Solvent Dependence of Lateral Charge Transfer in a Porphyrin Monolayer

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Supporting Information

ABSTRACT: Lateral charge transport in a redox-active monolayer can be utilized for solar energy harvesting. A model porphyrin system was chosen to study the influence of the solvent on lateral hole hopping, which plays a crucial role in the charge-transfer kinetics. We examined the influence of water, acetonitrile, and propylene carbonate as solvents. Hole-hopping lifetimes varied by nearly 3 orders of magnitude among solvents, ranging from 3 ns in water to 2800 ns in propylene carbonate, and increased nonlinearly as a function of added acetonitrile in aqueous solvent mixtures. These results elucidate the important roles of solvation, molecular packing dynamics, and lateral charge-transfer mechanisms that have implications for all dye-sensitized photoelectrochemical device designs.

Surface-bound monolayers of redox active molecules on semiconductors are known to transport charges laterally along the surface.1–4 This property has been studied and exploited for various uses, namely in relation to molecular dye-sensitized semiconductors for solar energy conversion.5–14 In these systems, photoexcitation of the dye induces electron transfer to the n-type semiconductor, leaving an electron hole on the molecule. This hole can diffuse among the surface-bound molecules in competition with electron recombination processes. Recent reviews discuss current developments in this research area.15,16

In solid-state dye-sensitized solar cells, lateral hole diffusion allows charges to more efficiently access the hole-transport species, generally Spiro-MeOTAD.8 In photoelectrocatalytic cells, lateral hole transport facilitates oxidation of a catalyst dispersed among the dyes.17 The surface-bound monolayer functions as a light-harvesting system and electronically connects the catalyst to multiple dye molecules. Lateral charge transport is a promising route toward overcoming the diffuse nature of sunlight and low efficiencies in multielectrode photoelectrocatalytic cell designs.18–20 Under standard AM1.5G sunlight, which corresponds to 100 mW/cm², a dye absorbs approximately one photon per second,21,22 with photoinduced charges recombining orders of magnitude faster.23 Therefore, in order to use sunlight to perform a multielectron catalytic reaction such as the four-electron oxidation of water, a significant number of dyes must work in conjunction with a single catalyst. Maximizing lateral charge-transfer rates is thus essential to increasing solar photoelectrocatalytic device efficiencies.

In a dye–catalyst monolayer system, faster lateral charge transport competes more effectively with charge recombination between the photoinjected electron and oxidized dye, allowing a larger number of dye molecules to electronically access a single catalyst. In past work, we developed a porphyrin system to study the various factors affecting lateral charge transfer.1 The design in Figure 1 depicts a light absorbing monolayer of freebase porphyrin (monomer H2PF8) on a SnO2 nanoparticulate film. Randomly dispersed among this monolayer are porphyrin dyads consisting of a surface-bound freebase porphyrin dyad, H2PF8−ZnPF8, that act as thermodynamic sinks for the hole, similar to a catalyst. Previous research has shown that photons absorbed by ZnPF8 efficiently transfer that energy to the freebase moieties.1 The photoexcited H2PF8 monomer or dyad moiety (1) transfers an electron to the SnO2 conduction band (2), yielding a charge-separated state with a conduction band electron and porphyrin radical cation (hole). The hole is then able to hop among neighboring porphyrins (3) in competition with recombination processes. Hole transfer to the H2PF8 moiety of the dyad results in intramolecular charge transfer to the zinc porphyrin on the subnanosecond time scale (4). The hole is then trapped and undergoes normal recombination processes with a semiconductor electron.

Using nanosecond transient absorption spectroscopy (ns-TAS), we were able to monitor the relative ratios of oxidized freebase and zinc porphyrin species in the monolayer as a function of time, which can be simulated using a hole-transfer model (see the Supporting Information for model description).
We previously analyzed different ratios of monomer and dyad porphyrin monolayers (∼90% theoretical monolayer surface coverage) on SnO₂ films under aqueous conditions. Our model indicated that lateral hole diffusion across the freebase porphyrin monolayer occurs via a random-walk process. While this was expected, the rate of 4.0 × 10⁷ s⁻¹ (τ = 25 ns) for each hole-hopping event was unexpectedly fast. The lateral hole-hopping rates were closer to those for intramolecular charge transfer rather than an intermolecular process and outside the range of lateral hole-hopping rates studied in other porphyrin and ruthenium polypyridyl systems. In those studies, hole-hopping lifetimes ranged between hundreds of nanoseconds for the fastest systems to systems where no lateral transfer was observed at all.

Monolayer samples with a 25:1 ratio of H₂PF₈ (monomer) to H₂PF₈−ZnPF₈ (dyad) were prepared. At this ratio, we can use ns-TAS to study lateral charge-transfer rates ranging from reciprocal nanoseconds to reciprocal milliseconds. Ultraviolet (UV)–visible spectra of the molecules and the dye-sensitized SnO₂ films are shown in Figure 2. Slight spectral shifts between the solution and dry surface-bound material are expected and observed, and it is apparent that there is only a small percentage of dyad on the surface (see Figure S2).

The samples were immersed in different solvents [water, acetonitrile (ACN), water/acetonitrile mixtures (vol %), and propylene carbonate (PC)] containing 100 mM LiClO₄ as electrolyte. They were analyzed by using ns-TAS, which entailed photoexciting the samples with 515 nm light and probing at 560 nm. The freebase H₂PF₈ is primarily photoexcited at the chosen pump wavelength and injects an electron into the SnO₂ conduction band. The ns-TAS signature of the H₂PF₈ radical cation (hole) shown in Figure 3 is then observed as a positive value at the probe wavelength. The hole randomly traverses the monolayer and can either recombine with SnO₂ conduction band electrons or

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**Figure 1.** Model system for studying lateral charge transport. R₁ = 2,3,5,6-tetrafluorophenyl; R₂ = 4-carbomethoxyphenyl; R₃ = 4-trifluoroacetamidophenyl.

**Figure 2.** UV–visible spectra in the Q-band region of model porphyrins in solution and dye-sensitized SnO₂. Spectra of monomer (black solid line) and dyad (red solid line) in ethyl acetate solvent were obtained from ref 1, and the absorbance of a 25:1 monomer-to-dyad sample was obtained using a dry SnO₂ film (black dotted line).

**Figure 3.** TA map of monomer (solid lines) and dyad (dashed lines) porphyrin radical cations. Data were normalized at the probe wavelength of 560 nm.
reside on the H$_2$PF$_8$ moiety of a dyad, in which case rapid intramolecular charge transfer produces the ZnPFP$_8$ radical cation, which has a negative TA signature at 560 nm due to a bleach of the zinc porphyrin Q-band. The trapped hole on ZnPFP$_8$ then undergoes recombination and the signal decays. The observed signal as a function of delay time is the sum of H$_2$PF$_8$ and ZnPFP$_8$ radical cation contributions. The ratio of extinction coefficients at 560 nm for the ZnPFP$_8$ (negative signal) and H$_2$PF$_8$ (positive signal) porphyrin radical cations was previously determined to be $-3.5$.$^1$

Monolayers of pure monomer or dyad on SnO$_2$ in each solvent were analyzed by ns-TAS to determine their recombination rates (see Figure S3). The decay of signal due to recombination at early delay times is well-described by a stretched exponential decay function corresponding to the Kohlrausch–Williams–Watts (KWW) kinetic model, which is commonly used for molecular dyes on metal oxides.$^{25}$

Transient absorption spectra revealed only minor solvatochromic shifts among solvents (Figures 3 and S4). For the mixed-species monolayers, it is apparent in Figure 4 that the ns-TAS signals transition from positive values toward negative values much faster for samples in water than those with added acetonitrile or in the pure organic solvents. Signals for all solvents eventually become negative before decaying back toward the baseline by recombination processes (see Figure S5). As was observed previously,$^7$ the dyad has different recombination kinetics when dispersed in a H$_2$PF$_8$ monomer monolayer compared to a pure monolayer of dyad, and this is the case in all the solvents examined in this study. The ZnPFP$_8$ solvation sphere is entirely that of solvent when dispersed in a monolayer of H$_2$PF$_8$ monomer, while in the pure monolayer of dyad it is surrounded by other ZnPFP$_8$ moieties. This solvent accessibility leads to a much slower recombination rate in the more isolated dyads.$^{25}$

Data shown in Figure 4, along with the recombination kinetics for both monomer and dyad, were input into a random-walk model previously described to simulate the transient absorption spectra and calculate the hole-hopping rate.$^7$ The fits are presented in Figure 4, with a longer time window shown in Figure S5. The observed hole-hopping lifetimes are presented in Table 1. The stretched exponential decay recombination dynamics were relatively consistent among solvents. For the monomer, values of $\beta$ vary between 0.20 and 0.25, and recombination rates vary between $1.0 \times 10^5$ and $1.3 \times 10^6$ s$^{-1}$. For the dyad, recombination rates vary between $0.5 \times 10^3$ and $2.3 \times 10^3$ s$^{-1}$ with a fixed $\beta$ value of 1 corresponding to a single exponential decay in the KWW model.

While recombination rates are similar among solvents, the hole-hopping lifetimes span nearly 3 orders of magnitude from 3 to 2800 ns. We obtain 3 ns per hop in water, which is similar to our previously published value (25 ns),$^1$ and the hole-hopping lifetime increases significantly as the proportion of acetonitrile is increased. The results in acetonitrile (1800 ns) and propylene carbonate (2800 ns), which are polar organic solvents, are similar to each other. This contrasts with previous reports in the literature, as comparable rates are obtained for ruthenium bipyridine analogs in both water and acetonitrile.$^{5,26}$

The wide range of calculated lateral charge-transfer rates in the water/acetonitrile mixtures provides insight into the dynamics. According to Marcus theory of electron transfer in a classical nonadiabatic system, electron self-exchange rates are strongly dependent on the solvent because of inherent differences in outer-sphere reorganization energies.$^{27,28}$ The rate, $k$, of electron self-exchange (i.e., when $\Delta G^0 = 0$) is described in its most general form in eq 1 as

$$k = \frac{2\pi}{\hbar} \left| V \right|^2 \frac{1}{\sqrt{4\pi \beta k_b T}} \exp \left( -\frac{\lambda}{4k_b T} \right)$$

This relates the rate of electron transfer ($k$) to the electronic coupling ($V$) between the donor and acceptor, the reorganization energy of the reaction ($\lambda$), the Boltzmann constant ($k_b$), and the temperature ($T$). In solvents with higher polarity, $\lambda$ generally increases, causing $k$ to decrease. Our results are the opposite of this general trend and complement recent results by Moia and co-workers using a different molecular system.$^{11}$ Multiple explanations can be hypothesized for the unexpected results, but most can be discounted outright because of the magnitude of the change in hole-hopping rates. One possible explanation for the wide discrepancy is solvent-dependent ion pairing of the H$_2$PF$_8$ moiety. Ion pairing is a complicated phenomenon that has been theorized and experimentally observed to significantly affect charge-transfer rates by increasing outer-sphere reorganization energies.$^{29–34}$ The strength of ion pairing correlates with spectral shifting of the ion, in this case that of the monomer porphyrin radical

Table 1. Lateral Charge-Transfer Rates and Model Kinetic Parameters$^a$

<table>
<thead>
<tr>
<th>solvent</th>
<th>$k_{\mathrm{exp}}$ (sec$^{-1}$)</th>
<th>$\tau_{\mathrm{mono}}$ (ns)</th>
<th>$k_{\mathrm{rec,mono}}$ (sec$^{-1}$)</th>
<th>$\beta_{\mathrm{mono}}$</th>
<th>$k_{\mathrm{rec,dyad}}$ (sec$^{-1}$)</th>
<th>$\beta_{\mathrm{dyad}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>H$_2$O</td>
<td>$3.6 \times 10^3$</td>
<td>3</td>
<td>$1.3 \times 10^3$</td>
<td>0.25</td>
<td>$2300$</td>
<td>1.0</td>
</tr>
<tr>
<td>25% ACN</td>
<td>$8.9 \times 10^2$</td>
<td>11</td>
<td>$1.3 \times 10^2$</td>
<td>0.25</td>
<td>$1500$</td>
<td>1.0</td>
</tr>
<tr>
<td>50% ACN</td>
<td>$5.0 \times 10^2$</td>
<td>20</td>
<td>$1.0 \times 10^2$</td>
<td>0.20</td>
<td>$1200$</td>
<td>1.0</td>
</tr>
<tr>
<td>75% ACN</td>
<td>$2.5 \times 10^2$</td>
<td>40</td>
<td>$8.0 \times 10^1$</td>
<td>0.25</td>
<td>$1000$</td>
<td>1.0</td>
</tr>
<tr>
<td>ACN</td>
<td>$5.6 \times 10^2$</td>
<td>1800</td>
<td>$4.0 \times 10^1$</td>
<td>0.23</td>
<td>$700$</td>
<td>1.0</td>
</tr>
<tr>
<td>PC</td>
<td>$3.6 \times 10^2$</td>
<td>2800</td>
<td>$2.6 \times 10^2$</td>
<td>0.25</td>
<td>$500$</td>
<td>1.0</td>
</tr>
</tbody>
</table>

$^a$Acetonitrile solvent mixtures with water are volume %.
cation shown in Figure S4. The relatively minor observed spectral shift of 5–10 nm among the solvents containing LiClO₄ electrolyte can be attributed to general solvatochromism. Therefore, ion pairing need not be considered as a possibility in our system.35,36

We believe hydrophobic clustering provides the simplest explanation for the significant differences in solvent-dependent hole-hopping rates. Shown in Figure 5, hydrophobic clustering occurs when water is excluded from the solvation sphere of the largely hydrophobic porphyrin. Because the porphyrin is not soluble in water, and the surface-anchoring carboxylic acid has been shown to be unstable,37,38 the porphyrins could undergo a hydrophobic collapse into a more well-packed monolayer, thereby excluding solvent waters. This solvent exclusion process could be driven by the entropy gained from the loss of solvation waters, similar to micellization.39 Thus, in water, the porphyrins are actually solvated by the relatively nonpolar appendages of the surrounding porphyrins, a conclusion supported by our results that agrees with Marcus electron-transfer theory.

In addition to merely altering the solvation environment, rearrangement to a more well-packed monolayer would decrease the distance between porphyrins. While previous results have shown that the initially formed monolayer is relatively well-packed,40 removing a single shell of hydration or even interstitial solvent molecules from each porphyrin would decrease the interporphyrin tunneling distance by a significant amount. These two factors both exponentially increase the charge-transfer rate according to the Marcus theory of electron transfer and can account for the wide range of hole-hopping rates in the solvent systems.28

The generalized form of Marcus electron-transfer theory in eq 1 can be further differentiated for our specific case. Equations 2 and 3 provide a summary of the effects of distance, tunneling medium, and reorganizational energies on electron-transfer rates. In eq 2, the electronic coupling matrix element (V), which is directly proportional to the rate of electron transfer (k), increases exponentially with decreasing distance (R) between porphyrins beyond van der Waals separation (R₀).28 Altering the solvation environment also modifies the medium-dependent scalar for electron tunneling ($\beta$), which becomes smaller in more nonpolar solvents, further increasing the electron-transfer rate as water is replaced by a less polar environment.40 In eq 3, k is also exponentially dependent on the reorganizational energy (\lambda), which is generally dominated by outer-sphere solvent dynamics.41 If solvent is excluded and replaced by rigid aryl functionalities, a significant decrease in $\lambda$ is expected with a concomitant increase in k. A caveat to this assessment is the unknown nature of the metal oxide surface, where solvation and surface potentials could affect the local environment around the porphyrin.

$$k \propto V \propto \exp \left( -\frac{\beta}{2} (R - R_0) \right)$$

(2)

$$k \propto \exp \left( -\frac{\lambda}{4} \right)$$

(3)

Our results in the pure organic solvents, which we believe solvate porphyrins in a more ideal way, allow direct comparison to hole-hopping kinetics in other model systems using porphyrins or inorganic coordination complexes. The average hole-hopping lifetimes in acetonitrile and propylene carbonate of 1800 and 2800 ns, respectively, are similar to that observed by others.7,6 Thus, our results in aqueous solutions are extraordinary, while those in organic solvents are fairly typical. Because of the similarity of the lifetimes in these pure organic solvents, there is not enough information to conclusively show that the system follows classical Marcus theory where charge-transfer rates increase with decreasing solvent Pekar factor (given by $1/\epsilon_m - 1/\epsilon_0$ and is 0.53 for acetonitrile and 0.48 for propylene carbonate) or with decreasing solvent longitudinal relaxation time (acetonitrile, 0.2 × 10⁻¹² s; propylene carbonate, 2.6 × 10⁻¹² s).42 Lifetimes measured in additional organic solvents would be needed, but these initial results suggest it is the latter.

From the lateral charge-transfer model, we can estimate the harvesting efficiency of the photoinduced charges by determining the percentage of holes that become localized on the dyad prior to recombination. In the case of water, 77% of holes become localized while only 19% are localized in acetonitrile and 17% in propylene carbonate. The higher rate of lateral charge transfer under aqueous conditions makes the overall efficiency of hole hopping to the dyad relatively high. In a monolayer with porphyrin—dyad ratio of 25:1, a majority of the charge carriers produced by absorption of solar energy can be collected by the dyad which comprises less than 4% of the monolayer. This is an important result for the development of aqueous photoelectrocatalytic devices where multiple dyes must electronically interact with a single catalyst.

In summary, we describe the solvent dependence of lateral charge transfer in a porphyrin-based model system, generating new understanding of the roles of solvation and the complexity of surface dynamics on the kinetic rates. Our previous results suggesting relatively fast hole hopping in water have been confirmed with a lifetime of 3 ns in the system studied with aqueous LiClO₄ electrolyte, and this lifetime is shown to increase dramatically toward 1800 ns as the solvent fraction of acetonitrile is increased. Our hypothesis is that the porphyrin monolayer reorganizes to exclude solvent, thereby increasing the hole-hopping rate by decreasing the electron tunneling parameter ($\beta$), reorganizational energy ($\lambda$), and likely the intermolecular porphyrin distances. These results provide new insights into lateral charge-transfer mechanisms and are of interest to those studying redox-active monolayers in dye-sensitized systems. The results are particularly promising for
the prospect of exploiting lateral charge transfer in aqueous environments, such as photoelectrocatalytic water oxidation.

**ASSOCIATED CONTENT**

* Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsenergylett.6b00583.

Experimental procedures, description of the model, characterization of samples, and additional transient absorption data (PDF)

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Notes

The authors declare no competing financial interest.

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