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Chemisorption of Atomically Precise 42-Carbon Graphene Quantum Dots on Metal Oxide Films Greatly Accelerates Interfacial Electron Transfer

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ABSTRACT: Graphene quantum dots (GQDs) are emerging as environmentally friendly, low-cost, and highly tunable building blocks in solar energy conversion architectures, such as solar (fuel) cells. Specifically, GQDs constitute a promising alternative for organometallic dyes in sensitized oxide systems. Current sensitized solar cells employing atomically precise GQDs are based on physisorbed sensitizers, with typically limited efficiencies. Chemisorption has been pointed out as a solution to boost photoconversion efficiencies, by allowing improved control over sensitizer surface coverage and sensitizer-oxide coupling strength. Here, employing time-resolved THz spectroscopy, we demonstrate that chemisorption of atomically precise C42-GQDs (hexa-peri-hexabenzocoronene derivatives consisting of 42 sp² carbon atoms) onto mesoporous metal oxides, enabled by their functionalization with a carboxylate group, enhances electron transfer (ET) rates by almost 2 orders of magnitude when compared with physisorbed sensitizers. Density functional theory (DFT) calculations, absorption spectroscopy, and valence band X-ray photoelectron spectroscopy reveal that the enhanced ET rates can be traced to stronger donor–acceptor coupling strength enabled by chemisorption.

Graphene quantum dots (GQDs) are nanosized graphene fragments, which have nonzero, size-dependent bandgaps due to quantum confinement effects. Moreover, GQDs are metal-free and hence potentially low-cost and environmentally friendly. These features have motivated researchers to apply GQDs in solar energy conversion schemes (e.g., solar cells†–‡ and photocatalytic devices§,∥). GQDs are typically prepared by hydrothermal treatment of graphene or small molecules; although certain control of GQD size has been achieved following this synthesis protocol,‡ samples are generally defined by broad absorption features induced by inhomogeneous broadening (i.e., samples do not consist on a narrow distribution of chemical structures), an aspect that is detrimental for optoelectronic applications.

Alternatively, large polycyclic aromatic hydrocarbons (PAHs) have been synthesized in the field of organic chemistry over the last decades, with hexa-peri-hexabenzocoronene (HBC), consisting of 42 sp² carbon atoms, being a representative example.¹⁰,¹¹ Recently, Yan et al. reported the synthesis of large PAHs consisting of 132, 168, and 170 sp² carbon atoms and referred to them as colloidal GQDs.¹²,¹³ Such PAHs, sometimes also called nanographenes, can indeed serve as atomically precise zero dimensional GQDs, owing to their well-defined size- and shape-dependent optoelectronic properties as predicted by theoretical predictions.¹⁴–¹⁶ Yan et al. also reported the use of such well-defined GQDs as absorbers in a sensitized solar cell geometry.¹⁷ However, these initial solar cell devices revealed low photoconversion efficiencies, mainly linked with low short-circuit currents. The poor photocurrent produced in the cells was tentatively correlated with the low affinity of the employed GQDs sensitizers, which were physisorbed onto the mesoporous oxide surface. In a follow up report from the same group, theoretical studies have shown that chemisorption, instead of physisorption, of sensitizers should favor donor–acceptor coupling and hence boost electron transfer at GQD/oxide interfaces.¹⁸ Furthermore, theoretical studies have shown that chemisorption, instead of physisorption, of sensitizers should favor donor–acceptor coupling and hence boost electron transfer at GQD/oxide interfaces.¹⁸,¹⁹,²⁰ Although all these works have suggested that chemisorption of GQDs...
onto metal oxides might improve photoconversion efficiencies in sensitized systems, there is at present no experimental evidence to support that claim. Here, we quantify interfacial electron transfer (ET) rates for atomically precise GQDC42 (HBC with 42 sp² carbon atoms) chemisorbed and physisorbed on mesoporous SnO₂ by optical pump-terahertz probe (OPTP) spectroscopy. We demonstrate that sensitizer chemisorption onto the oxide electrode substantially improves ET rates induced by strong overlap (hybridization) between donor and acceptor wave functions, which is triggered by the functionalization of the GQDC42 by a carboxylate group. This claim is directly evident from the OPTP data and is further supported by absorption spectroscopy, valence band X-ray photoelectron spectroscopy (VBXPS) and density functional theory (DFT) calculations.

The molecular structures of the two GQDC42 samples used in this study are shown in Figure 1. For clarity, we name our samples as GQDC42 and GQDC42-PhCOOH for the sensitizer without and with a phenyl carboxylic acid functional group, respectively. The GQDC42 sample was prepared following a protocol described previously and the synthesis of GQDC42-PhCOOH is described in the Supporting Information (SI). In brief, GQDC42-PhCOOH was synthesized starting from a hexaphenylbenzene derivative bearing bromo- and iodo-groups through a Suzuki-coupling reaction to selectively introduce a phenyl ester moiety and subsequent cyclodehydrogenation reaction to convert the hexaphenylbenzene core into HBC. Finally, the ester group was hydrolyzed under basic conditions to the desired carboxylic acid anchoring group. The synthesized samples were diluted in toluene, and the obtained suspensions were employed for sensitizing the mesoporous oxide films (see the SI for sample preparation details). The functionalized films were subsequently characterized by OPTP spectroscopy; the measurements were performed under nitrogen conditions to prevent any potential sensitizer photo-oxidation.

OPTP spectroscopy is a powerful tool to investigate ultrafast interfacial dynamics for dye- and QD-sensitized oxide systems. As the employed THz probe (∼1.5 THz bandwidth) is primarily sensitive to free carrier motion (i.e., photoconductivity), an OPTP measurement in a sensitized oxide neatly probes the emergence of photoconductivity in the oxide electrode after selective excitation of the sensitizer. As such, it resolves, in time, the arrival of electrons from the sensitizer’s populated molecular orbitals (e.g., LUMO) into the oxide’s conduction band (CB). Figure 2a shows normalized OPTP dynamics for GQDC42 and GQDC42-PhCOOH sensitizing SnO₂ mesoporous films (blue diamonds and red circles respectively; 400 nm pump excitation, 0.6 mJ/cm²); these dynamics were collected in the linear single-exciton regime (see Figure S1). The lack of response under 400 nm pump excitation of a bare SnO₂ film is also presented in Figure 2a (black triangles). As evident from Figure 2a, the sensitization of mesoporous SnO₂ by the sensitizer functionalized with a phenyl carboxylate group (GQDC42-PhCOOH, Figure 1), which is expected to

Figure 1. Chemical structures of the graphene quantum dots, GQDC42 and GQDC42-PhCOOH, analyzed in this study; they differ on the presence of a carboxylate functional group which is expected to serve as a covalent link to the oxide electrode.

Figure 2. (a) OPTP dynamics of GQDC42 (open blue diamonds) and GQDC42-PhCOOH (open red circles) sensitizing SnO₂ films (400 nm pump excitation, 0.6 mJ cm⁻²). Traces are normalized to the plateau of the biexponential fits (solid red lines). Black open triangles are OPTP dynamics for a bare SnO₂ film. (b) Frequency-resolved complex photoconductivity for both sensitized systems (1 ns after photoexcitation); solid and dashed lines correspond to Drude–Smith fits for the real and imaginary components of the frequency dependent conductivity.
chemisorbed at the oxide surface, results in faster ET rates when compared with the sensitizer lacking functionalization (GQD-C42 in Figure 1), which is expected to physisorb at the oxide surface. Both traces can be well described phenomenologically by a biphasic exponential model (solid red lines in Figure 2), providing time constants of $t_1 = 12 \pm 1$ ps and $t_2 = 2025 \pm 450$ ps for GQD-C42 sensitized SnO$_2$ films; and $t_1 = 0.2 \pm 0.1$ ps and $t_2 = 51 \pm 3$ ps for GQD-C42-PhCOOH sensitized SnO$_2$ films. Biphasic ET dynamics are a common observable for dye- and QD- sensitized oxide systems that have been generally explained in terms of two distinct transfer channels toward the oxide electrode. They might be linked with "hot" and "cold" ET channels or, alternatively, to “cold” ET channels induced by two donor–acceptor interfacial conformations (i.e., providing distinct donor–acceptor energetics). Biphasic dynamics have been also rationalized by considering effects induced by molecules loosely attached (physisorbed) to the surface or those present in form of the aggregates. Even though it is difficult to rule out any of these scenarios from the current data, the fact that both ET components become faster when QDs are chemisorbed onto the oxide matrix support the view that two ET channels define our interfacial dynamics. A deeper analysis of the nature of the biphasic signals is underway and will be reported elsewhere.

To validate the conclusion of faster ET rates for the chemisorbed sensitizers inferred from the OPTP data, we analyze the frequency-resolved complex photoconductivity in the samples. This approach can tell whether OPTP dynamics indeed refer uniquely to electrons populating the oxide conduction band. Figure 2b presents the real and imaginary (closed and open symbols) components of the complex conductivity for GQD-C42 (blue) and GQD-C42-PhCOOH (red) sensitized oxide films (1 ns after excitation, 1.5 THz bandwidth). As evident from Figure 2b, the complex photoconductivity spectra for chemisorbed and physisorbed dots onto the oxide overlap quite well, indicating that the nature of the monitored photoconductivity for both samples is identical, as expected for electrons in SnO$_2$. The resolved complex spectra can be well described by the phenomenological Drude–Smith (DS) model:

$$\sigma(\omega) = \frac{n_0^2 \varepsilon_0 \varepsilon_r}{1 - i \omega \tau_\sigma} \left(1 + \frac{c}{1 - i \omega \tau_c}\right)$$

where $n_0$, $\varepsilon_0$, $\varepsilon_r$, $\tau_\sigma$, and $c$ refer to the plasma frequency, vacuum permittivity, scattering time and localization factor, respectively. Within the DS model, the measurable frequency resolved photoconductivity can be attributed to free carriers experiencing preferential backscattering at the boundaries of the nanocrystalline mesoporous oxide electrode. The strength of the localization is parametrized by $c$, which ranges between 0 (free carrier, Drude response) and 1 (representing full localization of charges). Best fit to the data using eq 1 for both samples provides the same fitting parameters within error; a scattering rate of $\tau_\sigma = 46 \pm 2$ fs and localization factor of $c = -0.69 \pm 0.1$. These results are in good agreement with previous values inferred for nanostructured SnO$_2$ films and, hence, demonstrate that the monitored signal in Figure 2a refers uniquely to electrons populating the oxide conduction band. These results support our conclusion that the changes in the monitored interfacial ET rates can be traced uniquely to the presence of the phenyl carboxylate group functionalizing the GQD-C42-PhCOOH, allowing for chemisorption. Further more, these results reveal that electron transport within the analyzed SnO$_2$ electrode is unaffected by the type of functionalization (chemisorption or physisorption).

To check the generality of our observation that ET is accelerated for chemisorbed sensitizers, we also measured interfacial ET dynamics for both sensitizers onto mesoporous ZnO and TiO$_2$ films (Figure S3). For ZnO samples, we observed as well faster ET rates for the chemisorbed sensitizers, analogous to the case of SnO$_2$ based samples (Figure 2). For titania, ET occurs faster than our experimental sub-picosecond time resolution for both sensitizers, so we are unable to determine changes in ET rates for this set of samples. The overall slower ET rates observed for ZnO and SnO$_2$ electrodes when compared to titania, generally explained in terms of larger density of states for the latter improving donor–acceptor coupling strength, agree qualitatively with those reported for organometallic dye sensitizers onto the same electrodes. Overall, the results support the view that the presence of a phenyl carboxylate group facilitates electron transfer toward the electrode independently of its nature.

To rationalize our finding of faster ET for chemisorbed GQD-C42, we investigate the nature of the interaction between sensitizer donor and oxide acceptor. Figure 3 presents the relative energy level alignments. For the SnO$_2$ electrode, the energy positions are obtained from ultraviolet photoelectron spectroscopy (UPS, see Figure S5) and absorbance. For the two molecules, the highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO) were obtained from gas-phase DFT calculations. When the interaction of the metal oxide with the molecules is considered, the frontier orbital distributions of HOMO and LUMO for both sensitizers are also presented. The Fermi level for the tin oxide sample is within the oxide CB, indicating strong n-type character.
respectively) along with the bare oxide solution (dashed lines). The spectrum of bare SnO$_2$ GQDC42-PhCOOH and GQDC42 sensitizers (red and blue line (solid black line) and oxide electrodes functionalized by GQDC42 remains essentially identical upon sensitization, a PhCOOHs, respectively. Whereas the peak absorbance for GQDC42 sensitized and the bare tin oxide electrodes reveal respectively). As evident from the plot, the physisorbed GQDC42 orbitals and the oxide conduction band, we analyzed spectroscopy (VBXPS) data for a bare SnO$_2$ mesoporous electrode (dotted and solid red lines in Figure 4a). A similar blue shift in the absorbance after sensitization has been previously reported for dye-sensitized electrode systems.

To reveal a stronger interaction between chemisorbed GQD$_{C42}$ orbitals and the oxide conduction band, we analyzed the samples by optical absorption and valence band X-ray photoelectron spectroscopy. In Figure 4a, we present the UV−vis absorbance spectrum of both sensitized systems (solid red and blue lines for chemisorbed and physisorbed GQD$_{C42}$, respectively); the absorption profiles peak at 3.45 and 3.39 eV for GQD$_{C42}$ and GQD$_{C42}$-PhCOOHs, respectively. Whereas the peak absorbance for GQD$_{C42}$ remains essentially identical upon sensitization, a clear blue shift is resolved for the peak absorbance of the GQD$_{C42}$-PhCOOH sensitizers after binding to the oxide electrode (dotted and solid red lines in Figure 4a). A similar blue shift in the absorbance after sensitization has been previously reported for dye-sensitized films and explained in terms of the strong coupling induced by deprotonation of the dyes$^{42−45}$ induced by chemisorption; in good agreement with our expectations.

In Figure 4b, we present valence band X-ray photoelectron spectroscopy (VBXPS) data for a bare SnO$_2$ mesoporous film (solid black line) and oxide electrodes functionalized by GQD$_{C42}$-PhCOOH and GQD$_{C42}$ sensitizers (red and blue line respectively). As evident from the plot, the physisorbed GQD$_{C42}$ sensitized and the bare tin oxide electrodes reveal identical electron binding energies. However, the electron binding energy is reduced by approximately 2 eV upon chemisorption of GQD$_{C42}$-PhCOOHs, indicating significant/efficient modification/decoration of the surface of the oxide, which was absent upon the physisorption (the results were reproducible within $\sim$50 meV for samples produced in three different batches). The observed change in electron binding energy can be explained by chemical modification at the oxide interface (e.g., deprotonation of COOH groups able to modify the Fermi energy at the oxide) and/or by dipolar effects (modifying vacuum levels induced by the presence of a surface electric field induced by sensitization).$^{46}$ In any case, it reveals a strong donor−acceptor interaction for the chemisorbed GQD$_{C42}$, in qualitative agreement with absorbance spectra shown in Figure 4a and the expected closer proximity of frontier orbitals inferred by DFT calculation (Figure 3). All these results support qualitatively the experimentally resolved effect on interfacial OPTP dynamics, i.e. the main conclusion of this work: chemisorption of GQD$_{C42}$ enabled by their functionalization with a carboxylate group, substantially enhances ET rates.

In summary, we investigated the rates of electron transfer from physisorbed and chemisorbed graphene quantum dot (GQD) sensitizers to metal oxides, using hexa-peri-hexabenzocoronene derivatives as atomically precise GQDs. An increase of ET rates as large as 2 orders of magnitude is observed for chemisorbed sensitizers when compared with physisorbed ones. Accelerated electron transfer is correlated with enhanced donor−acceptor coupling (i.e., wave function hybridization). These results demonstrate that functionalization of GQDs with anchoring head groups represents a potential path for improved photoconversion efficiencies in carbon-based sensitizer/oxide electrodes employed in solar energy conversion schemes.

**ASSOCIATED CONTENT**

* Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcllett.9b00399.

Materials and methods; OPTP dynamics of GQDs sensitized SnO$_2$ in linear single-exciton regime; measurement artifact in frequency-resolved photoconductivity; OPTP dynamics of GQDs Sensitized ZnO/TiO$_2$; frequency-resolved photoconductivity of GQDs sensitized ZnO/TiO$_2$; UPS and absorption spectrum of mesoporous SnO$_2$ Film (PDF)

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