How mobile are dye adsorbates and acetonitrile molecules on the surface of TiO$_2$ nanoparticles? A quasi-elastic neutron scattering study

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Motions of molecules adsorbed to surfaces may control the rate of charge transport within monolayers in systems such as dye sensitized solar cells. We used quasi-elastic neutron scattering (QENS) to evaluate the possible dynamics of two small dye moieties, iisonicotinic acid (INA) and bis-iisonicotinic acid (BINA), attached to TiO$_2$ nanoparticles via carboxylate groups. The scattering data indicate that moieties are immobile and do not rotate around the anchoring groups on timescales between around 10 ps and a few ns (corresponding to the instrumental range). This gives an upper limit for the rate at which conformational fluctuations can assist charge transport between anchored molecules. Our observations suggest that if the conformation of larger dye molecules varies with time, it does so on longer timescales and/or in parts of the molecule which are not directly connected to the anchoring group. The QENS measurements also indicate that several layers of acetonitrile solvent molecules are immobilized at the interface with the TiO$_2$ on the measurement time scale, in reasonable agreement with recent classical molecular dynamics results.

The properties of many materials can be tuned by exposing them to carefully chosen chemical functional groups. In particular, the functionalization of surfaces has extensive applications ranging from biology$^{1-4}$ to molecular electronics$^{5-7}$. For example, sensitizing semiconductor nanocrystals with dye molecules has led to the development of new types of solar fuel photoelectrodes$^{8,9}$ and dye sensitized solar cells (DSSC)$^{9,10}$. In these systems, the light absorbing properties of the dyes are combined with the electronic conductivity of the metal oxide substrate (often TiO$_2$). However, the organic/inorganic interface created is often complex, making its properties difficult to predict. For example, the standard role for a dye molecule in a DSSC is to act as a centre for light absorption, then to inject the photoexcited electron into the TiO$_2$ substrate, and to finally accept an electron from the surrounding electrolyte into the hole left on the dye following injection. However, in addition to these functions, lateral hole diffusion can occur in the dye monolayer by charge transfer between molecules anchored to adjacent sites on the TiO$_2$ surface$^{11-20}$. These interesting phenomena have been shown to improve the performance of solid state DSSCs$^{11,12}$, and may play a role in recombination within liquid DSSCs and devices for water splitting$^{21}$. Indeed, DSSCs can work without a separate hole transporting phase, with holes being collected solely via the dye monolayer$^{18}$. Thus, identifying the factors controlling the kinetics of this process is of interest for the design of dye sensitized electronic devices.

We have recently shown that the kinetics of lateral hole diffusion appear to be influenced by the nature of the surrounding medium$^{22,23}$, intermolecular interactions$^{24}$, and – of particular relevance for this study – the

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structural rearrangement of anchored molecules at the surface. Structural and vibrational dynamics of sensitized metal oxide surfaces have been investigated both theoretically with molecular dynamics and experimentally with X-ray, NMR, IR and photoelectron spectroscopy. However, these techniques typically only probe the sub-ps timescale. To be relevant to charge transport, we are interested in the timescale on which a charge transfer event occurs, typically $10^{-8}$ s or longer in these systems (dispersive transport is likely on a microscopic scale in these systems). This is much harder to characterize, due to both the static and dynamic disorder of the organic film on the surface.

In this study we address the question: does the conformational rearrangement of dyes anchored to TiO$_2$ occur on timescales comparable to hole hopping events? To investigate this problem, we used Quasi-Elastic Neutron Scattering (QENS) to probe sensitized TiO$_2$ nanoparticles. The wavelength of the ‘thermal’ neutrons used and range of scattering angles allow us to investigate structures with length scales from 0.5 to 5.5 Å. We selected a back-scattering spectrometer that can detect changes in the scattered neutron energy corresponding to timescales between tens of ps and a few ns, spanning the relevant window.

As a model experimental system, we examined TiO$_2$ nanoparticles (21 nm diameter) sensitized with a complete monolayer of one of two different dye moieties. These were isonicotinic acid (INA) which is a carboxylated pyridine, and the bipyridyl equivalent (BINA), which have one and two anchoring groups respectively (see Fig. 1). Since INA and BINA form the anchoring moieties in more complex monodentate dyes, such as indolene, and bidendate dyes, such as those based on ruthenium or cobalt. In selected experiments, the sensitized particles were immersed in acetonitrile since acetonitrile is commonly used as the solvent in DSSC electrolytes. The neutron scattering signals from the sensitized TiO$_2$ samples, measured either as a dried powder, or combined with acetonitrile to make a paste, were compared to the scattering signal from unsensitized TiO$_2$ nanoparticulate powder or paste. Since a significant proportion of the signal arises from the incoherent scattering contribution of hydrogen nuclei, analysis of the data should give information about the motion of individual molecules containing hydrogen such as the adsorbates.

Possible kinetics of the rotation of a dye’s body around the axis parallel to the TiO$_2$ surface are illustrated in Fig. 1d. Due to the steric hindrance of the surrounding molecules, we expect that the adsorbates do not lie flat on the surface but are likely to adopt tilted configurations (blue lines in Fig. 1e) and may ‘flap’ between these tilted positions at some rate, $\tau^{-1}$. The dynamics of solvent itself could also influence the properties of the dye sensitized substrates.

In the first section, we present experimental results that indicate that acetonitrile solvent molecules close to the TiO$_2$ surface are likely to be immobilized within the measurement time window. These observations also serve as a partial control for the second section of the results where our measurements indicate that both INA and BINA anchored to the TiO$_2$ nanoparticle surfaces are immobile within the time range of the measurement.
Results

Influence of the TiO$_2$ on solvent (acetonitrile) diffusion. In this section we compare the scattering spectra obtained from pure hydrogenated acetonitrile (hACN) with that obtained from a paste of approximately 1:1 by weight of hACN and TiO$_2$ nanoparticles. This allows us to test the possibility that the nanoparticles influence the diffusion of the ACN molecules.

The QENS spectra of pure acetonitrile could be well described by a self-diffusion model incorporating the translational motion of the ACN molecules \cite{40,41}. Figure 2a shows an example of the full width half maximum (FWHM) of the Lorentzian function fitted to the broadening of the scattering peak attributed to translational motion of the hACN molecules ($\Gamma_{\text{trans}}$) plotted against the square of momentum magnitude transferred, $Q^2$, at 250 K (an example fit is shown in Figure S1). For sufficiently low momentum transfer, the data are well described by a Fickian model of self-diffusion which predicts a linear dependence between the FWHM and $Q^2$ in the limit of low momentum transfer:\cite{40,42}

$$\lim_{Q^2\to0} \Gamma_{\text{trans}}(Q) = \frac{\hbar}{2} D_{\text{trans}}$$

where $\hbar$ is Planck’s constant divided by $2\pi$. The following self-diffusion coefficients are obtained at 250 K, 270 K and 290 K: $D_{\text{trans}} = 2.41 \pm 0.02 \times 10^{-5}$ cm$^2$/s, $3.12 \pm 0.05 \times 10^{-5}$ cm$^2$/s and $3.95 \pm 0.07 \times 10^{-5}$ cm$^2$/s (see also Table S1 for the time constants corresponding to the plateaus at high $Q$ values which can be interpreted in terms of expected methyl group rotation). The resulting Arrhenius activation energy of these values is 0.08 eV, enabling us to confirm by extrapolation that the diffusion coefficients we observe are consistent with those derived previously for acetonitrile at 298 K with QENS ($D_{298K} = 4.2 \pm 0.2 \times 10^{-5}$ cm$^2$/s)\cite{40,43} and NMR ($D_{298K} = 4.04 \times 10^{-5}$ cm$^2$/s)\cite{44}.

While the QENS spectra of the hACN + TiO$_2$ nanoparticle paste indicated similar broadening to that of the pure hACN, the hACN diffusion coefficient in the paste was reduced by 5–10% by the presence of the TiO$_2$ (Fig. 2a and Table S2). Comparison of the hACN + TiO$_2$ with the pure hACN spectra also indicated a larger relative elastic contribution to the total scattering intensity in the presence of TiO$_2$ (see Fig. 2b). Since only...
approximately 2% of the elastic scattering can be attributed to the presence of TiO2 (Table S2), this suggests that about 8% of the hACN maybe immobilized on the measurement timescale by the presence of the TiO2. We note that the minimum Q value detectable by the OSIRS instrument is 0.18 Å\(^{-1}\), and expect that if lower Q values were detectable then elastic scattering fraction would extrapolate to 1 at sufficiently low Q.

Acetonitrile has a surface density of approximately 24 Å\(^2\) per molecule\(^{45}\), so we infer the immobile fraction of the 1.6 g of solvent in the hACN\(^+-\)TiO2 sample corresponds to about 4.5 \(\times\) 10\(^2\) m\(^2\) assuming negligible evaporation during preparation. We estimate the exposed surface of the 1.5 g of 21 nm diameter TiO2 nanoparticles in the sample to be approximately 1.1 \(\times\) 10\(^2\) m\(^2\), a similar order of magnitude. This is consistent with the immobile fraction corresponding to around 3–4 immobile layers of acetonitrile molecules coating the TiO2 surface, with the remaining fraction experiencing close to bulk liquid diffusion. These observations are in reasonable agreement with first principles molecular dynamics simulations by Schiffman \textit{et al.}\(^{27}\), who predicted the formation of several structured solvent layers at the TiO2 interface which suppress diffusion.

\textbf{Adsorbate dynamics.} We now examine the possible dynamics of the dye moieties adsorbed on the TiO\(_2\) nanoparticles. Initially, we compare sensitized and unsensitized TiO\(_2\) without solvent. The uptake of INA and BINA was estimated from UV-vis measurements to be about 0.016 and 0.018 grams per gram of TiO2, respectively (1.6% and 1.8% by weight without solvent). This is consistent with complete coverage of the nanoparticles by a monolayer of the adsorbates.

A comparison of the QENS spectra for TiO\(_2\), TiO\(_2\) sensitized with INA, and TiO\(_2\) sensitized with BINA is shown in Fig. 3, indicating no significant difference in the spectra. This was the case for all Q groups at temperatures ranging from 250 to 290 K.

20% and 17% of the scattering signal in the sensitized samples should arise from incoherent contributions from INA and BINA respectively (see Table S1). Consequently, if the dye moieties were mobile within the spatio-temporal window probed by the measurement we would expect a measurable broadening of the elastic peak in the sensitized samples relative to the control sample of up to around 20% of the scattering intensity. Since no difference was observed, we conclude that INA and BINA do not move within the time window of the measurement, spanning approximately 10 ps and a few ns.

Next we examined whether the presence of acetonitrile surrounding the sensitized particles could allow motion of the moieties. To reduce the effect of incoherent scattering from hydrogen in the solvent which would mask any motion of the adsorbates, a 1:1 by weight ratio of deuterated acetonitrile (dACN) to BINA sensitized TiO\(_2\) particles was used. This was compared to a similar weight ratio of dACN and unsensitized TiO\(_2\). From Table S2 we expect incoherent scattering from BINA to contribute to only 5% of the total scattering signal in the sensitized sample with solvent. Thus even if BINA is mobile within the spatio-temporal window probed, we would expect its contribution to QENS measurement to be small and difficult to distinguish from the large contributions from dACN (77%) and TiO\(_2\) (18%). The magnitude of the QENS spectra from a pure dACN control sample was around 17% that of hACN, as expected due to the significantly lower incoherent scattering strength of deuterium relative to hydrogen. A comparison of QENS spectra of the mixture of dACN + BINA + TiO\(_2\), dACN + TiO\(_2\) and dACN is shown Figure S2.

In an effort to remove the contribution from dACN to these data, the scattering spectrum of pure dACN multiplied by its expected contribution (inferred from its weight) in the sample was subtracted from the spectrum of the sample mixtures. The resulting spectra were then normalized. As above, no appreciable difference can be seen between the sensitized and unsensitized mixtures (see Fig. 4). Thus, given the experimental uncertainty, we see no evidence for the motion of BINA within the spatio-temporal window probed between 270 and 290 K, which is consistent with the observations from the INA and BINA sensitized samples in the absence of solvent.
Discussion
We observed that TiO₂ nanoparticles immersed in acetonitrile immobilise a fraction of acetonitrile molecules corresponding to several monolayer coverage of the particles. The remaining mobile fraction experiences close to bulk diffusion, consistent with previous computational work. We believe that polar solvents (such as acetonitrile) will result in a strong interaction with the TiO₂ surface leading most molecules in contact with the surface being immobilised. Apolar/aprotic solvent are less likely to form completely immobilized interfacial layers. It is possible that the adsorption of dye molecules to the TiO₂ surface might reduce the immobilisation acetonitrile however we did not have a straightforward approach to distinguish between these two immobilized components with our experiments.

The monolayers of the dye moieties, INA and BINA, anchored to the TiO₂ were immobile on the timescale resolvable with OSIRIS (tens of picoseconds to few nanoseconds). If structural rearrangements of these molecules occur, they do so on longer timescales. Extrapolation of our observations to larger dyes suggests that molecules containing anchoring moieties similar to INA and BINA, packed on the TiO₂ surface with a similar coverage, are unlikely to undergo significant motions articulated from the anchoring point within this window. Clearly, more complex dye molecules could allow additional degrees of freedom around which motion of other moieties are be possible. This would be necessary (on timescales exceeding nanoseconds, possibly up to microseconds) if changes in molecular conformation explain the observed charge diffusion rates. A wider temporal resolution and smaller Q-range would be necessary to resolve slower and rotational relaxations of larger molecules. Other techniques such as nuclear magnetic resonance relaxation might also yield information on the possible molecular dynamics on longer timescales.

Methods
Sample preparation: 1.2 grams of 2,2′-Bipyridyl-4,4′-dicarboxylic acid (BINA) was added to 400 mL of N,N-Dimethyl formamide (DMF) in a 500 mL bottle. Alternatively, 1.2 grams of Pyridine-4-carboxylic acid (INA) was added to 400 mL ethanol. The solution was then heated to 70 °C and stirred at 700–800 rpm until complete dissolution of the dye. 3 grams of Titanium (IV) oxide (TiO₂) anatase nanopowder (∼21 nm particle size, Sigma Aldrich) was added to the solution. The suspension was centrifuged at 4000 rpm for 5 minutes, followed by three ethanol washes, each followed by further centrifugation at 4000 rpm for 5 minutes to ensure that no residue of the high boiling point DMF (Tboil(MeOH) = 153 °C) remained in the sample. The mass of dye adsorbed to the TiO₂ was determined by UV-vis spectrophotometry of the dye solution before and after the addition of the nanoparticles. The samples were stored in ethanol overnight. Just before the loading of the sample cans, the samples were dried in two steps: hotplate at 80 °C and vacuum oven at 80 °C (Tboil(MeOH) = 78 °C) for approximately half an hour each. Finally, about 1.5 grams of dried powder was either transferred into the same mass of solvent (hydrogenated or deuterated acetonitrile) or directly into the sample sachet made of aluminium foil. For the samples with solvent, the resulting paste was spread onto the sachet. The sachets were rolled into the OSIRIS cylindrical sample cans made of aluminium where the sample was contained within an annulus 0.5–1 mm thick with a circumference of around 70 mm and height 50 mm.

Quasi-elastic neutron scattering (QENS) measurements were performed at OSIRIS, a high resolution (25 μeV) spectrometer within the ISIS pulsed neutron source facilities in the Rutherford Appleton Laboratory (Oxfordshire). The dynamic energy range and momentum transfer range (Q-range) are −0.4–0.4 meV and 0.18–1.8 Å⁻¹ respectively. The analysing energy is 1.84 meV. Calibration measurements were made using a purely incoherent scattering vanadium sample. Including controls, measurements were performed on samples of pure hydrogenated acetonitrile, pure deuterated acetonitrile, TiO₂ nanoparticle powder, TiO₂ nanoparticle paste with hydrogenated acetonitrile, TiO₂ nanoparticle paste with deuterated acetonitrile, INA or BINA sensitized TiO₂ nanoparticles, and BINA sensitized TiO₂ nanoparticle paste with deuterated acetonitrile. All the aforementioned
measurements were made at three temperatures 250, 270 and 290 K which are above the melting point of acetonitrile (228 K). The beam intensity at the sample was $2.7 \times 10^7$ neutrons cm$^{-2}$ s$^{-1}$ at 150 μA ISIS beam current, and samples were measured for 30 μAh.

The QENS spectra $S_{\text{raw}}(Q, E')$, for each sample were extracted from the raw QENS data ($I_{\alpha}$) with the software package Mantid (data reduction)\(^47\). The data from different ranges of detector angles were combined into ten groups to improve the measurement statistics.

The possible dynamics of molecules can be resolved by analysing the intensity of the spectrum of scattered neutrons from each nuclei, $S_{\text{coh}}$ which parametrically depends on the scattering regime $i$ (i.e. coherent or incoherent) and the type of the nuclei in the sample $j$. $S_{\text{coh}}$ is related to the intensity of the scattered neutron beam, $I_{\alpha}$ through\(^41,42\):

$$I_{\alpha}(Q, E') = I_0(E_0) \frac{k_0}{k_q} \sum_i \alpha_i S_{\text{coh},i}(Q, E') \otimes R(Q, E')$$

(2)

where $I_0$ is the incident neutron flux, $(E_0, k_0)$ and $(E', k_q)$ are the incident and scattered neutron beam energy and wavevectors respectively, $Q$ is the magnitude of the momentum transfer $(Q = |k_q - k_0|)$, and $\alpha_i$ is a factor accounting for the relative scattering magnitude of each nucleus $i$ in regime $i$. The scattered signal is convolved with the instrument response function, $R(Q, E')$, which results in a spread of energies for all scattering events. For coherent scattering, $\alpha_{\text{coh}} = B^2$ while for incoherent scattering, $\alpha_{\text{inc}} = B - B^2$ where $B$ and $B^2$ are the scattering length and spatially averaged scattering length of each nuclei\(^41,42\). These scattering lengths were used to calculate the expected contributions to the total scattering signal from the different samples examined in this study (Table S2).

For hydrogenated samples, such as acetonitrile (hydrogenated ACN was used for measurements of solvent diffusion), INA and BINA, a major contribution to the total intensity is the incoherent scattering signal of hydrogen\(^40–42\). Consequently, the scattering intensity can often be approximated by:

$$I_{\alpha}(Q, E') \approx I_0(E_0) \frac{k_0}{k_q} \alpha_{\text{inc},i} S_{\text{inc},i}(Q, E') \otimes R(Q, E').$$

(3)

However, due to the low concentration of the dye moieties in this study the contribution to the scattering spectrum from the other nuclei was often significant. The relative contribution to the scattering signal from each component in the different samples is shown in Table S2, this was calculated from the scattering cross sections and molar fraction of the different nuclei in the samples.

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Author Contributions

X.L. and P.B. prepared the samples, V.G.S. and P.B. collected the QENS data, V.V. and V.G.S. analysed the results with assistance from J.T.C., J.N. and P.B. V.V., V.G.S. and P.B. prepared the manuscript with contributions from all other authors.

Additional Information

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