Proton-Coupled Electron Transfer

Distance Dependence of Bidirectional Concerted Proton–Electron Transfer in Phenol-Ru(2,2'-bipyridine)₃²⁺ Dyads

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Abstract: Proton-coupled electron transfer (PCET) was investigated in three covalent donor–bridge–acceptor molecules with different bridge lengths. Upon photoexcitation of their Ru(bpy)₃²⁺ (bpy=2,2'-bipyridine) photosensitizer in acetoni-trile, intramolecular long-range electron transfer from a phenolic unit to Ru(bpy)₃²⁺ occurs in concert with release of the phenolic proton to pyrrolidine base. The kinetics of this bidirectional concerted proton–electron transfer (CPET) reaction were studied as a function of phenol–Ru(bpy)₃²⁺ distance by increasing the number of bridging *p*-xylene units. A distance decay constant (β) of 0.67±0.23 Å⁻¹ was deter-

Introduction

The distance dependence of "simple" electron transfer has been thoroughly explored in proteins,^[1] DNA,^[2,3] self-assembled monolayers,^[4] and donor–bridge–acceptor molecules.^[5,6] An important open question is how the distance between an electron donor and an electron acceptor affects the rates of proton-coupled electron transfer (PCET) reactions, particularly when electron and proton transfer occur in concerted fashion and when the electron and the proton are transferred into different directions. Long-range PCET reactions of this type are biologically relevant, for instance in photosystem II, in ribonucleotide reductase, and in DNA.^[7-9] We are unaware of prior investigations of the distance dependence of bidirectional concerted proton-electron transfer (CPET) reactions.

The rate constant for a PCET reaction can be expressed in a similar way as the rate constant for "simple" electron transfer, that is, in analogy to what is commonly known as the Marcus equation.^[10] In Equation (1), $H_{AB,PCET}$ is the electronic coupling between the PCET reactant and product states, λ_{PCET} is the reorganization energy associated with the overall PCET process, and ΔG^{0}_{PCET} is the reaction free energy.^[11] The interplay between these three factors governs the PCET rate constant (k_{PCET}).

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mined. The distance dependence of the rates for CPET is thus not significantly steeper than that for ordinary (i.e., not proton coupled) electron transfer across the same bridges, despite the concerted motion of oppositely charged particles into different directions. Long-range bidirectional CPET is an important reaction in many proteins and plays a key role in photosynthesis; our results are relevant in the context of photoinduced separation of protons and electrons as a means of light-to-chemical energy conversion. This is the first determination of β for a bidirectional CPET reaction.

In many cases of "simple" electron transfer the distance dependence of reaction rates is dominated by the electronic coupling term $(H_{AB})^{[1,5]}$ but in other cases changes in reorganization energy (λ) or reaction free energy (ΔG^0) with increasing donor–acceptor distance are more important.^[12,13] It is not a priori clear which one of the three factors ($H_{AB,PCET}$, λ_{PCET}) will dominate the distance dependence of a bidirectional CPET reaction. On the one hand, increasing the electron donor/electron acceptor distance is expected to have a significant influence on $H_{AB,PCET}$ ^[11] On the other hand, it is known that bidirectional CPET reactions can be associated with significant-ly larger reorganization energies (λ_{PCET}) than "simple" electron-transfer reactions,^[14] and significant changes in λ_{PCET} with distance could strongly influence CPET rates.

$$k_{\text{PCET}} = \frac{2\pi}{\hbar} H_{\text{AB,PCET}}^2 \frac{1}{\sqrt{4\pi\lambda_{\text{PCET}}RT}} \exp\left[-\frac{\left(\Delta G_{\text{PCET}}^0 + \lambda_{\text{PCET}}\right)^2}{4\lambda_{\text{PCET}}RT}\right]$$
(1)

In the work presented herein we aimed to explore these basic aspects by using the phenol-Ru(bpy)₃²⁺ (bpy=2,2'-bipyridine) dyads shown in Scheme 1. They are comprised of a phenolic



Scheme 1. Molecular structures of the PhOH- xy_n -Ru²⁺ molecules (n = 1-3) investigated in this work and illustration of their light-induced bidirectional CPET reaction involving pyrrolidine. ET = electron transfer, PT = proton transfer.

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unit acting as a combined electron-proton donor, *p*-xylene bridges of variable length, and a ruthenium photosensitizer as an electron acceptor. The proton acceptor is a base (pyrrolidine) which is added separately to the solution. Phenols have received much attention for mechanistic PCET studies,^[15-17] and there have been several investigations of donor-bridge-acceptor molecules incorporating phenols.^[18-24] Long-range PCET reactions have received increasing attention recently,^[24-28] but to our knowledge the dependence of the rates for bidirectional CPET on the electron donor/electron acceptor distance has never been investigated before. Given the biological relevance of this reaction type a systematic investigation of the model compounds from Scheme 1 seemed to be a worthy research endeavor.

Results and Discussion

Basic photochemistry and electrochemistry of the dyads

In pure acetonitrile photoexcitation of the three dyads from Scheme 1 leads to ³MLCT emission of the Ru(bpy)₃²⁺ moieties, but there is no photochemistry. In absence of any base the ³MLCT excited state lifetimes of the dyads are within experimental accuracy the same as for isolated Ru(bpy)₃²⁺. Although a significantly better oxidant in the ³MLCT state than in the electronic ground state, Ru(bpy)₃²⁺ is obviously not strongly oxidizing enough to convert the phenols to phenoxyl radical cations (PhOH⁺). This finding can be understood on the basis of the electrochemical data in Table 1 which were extracted

Table 1. Electrochemical potentials (in [V] vs. Fc^+/Fc in CH_3CN with 0.1 M
$Bu_4NPF_6\!)$ of the individual redox-active moieties of the dyads and some
reference molecules.

Molecule	Ru ^{III/II}	bpy/bpy ⁻	PhOH ⁺ /PhOH	PhO'/PhO-
PhOH-xy ₁ -Ru ²⁺	0.90	-1.73	0.88	-0.54
PhOH-xy ₂ -Ru ²⁺	0.90	-1.72	0.90	-0.58
PhOH-xy ₃ -Ru ²⁺	0.90	-1.72	0.86	-0.58
Ru(bpy) ₃ ²⁺	0.90	-1.73		
2,4,6- <i>t</i> Bu ₂ PhOH			1.18 ^[a]	-0.70 ^[a]
These data were extracted from the cyclic voltammograms shown in Fig-				

ure S1 in the Supporting Information. [a] From Ref. [15].

from cyclic voltammetry measurements (Figure S1 in the Supporting Information). Reduction of Ru(bpy)₃²⁺ is ligand based and occurs at approximately -1.7 V vs. Fc⁺/Fc in CH₃CN in all cases considered here. Oxidation of phenol (PhOH) to phenoxyl radical cation (PhOH⁺) occurs at approximately 0.9 V vs. Fc⁺/Fc in the three dyads. Given a ³MLCT energy (ΔE_{MLCT}) of 2.1 eV,^[29] one estimates a free energy of $\Delta G^0_{ET1} = -e \times [E(bpy/bpy^-) - E(PhOH⁺/PhOH)] - \Delta E_{MLCT} \approx +0.5$ eV for photoinduced electron transfer.^[30] Such driving-force estimates are usually accurate to ± 0.1 eV. In the present case it seems safe to conclude that "simple" electron transfer from phenol to photoexcited Ru(bpy)₃²⁺ in pure CH₃CN does not occur at measurable

rates because it is too endergonic. Pyrrolidine does not deprotonate our charge-neutral phenols to a significant extent (see pK_a values below).

The situation changes completely when base is present because the neutral phenoxyl radical (PhO') instead of phenoxyl radical cation (PhOH⁺) can be formed upon photoexcitation of the Ru(bpy)₃²⁺ photosensitizers. In other words, phenol oxidation becomes thermodynamically possible when combined with proton release (see below for driving-force estimates). In acetonitrile solution, addition of 4-aminopyridine quenches the ³MLCT excited state of the PhOH-xy₁-Ru²⁺ dyad but has no effect on the longer congeners. The stronger base pyrrolidine leads to a photoreaction in all three dyads. We have therefore chosen pyrrolidine as a base for our PCET investigations. Importantly, up to concentrations of 1 M in CH₃CN pyrrolidine does not have any noticeable influence on the ³MLCT decay of the Ru(bpy)₃²⁺ reference complex (Figure S2 in the Supporting Information).

Photochemistry in presence of pyrrolidine

Transient absorption spectra of the three dyads in aerated CH_3CN obtained in presence of excess pyrrolidine are shown in Figure 1 a (solid black lines). These spectra were measured



Figure 1. a) Solid black lines: Transient absorption spectra measured after excitation of the three dyads in aerated CH_3CN (2×10^5 M) in presence of pyrrolidine at 532 nm. Detection occurred by time averaging over a period of 200 ns starting immediately after excitation with pulses of ca. 10 ns duration. The pyrrolidine concentrations were: 25 mM (PhOH-xy₁-Ru²⁺), 96 mM (PhOH-xy₂-Ru²⁺), and 3 M (PhOH-xy₃-Ru²⁺). Dashed grey traces: Difference spectra obtained by subtraction of the solid black traces in Figure 1 b from the dotted grey traces in Figure 1 b. b) Solid black traces: UV/Vis absorption of the dyads in pure CH₃CN. Dotted grey traces: UV/Vis spectra of the deprotonated dyads in CH₃CN.

after selective excitation of the Ru(bpy)₃²⁺ moieties at 532 nm. The dyad concentrations were always 2×10^{-5} M, but with increasing bridge length increasing pyrrolidine concentrations were necessary in order to induce efficient photochemistry. For PhOH-xy₁-Ru²⁺ a pyrrolidine concentration of 25 mM was sufficient whereas for PhOH-xy₂-Ru²⁺ and PhOH-xy₃-Ru²⁺ concentrations of 96 mM and 3 M, respectively, were employed.



Under these conditions, conversion to the photoproducts was essentially complete after the 10 ns duration of the laser excitation pulses in all three cases. Measurement of the transient absorption data in Figure 1a occurred by time integration over a period of 200 ns. The three resulting spectra are relatively similar to each other and feature a prominent band near 365 nm in addition to weaker bands at longer wavelengths. The shape of the transient absorption spectra can be understood readily on the basis of the UV/Vis data shown in Figure 1b. The solid black traces in Figure 1b are the absorption spectra of the PhOH-xy_n-Ru²⁺ dyads in pure CH₃CN, whereas the dotted grey traces are the absorption spectra of their deprotonated forms (PhO⁻- xy_n -Ru²⁺). Deprotonation occurred by addition of excess tetrabutylammonium hydroxide to the CH₃CN solutions. When subtracting the solid black traces from the dotted grey traces in Figure 1b, one obtains the difference spectra shown as dashed grey traces in Figure 1a. The similarity between these derived difference spectra and the experimental transient absorption spectra (solid black lines in Figure 1a) is evident, and we conclude that the photoproducts which are detected on the nanosecond timescale are comprised of the deprotonated phenols and Ru(bpy)₃²⁺ in the electronic ground state. It is as if the three dyads from Scheme 1 merely acted as photoacids.

Excitation at 532 nm occurs selectively into the $\text{Ru}(\text{bpy})_3^{2+}$ moieties. The phenols have insignificant extinction at this wavelength, and it is not obvious why the phenolic units should become more acidic upon excitation of the $\text{Ru}(\text{bpy})_3^{2+}$ moieties which are more than 12 Å away. However, the apparent photoacid behavior can be explained by a reaction sequence of concerted proton–electron transfer (CPET) followed by "simple" electron transfer (back-ET) in the reverse direction (Scheme 2).^[31] In order to understand this reaction sequence it



Scheme 2. Energy-level diagram established on the basis of the electrochemical potentials in Table 1 and the acidity constants in Table 2 (see the Supporting Information for details).

is necessary to work out the thermodynamics of the different photochemical reaction pathways for the PhOH- xy_n -Ru²⁺ dyads in presence of pyrrolidine (pyr).

The starting point in the energy-level diagram of Scheme 2 is the ³MLCT state 2.1 eV above the ground state.^[29] As discussed above, "simple" photoinduced electron transfer to produce phenoxyl radical cations and reduced ruthenium complex

(pyr/PhOH⁺-xy_n-Ru⁺ state at 2.6 eV) is endergonic by 0.5 eV. The absence of any photochemical reaction in pure CH₃CN without base strongly suggests that the pyr/PhOH⁺-xy_n-Ru⁺ photoproduct is never formed, not even as a short-lived intermediate when base is present. However, on the basis of Equation (2) photoinduced CPET producing protonated pyrrolidine (pyrH⁺), phenoxyl radical (PhO⁺), and reduced ruthenium complex (Ru⁺) is estimated to be exergonic by approximately 0.8 eV.

$$\Delta G^{0}_{PT1} = 0.059 \text{ eV} \times \left[pK_{a}(PhOH^{+}) - pK_{a}(pyrH^{+}) \right]$$
(2)

Specifically, Equation (2) is used to estimate the free energy associated with protonation of pyrrolidine by phenoxyl radical cation.^[32] The p K_a of PhOH⁺ in CH₃CN is -3 and the p K_a of the conjugate acid of pyrrolidine (pyrH⁺) in CH₃CN is 19.56 (Table 2),^[15,33] resulting in $\Delta G^0_{PTI} \approx -1.3$ eV. Consequently, the

Table 2. Acidity constants in CH ₃ CN.			
Acid	р <i>К</i> а		
pyrH ⁺ PhOH PhOH ⁺	$19.56^{[a]}\ 28^{[b]}\ -3^{[b]}$		
[a] From Ref. [33]. [b] Values for 2,4,6-tBu ₃ PhOH from Ref. [15].			

pyrH⁺/PhO⁻xy_n-Ru⁺ state must be approximately 1.3 eV below the pyr/PhOH⁺-xy_n-Ru⁺ state, that is, at 2.6 eV–1.3 eV = 1.3 eV. This is 0.8 eV below the initially populated ³MLCT state. Thus, the CPET reaction in which phenol oxidation by photoexcited Ru(bpy)₃²⁺ occurs in concert with release of the phenolic proton to pyrrolidine (arrow marked with "CPET" in Scheme 2) has ΔG^0_{CPET} =-0.8 eV. It seems reasonable to conclude that this is indeed the first reaction step occurring after photoexcitation.

The pyrH⁺/PhO⁻xy_n-Ru⁺ photoproduct at 1.3 eV is composed of a strongly reducing ruthenium complex (Ru(bpy)₃⁺) connected to a relatively good electron acceptor (PhO⁻). Using Equation (3), one can estimate the reaction free energy associated with intramolecular electron transfer from Ru(bpy)₃⁺ to PhO⁻.

$$\Delta G^{0}_{\text{ET2}} = -e \times [E(\text{PhO}^{\cdot}/\text{PhO}^{-}) - E(\text{bpy}/\text{bpy}^{-})]$$
(3)

With the electrochemical potentials in Table 1, Equation (3) yields $\Delta G^{0}_{ET2} \approx -1.1$ eV. In other words, after initial CPET thermal electron transfer from Ru(bpy)₃⁺ to PhO' is exergonic by 1.1 eV. This second reaction step (downward arrow in Scheme 2 labeled "back-ET") leads to the photoproducts which are detected by transient absorption spectroscopy in Figure 1a, that is, to Ru(bpy)₃²⁺ in the ground state and phenolate. Using the thermodynamic cycle described above, one arrives at the conclusion that the pyrH⁺/PhO⁻-xy₀-Ru²⁺ state is

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approximately 0.2 eV above the ground state. Alternatively, using the p K_a values of phenol (28)^[15] and the conjugate acid of pyrrolidine (19.56)^[33] in CH₃CN one arrives at the conclusion that the respective state is 0.5 eV above the ground state (ΔG^0_{PT2}). The deviation between the two energy estimates for the same state (0.2 eV vs. 0.5 eV) simply reflects the approximate character of these thermodynamic considerations and the uncertainty in the electrochemical potentials and acidity constants.

The conclusion from this section is that the apparent photoacid behavior of the three dyads in presence of pyrrolidine is due to a sequence of two consecutive reaction steps: Photoinduced CPET followed by electron transfer in the reverse direction (two thick grey arrows in Scheme 2).^[31] The pyrH⁺/PhO⁻xy_n-Ru⁺ intermediate (at 1.3 eV) produced after initial CPET escapes detection because it decays more rapidly than it is formed.^[34] In other words, CPET is the rate-determining step leading to the observable $pyrH^+/PhO^--xy_n-Ru^{2+}$ photoproducts. This view is consistent with previously reported rates exceeding 10⁹ s⁻¹ for "simple" electron transfer over three adjacent *p*-phenylene units as well as with the relative sluggishness of CPET reactions in related phenol-Ru(bpy)₃³⁺ and phenolrhenium(I) systems.^[14,23,24,35-38] Some of the H/D kinetic isotope effects reported below provide additional support for the conclusion that CPET is rate-determining.

Kinetics of the photoreaction in presence of pyrrolidine

In order for the photoinduced CPET reaction to occur, pyrrolidine must be in immediate proximity to the phenolic units because the proton cannot tunnel very far.^[39] It is therefore reasonable to expect CPET to occur predominantly from the hydrogen-bonded encounter adducts on the right-hand side of the chemical equilibrium in Equation (4), in analogy to prior studies of PCET with hydrogen-bonded phenols.^[22, 31, 39-46]

$$pyr+PhOH-xy_n-Ru^{2+} \rightleftharpoons pyr \cdots PhOH-xy_n-Ru^{2+}$$
(4)

Any analysis of the CPET kinetics must take this equilibrium into account. The CPET kinetics of the three dyads can be measured by monitoring the transient absorption signal at 450 nm as a function of time after excitation at 532 nm with laser pulses of 10 ns width. Exemplary bleach recoveries for the PhOH-xy₁-Ru²⁺ dyad are shown in Figure 2, all others are reported in Figure S3 in the Supporting Information. Immediately after the laser pulses one detects a bleach at 450 nm due to the disappearance of the ¹MLCT absorption as a consequence of Ru(bpy)₃²⁺ excitation.^[47] In absence of pyrrolidine the bleach recovery time is about 200 ns in aerated CH₃CN for all three dyads, corresponding to relaxation of the ³MLCT state to the ground state like in free $Ru(bpy)_3^{2+}$. In presence of pyrrolidine the bleach recovery becomes significantly more rapid (Figure 2, Figure S3 in the Supporting Information) due to the sequence of photoreactions (CPET, back-ET) discussed in the previous paragraph. Since the CPET step is rate determining,



Figure 2. Temporal evolution of the transient absorption signal of PhOH-xy₁- Ru^{2+} (2×10⁻⁵ M) in aerated CH₃CN at 450 nm after excitation at 532 nm with pulses of ca. 10 ns duration. Pyrrolidine concentrations are given in the inset.

these bleach recovery times essentially reflect the kinetics of the CPET reaction.

The observed rate constants (k_{obs} , extracted from single exponential fits to the data in Figure 2 and Figure S3 in the Supporting Information) are plotted as a function of pyrrolidine concentration in Figure 3. Squares, circles, and triangles are used to visually set apart data for dyads with 1, 2, and 3 *p*-xylene spacers; open symbols represent data for ordinary dyads, filled symbols represent data obtained for their deuterated analogs in presence of deuterated pyrrolidine. In this context "deuterated" means replacement of the phenolic proton



Figure 3. Experimentally observable ³MLCT-depopulation rate constants (k_{obs}) as a function of pyrrolidine concentration for the three dyads and their deuterated analogues. The actual experimental data are shown in Figure 2 and Figure S3 in the Supporting Information. The solid lines are fits to the experimental data with Equation (5) as described in the text. Results from these fits are reported in Table 3.

by a deuteron and replacement of the easily exchangeable N– H proton of pyrrolidine by a deuteron. The most important observation is that upon bridge lengthening increasing pyrrolidine concentrations are necessary to induce similar reaction kinetics. This is a manifestation of the fact that the CPET reaction rate decreases with increasing bridge length.

The experimentally determined bleach recovery rate constants (k_{obs}) are principally a function of the intrinsic ³MLCT ex-

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cited-state decay rate constant (k_0) and the rate constant for CPET (k_{CPET}). Equation (5) further contains a rate constant for "simple" electron transfer (k_{ET}) and a k_Q [base] term describing direct ³MLCT excited-state quenching by the base.^[22,40,41]

$$k_{obs} = k_0 + k_{ET} + k_Q[base] + k_{CPET} \frac{K_A[base]}{1 + K_A[base]}$$
(5)

As discussed above, in absence of pyrrolidine the ³MLCT lifetime of the three dyads in aerated CH₃CN is within experimental accuracy the same as that of isolated $Ru(bpy)_3^{2+}$ hence we conclude that $k_{\rm FT}$ is negligible compared to k_0 . Addition of up to 1 M pyrrolidine to a 2×10^{-5} M solution of Ru(bpy)₃²⁺ in aerated CH₃CN has no influence on the ³MLCT emission and its lifetime (Figure S2 in the Supporting Information). From this we conclude that $k_Q \le 10^5 \text{ M}^{-1} \text{ s}^{-1}$. Fits with Equation (5) to the experimental data in Figure 3 are therefore possible with only two adjustable parameters: The CPET rate constant (k_{CPET}) and the association constant (K_A) describing the chemical equilibrium in Equation (4). We attempted to determine K_A independently on the basis of steady-state luminescence data as described previously (Figure S4 and S5 in the Supporting Information) but have not been able to extract meaningful K_A values from this procedure (see the Supporting Information).^[48]

Two-parameter fits to single sets of data from Figure 3 turned out to be not meaningful,^[49] and therefore we decided to fit all six data sets globally with the restriction to one common K_A value for all six cases. In other words, we assume that the equilibrium in Equation (4) is affected neither by the number of *p*-xylene spacers nor by deuteration. Our fits employed $k_0 = 5.2 \times 10^6 \text{ s}^{-1}$, $k_{\text{ET}} = 0 \text{ s}^{-1}$, and $k_Q = 10^5 \text{ M}^{-1} \text{ s}^{-1}$ as non-adjustable parameters and produced the k_{CPET} values reported in the third column of Table 3 and $K_A = 1.4 \pm 0.2 \text{ M}^{-1}$. The fits are shown as solid lines in Figure 3. The quality of these fits supports the CPET hypothesis based on the chemical equilibrium in Equation (4) and speaks against any stepwise proton transfer, electron transfer mechanisms.

A value of $1.4 \pm 0.2 \text{ M}^{-1}$ for K_A is in line with previously reported equilibrium constants of hydrogen-bonded phenols in benzonitrile.^[40,41] The H/D kinetic isotope effect (KIE) of 3.91 ± 0.95 (ratio between k_{CPET} of PhOH-xy₂-Ru²⁺ and PhOD-xy₂-Ru²⁺) found for the dyad with two *p*-xylene bridging units is significant and supports the hypothesis of a rate-determining CPET step. The two other dyads exhibit smaller KIEs $(1.26 \pm 0.40, xy_1$ system) and $(0.95 \pm 0.44, xy_3$ system), but the absence of a sig-

Table 3. CPET rate constants (k_{CPET}) as a function of phenol–ruthenium (center-to-center) distance (R_{DA}).					
Molecule	<i>R</i> _{DA} [Å]	$k_{CPET} [s^{-1}]$			
PhOH- xy_1 -Ru ²⁺	12.2	$(1.07\pm0.14) \times 10^{9}$			
PhOD- xy_1 -Ru ²⁺	12.2	$(0.85\pm0.12) \times 10^{9}$			
PhOH- xy_2 -Ru ²⁺	16.5	$(3.35\pm0.41) \times 10^{8}$			
PhOD- xy_2 -Ru ²⁺	16.5	$(0.86\pm0.10) \times 10^{8}$			
PhOH-xy ₃ -Ru ²⁺	20.8	(3.41±1.22)×10°			
PhOD-xy ₃ -Ru ²⁺	20.8	(3.58±1.14)×10 ⁶			

nificant KIE is not an argument against CPET.^[50] Trends in the magnitudes of KIEs associated with PCETs are generally difficult to rationalize because the KIE depends on many different parameters.^[11]

The distance dependence of the CPET rate constants is shown in Figure 4 in semilogarithmic representation; R_{DA} corresponds to center-to-center distances between the phenol and Ru(bpy)₃²⁺ moieties as obtained by molecular modeling. Linear



Figure 4. Distance dependence of k_{CPET} The solid lines are linear regression fits yielding the β -values reported in the text.

regression fits to the data in Figure 4 yields distance decay constants (β) of 0.67 ± 0.23 Å⁻¹ and 0.64 ± 0.06 Å⁻¹ for the PhOH-xy_n-Ru²⁺ and PhOD-xy_n-Ru²⁺ series, respectively. These β values are much in line with those extracted for "simple" (i.e., not proton-coupled) electron tunneling across oligo-*p*-xylene spacers (0.77 Å⁻¹, 0.52 Å⁻¹)^[51,52] and oligo-*p*-phenylene bridges.^[35,36,53,54] Extrapolation of these exponential fits to van der Waals contact distance between phenol and Ru(bpy)₃²⁺ reactants (R_{DA} =4 Å) yields k_{CPET} =4.5×10¹¹ s⁻¹ for ordinary phenol and k_{CPET} =1.8×10¹¹ s⁻¹ for deuterated phenol. These rates compare favorably to those of "simple" electron transfer between reactants in van der Waals contact.^[1,10]

Conclusion

The distance dependence of the rates for bidirectional CPET in the systems investigated here does not deviate drastically from that of ordinary (i.e., not proton-coupled) electron transfer. The dominant contribution to this distance dependence likely comes from $H_{AB,PCET}$ [Eq. (1)], which can be described as a product of electronic coupling matrix elements for the electron transfer ($H_{AB,ET}$) and proton transfer components ($H_{AB,PT}$) of the overall CPET reaction.[11,39] Increasing the electron transfer distance decreases $H_{AB,ET}$ whereas $H_{AB,PT}$ remains constant. From this perspective the observed similarity between the distance dependences of bidirectional CPET ($\beta = 0.67 \pm 0.23$ Å⁻¹) and ordinary electron transfer ($\beta = 0.52 - 0.77 \text{ Å}^{-1}$)^[51,52,55] across the same type of bridges is readily understandable. Changes in reorganization energy (λ_{PCET} [Eq. (1)]) with increasing electron donor/electron acceptor distance appear to be relatively minor. It is possible that reorganization energy effects are re-

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sponsible for the deviation from the exponential distance dependence of CPET rates observed here.^[12]

Since this is the first determination of β for a bidirectional CPET reaction it is delicate to make a general statement, but it seems that the dependence of rates for bidirectional CPET on electron donor/electron acceptor distance is not steeper than that of ordinary electron transfer rates, despite the concerted motion of oppositely charged particles into different directions. This finding is of key importance when attempting to perform light-driven separation of electrons and protons in order to build up charge gradients across natural or artificial membranes, similar to what occurs in photosystem II. We find that a long electron transfer distance (up to ca. 20 Å) is no obstacle to bidirectional CPET.

Experimental Section

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¹H NMR spectra were measured on a 400 MHz Bruker Avance instrument. The ¹H spectra were referenced relative to tetramethylsilane by using the solvent signals as internal standards. High-resolution mass spectra were recorded on a Finnigan MAT8200 mass spectrometer. Elemental analysis was performed using a Vario EL III CHNS analyzer from Elementar. For cyclic voltammetry a Versastat3-200 potentiostat from Princeton Applied Research was used, and a glassy carbon disk served as a working electrode. A silver wire was used as a counter-electrode, and a second silver wire was employed as a quasi-reference electrode. Proper referencing of the potentials occurred through addition of small amounts of ferrocene and by recording of the ferrocenium/ferrocene redox wave. Voltage sweeps were performed with rates of 100 mVs⁻¹. Optical absorption spectra were recorded on a Shimadzu UV-1800 instrument. Steady-state luminescence was measured on a Fluorolog322 instrument from Horiba Jobin-Yvon. Transient absorption was performed with an LP920-KS spectrometer from Edinburgh Instruments, equipped with an R928 photomultiplier and an iCCD camera from Andor. The frequency-doubled output from a Quantel Brilliant b laser was used as an excitation source. The duration of the laser excitation pulses was approximately 10 ns. Transient absorption spectra were generally recorded by time-averaging the signal over a detection period of 200 ns. Quartz cuvettes from Starna were employed for all optical spectroscopic experiments. For H/D kinetic isotope effect studies, the deuterated dyads were dissolved in pure CH₃CN and deuterated pyrrolidine was added. Synthetic protocols and product characterization data are given in the Supporting Information.

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