading to a final hole coupling value of 38 meV, and of 75% decrease for the GQD-cAZO interface, with the final value of 24 meV. The fact that we observe different degrees of geometrical distortions induced by the presence of thermal energy on the two interfaces helps us in rationalizing the coupling values for PHT. For the GQD-tAZO interface, we observe that almost all the molecules are in the fold conformation, with some minor amount in the open one (12% of the total). On the other hand, a higher percentage of molecules are found in their open conformation for the GQD-cAZO interface (17%) with a non-negligible number of molecules with geometries that lie somewhere in between open and fold conformations (Fig. S8 and S9 in the ESI[†]). Since we determined that the couplings for open conformations are generally lower compared to the fold ones (see Table S2, ESI[†]), we can rationalize the large drop in



coupling observed for GQD-*c*AZO and the relatively small decrease in the value for GQD-*t*AZO compared to the model interfaces.

The reorganization energy obtained using eqn (1) and (2) calculated over 35 frames for the PET process has values of 0.08 eV and 0.10 eV for GQD-*t*AZO and GQD-*c*AZO, respectively. Considerably higher values of 0.19 eV and 0.23 eV are found for PHT, suggesting that a higher barrier must be crossed between the initial and final states to facilitate PHT than PET. Finally, a similar value of 0.05 eV has been computed for EET for both interfaces (Table 3). The driving force analysis shows a different trend for the two interfaces compared to the reorganization energy, with EET favored for the GQD-*t*AZO interface and PHT favored for the GQD-*c*AZO interface, suggesting that additional electronic factors rather than the pure geometrical distortions are responsible for the different computed Marcus rates.

The inclusion of thermal effects has interesting repercussions on the Marcus rates of the two interfaces. While this effect at the GQD-*t*AZO interface is negligible for the EET process, accounting for structural disorder induced by thermal fluctuations strongly enhances the PHT rate up to $1.7 \times 10^{12} \text{ s}^{-1}$, resulting in a strong competition between PHT and EET to dominate the photophysics of the system. Moreover, the PET rate is also enhanced, making the overall trend ($k_{\text{EET}} \sim k_{\text{PHT}} > k_{\text{PET}}$) different from the case at 0 K. If we compare the optimized GQD-*t*AZO interface and the current one in which thermal effects are considered, we observe that especially for PHT (and to a lesser extent also PET), the reorganization energy is strongly decreased, while driving force and coupling remain similar. This suggests that the reorganization of the *t*AZO molecules on the surface allows for a beneficial relaxation of this conformer, resulting in an increased PHT rate. Structural disorder induced by thermal energy has a strong impact also at the GQD-*c*AZO interface, for all processes. In fact, now the rates follow the $k_{\text{PHT}} \gg k_{\text{EET}} \gg k_{\text{PET}}$ trend, with a value of $6.5 \times 10^{11} \text{ s}^{-1}$ for PHT, much higher than that for EET ($1.4 \times 10^5 \text{ s}^{-1}$). Although the coupling for EET is strongly enhanced compared to the optimized interface and the PHT is decreased, the driving force for PHT is strongly decreased, while it is almost 7 times higher for EET. In addition, a stronger decrease in reorganization energy of 36% for PHT compared to a 30% decrease for EET ensures that the dominant process for the GQD-*c*AZO interface is PHT. Despite these large variations in values, all processes are faster at the GQD-*t*AZO interface compared to those at the GQD-*c*AZO interface. This confirms that *t*AZO is an active isomer for photoinduced processes, while *c*AZO can be considered to be less active.

Conclusions

A comprehensive study of photoinduced charge and energy transfer in azobenzene derivatives as well as their interfaces with graphene quantum dots is reported. By means of DFT calculations as well as MD simulations, the effects of conformational changes for the two isomers on the magnitude of PET, PHT and EET processes are investigated through the

calculation of reorganization energies, intra- and inter-molecular couplings, driving force and eventually Marcus transfer rates. For isolated AZO, the fold conformations are more stable compared to the open one and these are predominately present when performing MD simulations for the GQD-AZO interfaces as well. The comparison between the UV-Vis spectra of *cis* and *trans* isomers revealed that the azobenzene derivatives presented in this study are capable of undergoing photoisomerization, thereby presenting a unique opportunity to create an on/off switch by connecting them with GQDs.

We found that EET dominates the intramolecular process for both isomers of the isolated molecules, as well as the intermolecular process at the GQD-AZO interfaces, although for the latter system, PHT can be a competitive pathway. This effect is mainly due to the presence of the GQDs, which strongly decrease the reorganization energy at the interfaces of both isomers and enhance their hole coupling values. On the other hand, PET is strongly suppressed, due to a negligible electronic coupling (lower than 0.005 eV) between the moieties of the interfaces, as well as the sizable activation barrier characterizing the electron transfer in this case.

A striking decrease of reorganization energies when thermal effects are included in the model causes the enhancement of PET and PHT for both interfaces. This is a direct consequence of the presence of the GQDs that can accommodate an additional electron or hole without significant deformation of the interfacial geometrical structure. On the other hand, thermal effects strongly decrease the EET rates at the GQD-*c*AZO interface, favouring PHT instead, which becomes the fastest process (occurring on a ps time-scale). Crucially, the formation of the interface with the *trans* isomer of the AZO molecule ensures higher rates for all the processes compared to the *cis* isomer. Thus, *t*AZO can be considered as the active form of interest for photoinduced applications in comparison with the corresponding *c*AZO isomer. We would like to remark here that the presented results might depend on the specific system. Different functionalization of the AZO molecule might lead to different morphological and photophysical properties (such as energy level alignment) which need to be studied for each case. Moreover, this study highlights the importance of considering thermal fluctuations and structural dynamics as they play an important role in the determination of the kinetics of the different processes. Considering only the position of the energy levels as descriptors for the photophysical process taking place at interfaces might present an incomplete picture. This study paves the way to assess the photoisomerization transfer mechanism of low-dimensional material interfaces by means of a material design approach for selected applications in which one of the three processes is desired.

Author contributions

The manuscript was written through the contributions of all authors. All of the authors discussed the results and participated in writing the final version of the manuscript and have



given approval to the final version of the manuscript. S. O. proposed the research direction, supervised the project conceptually and experimentally and helped with the interpretation of the results. M. K. participated in the design of the research and performed the calculations, including all data collection and further analysis and interpretation. S. O. and S. G. supervised the preparation of the manuscript, including the interpretation of the results and participated in finalizing the paper.

Conflicts of interest

There are no conflicts to declare.

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