Solvent and Temperature Effects on Photoinduced Proton-Coupled Electron Transfer in the Marcus Inverted Region

Laura F. Cotter, Belinda Pettersson Rimgard, Giovanny A. Parada, James M. Mayer,* and Leif Hammarström*

**ABSTRACT:** Concerted proton-coupled electron transfer (PCET) in the Marcus inverted region was recently demonstrated (Science 2019, 364, 471–475). Understanding the requirements for such reactivity is fundamentally important and holds promise as a design principle for solar energy conversion systems. Herein, we investigate the solvent polarity and temperature dependence of photoinduced proton-coupled charge separation (CS) and charge recombination (CR) in anthracene–phenol–pyridine triads: 1 (10-(4-hydroxy-3-(4-methylpyridin-2-yl)benzyl)anthracene-9-carbonitrile) and 2 (10-(4-hydroxy-3-(4-methoxypyridin-2-yl)benzyl)anthracene-9-carbonitrile). Both the CS and CR rate constants increased with increasing polarity in acetonitrile:n-butynitrile mixtures. The kinetics were semi-quantitatively analyzed where changes in dielectric and refractive index, and thus consequently changes in driving force ($-\Delta G^0$) and reorganization energy ($\lambda$), were accounted for. The results were further validated by fitting the temperature dependence, from 180 to 298 K, in $n$-butynitrile. The analyses support previous computational work where transitions to proton vibrational excited states dominate the CR reaction with a distinct activation free energy ($\Delta G^0_{\text{CR}} \sim 140$ meV). However, the solvent continuum model fails to accurately describe the changes in $\Delta G^0$ and $\lambda$ with temperature via changes in dielectric constant and refractive index. Satisfactory modeling was obtained using the results of a molecular solvent model [J. Phys. Chem. B 1999, 103, 9130–9140], which predicts that $\lambda$ decreases with temperature, opposite to that of the continuum model. To further assess the solvent polarity control in the inverted region, the reactions were studied in toluene. Nonpolar solvents decrease both $\Delta G^0_{\text{CR}}$ and $\lambda$, slowing CR into the nanosecond time regime for 2 in toluene at 298 K. This demonstrates how PCET in the inverted region may be controlled to potentially use proton-coupled CS states for efficient solar fuel production and photooxid catalysis.

**INTRODUCTION**

The thermochemistry and kinetics of electron transfer (ET) and proton transfer (PT) are often intimately correlated in processes known as proton-coupled electron transfer (PCET) reactions.1–10 These associated electron–proton transfer reactions are critical to numerous fundamental energy conversion processes, from photosynthesis and respiration to combustion and fuel cells. Such processes may become even more favorable when high-energy intermediates can be bypassed via a concerted mechanism, where PT and ET occur in a single kinetic step (CPET).

Marcus theory, in its most archetypical form, predicts ET rate constants based on the reaction free-energy barrier ($\Delta G^0$), which depends on the intrinsic reorganization energy ($\lambda$) and the reaction driving force ($-\Delta G^0$):11

$$\Delta G^0 = \frac{(\Delta G^0 + \lambda)^2}{4\lambda}$$  (1)

Subsequent efforts by Levich, Jortner, Marcus, and others led to the development of a quantum mechanical description of ET reactions and rates.12–14 In the high-temperature limit, all solvent and solute modes can be treated classically, and one obtains the following expression for the nonadiabatic rate constant:

$$k_{\text{ET}} = \frac{|V_{el}|^2}{\hbar} \sqrt{\frac{\pi}{2k_B T}} \exp \left[ -\frac{(\Delta G^0 + \lambda)^2}{4\lambda k_B T} \right]$$  (2)

where $V_{el}$ is the electronic coupling between the reactants and products, $k_B$ is the Boltzmann constant, and $T$ is the temperature. As evident in eq 2, a remarkable prediction of Marcus theory is the bell-shaped free-energy dependence. Due to the quadratic relationship between $k_{\text{ET}}$ and the driving force, the rate constant reaches a maximum when $-\Delta G^0 = \lambda$ and then, counterintuitively, proceeds to decrease with a further...
Scheme 1. Structures of Anthracene—Phenol—Pyridine Triads 1—8 and Reaction Scheme for Photoinduced e–/H+ Charge Separation (CS) and CR

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increase in driving force. This regime, where $-\Delta G^\circ > \lambda$, is known as the Marcus inverted region (MIR).

Development of theories for ET and PT has been combined and extended to describe PCET reactions, treating the proton quantum mechanically, for a review of this development, see ref 3. The following expression has been obtained for nonadiabatic CPET in the high-temperature limit:

$$k_{\text{CPET}} = \sum_{\mu} P_\mu \sum_{\nu} \frac{|V_{\mu\nu}|^2 S_{\mu\nu}(R_{\text{PT}})|^2}{\hbar} \sqrt{\frac{\pi}{\lambda k_B T}} \exp \left( -\frac{\Delta G_{\mu\nu}^\circ + \lambda}{4\lambda k_B T} \right)$$

where the summations are over transitions between different proton vibrational states of the reactants ($\mu$) and products ($\nu$) weighted by the Boltzmann population ($P_\mu$) of a given reactant proton vibrational state. The coupling can be approximated by the electronic coupling, $V_{\mu\nu}$, multiplied by the proton vibrational wavefunction overlap between reactant and product states ($S_{\mu\nu}$). Thermal fluctuations result in a distribution of proton donor–acceptor distances, typically on the scale of 0.1–0.3 Å. Thus, $S_{\mu\nu}$ is an integral over various PT distances ($R_{\text{PT}}$).

Decades after the formulation of the Marcus theory, the first widely accepted experimental evidence of ET reactions in the MIR was obtained in the mid-1980s by Closs and Miller and co-workers for ground-state (GS) charge shift reactions. This was closely followed by a report of inverted ET in photoinduced charge recombination (CR) reactions by Wasielewski and co-workers. Specific details of the rate versus free-energy correlations in the MIR indicated the involvement of nuclear tunneling of medium-frequency modes (typically aromatic C–C vibrations). Contributions from such medium-frequency modes attenuate the inverted region effect, leading to a shallower decrease of $k_{\text{ET}}$ with increasing driving force than would be predicted from eq 2. A similar effect can be expected for CPET reactions due to contributions from high-frequency proton vibrationally excited states. The large electron–proton coupling, that is, the large shift in the equilibrium nuclear distance between reactants and products, was proposed to make nuclear tunneling to higher states even more important for CPET compared to the typical case of ET. Accordingly, it was originally predicted that the inverted region behavior would not be observed for CPET, even for extremely exergonic reactions.

In contrast to these predictions, we recently reported CPET reactions exhibiting the inverted region behavior within a series of anthracene–phenol–pyridine (An-PhOH-py) molecular triads 1–8 (Scheme 1). The series was designed to vary $\Delta G^\circ$ for photoinduced, proton-coupled CS and CR by substitution effects on the anthracene (electron acceptor) and pyridine (proton acceptor) moieties. Light excitation of the anthracene unit triggers ET from phenol to anthracene, concerted with PT of the phenolic proton to pyridine. The CS rate constant ($k_{\text{CS}}$) increases with increasing driving force as expected for a reaction in the normal region (eqs 2 and 3, $-\Delta G^\circ < \lambda$). The subsequent CR reaction for 1–3 reforms the GS reactants in a CPET reaction in the MIR ($-\Delta G^\circ > \lambda$). The CR rate constants ($k_{\text{CR}}$) decrease with increasing driving force within the series ($1 < 2 < 3$) as well as when the solvent polarity decreases from dimethylformamide (DMF, $\epsilon_r = 38.25$) to dichloromethane (DCM, $\epsilon_r = 8.93$). The strong solvent dependence is indicative of an inverted reaction since the effects of decreasing polarity on $\Delta G^\circ$ and $\lambda$ of a CR inverted reaction both result in shifting the reaction deeper into the MIR. Moreover, in spite of the earlier general predictions (see above), the observed inverted CR rates were satisfactorily modeled using the theory of Hammes-Schiffer and co-workers (eq 3), with contributions from proton vibrational excited states in the electronic GS products. The modeling used a description of the proton potentials as double wells that was more accurate in this case than the anharmonic or Morse potentials used for the previous general predictions.

In the original Marcus model for ET, the inverted region stems from the increase in the activation barrier with...
increasing driving force once \(-\Delta G^\circ > \lambda\). Quantum mechanical models provide an alternative view of the MIR, where activationless reactions to high product vibrational levels have a smaller vibrational wavefunction overlap when \(-\Delta G^\circ\) becomes larger. This is analogous to the origin of the energy gap law for nonradiative transitions. This would lead to the prediction of an inverted region ET rate constant that is independent, or nearly independent, of temperature. Theoretical modeling of the CR reactions in 1–3 in DCM according to eq 3 instead suggested that the dominant contributing reactant-to-product vibronic transitions (mainly around \(\mu = 0 \to \nu = 3\)) indeed have a significant barrier (\(\Delta G^\circ \approx 90\) meV in DCM), with \(-\Delta G_{\text{cr}}^\circ\) larger than \(\lambda\) (see the Supporting Information in ref 31). The barrierless transition \((0 \to 7)\) with \(-\Delta G_{\text{cr}}^\circ \approx \lambda\), on the other hand, has a negligible proton vibrational wavefunction overlap and therefore its contribution to \(k_{\text{CPET-CR}}\) is negligible. This explains why the MIR effect was observed for the CPET reactions of the An-PhOH-py triads.

Advancing the understanding of the conditions and parameters that allow for the MIR behavior in CPET reactions is of fundamental and general interest. Photoinduced CPET reactions in the MIR could enable the design of more efficient technologies for harvesting solar energy. This is particularly important for processes relying on the formation of charge-separated states (CSS) as energy-storing transient species, many of which implicate PCET steps, such as in photosynthesis. These highly reactive species are susceptible to energy-wasting CR reactions, which must be slow enough to allow for the productive, fuel-producing reactions to dominate. MIR kinetics, as proposed by Marcus and others, is thought to fulfill this function and is a fundamental principle operating in the primary CS reactions of photosynthesis. Although this hypothesis is widely accepted, it is rarely verified in natural systems. Therefore, it is important to understand what features allow for the MIR behavior and whether the MIR is a general phenomenon in CPET or if the An-PhOH-py triads are simply an exception to the rule.

The study presented here was designed to test and extend our understanding of inverted CPET in this system of An-PhOH-py triads. It focuses on the effects of temperature and solvent polarity on the photoinduced CPET reactivity of triads 1 and 2 using UV-vis femtosecond transient absorption (TA) spectroscopy. (Triad 3 also shows CR of the CSS with inverted region kinetics; however, it was excluded from the present studies due to its negligible solubility in nitrile solvents and toluene.) By using mixtures of similar solvents n-butylformamide (PrCN) and acetonitrile (MeCN), the solvent polarity was systematically varied to study the effects of modulating \(\Delta G^\circ\) and \(\lambda\) of the reactions. Temperature-dependent experiments in PrCN probed the influences of \(\Delta G^\circ\), \(\lambda\), Marcus barriers, and thermal state populations. Experiments in toluene (Tol) were conducted to extend the investigation to a nonpolar solvent, which should maximize the inverted region effect for CR by minimizing the reorganization energy and maximizing the driving force.

These studies give further support for and insight into the MIR behavior of the An-PhOH-py triads. The results herein demonstrate that long-lived (1H+/1e−) CSSs can be achieved via slow MIR recombination kinetics and therefore such states could, in principle, participate in follow-up chemical reactions. This proof of principle could therefore be of assistance in developing solar to chemical energy conversion schemes.

### EXPERIMENTAL DETAILS

Transient spectroscopy was performed with a 3 kHz 800 nm output of a Ti:sapphire amplifier (1.5 mJ, 45 fs FWHM, Libra, Coherent), which was split into pump (35%) and probe (65%). To achieve the 400 nm pump, 800 nm light was frequency-doubled using a 0.2 mm-thick BBO crystal (EKSMA Optics) prior to being chopped (1.5 kHz) in the sample compartment (Newport TAS). To avoid major effects of rotational depolarization, the pump was made pseudo-unpolarized using a depolarizer (Thorlabs). A white light supercontinuum probe was generated by focusing the light onto a 4 mm CaF2 crystal after passing through an 8 ns optical delay stage (Newport TAS). The probe spectra were recorded using a custom-made 200–1000 nm silicon diode array (Newport). Triads 1 and 2 from previous studies were dissolved in PrCN/MeCN (Sigma-Aldrich/Merck, >99.0% (GC)/spec. grade) mixtures, prepared in 1 mm × 10 mm quartz cuvettes, with an absorption of ∼0.1–0.15 at 400 nm measured using a Varian Cary 50 or 5000. The pump intensity was attenuated to 150 μW, and for each mixture, three scans were collected and averaged using 1000 ms integration time. Experiments with toluene (Tol, Merck, spectroscopic grade) followed the same procedure.

For temperature-dependent experiments in PrCN (Sigma-Aldrich, >99.0% (GC)), the solvent was dried overnight over molecular sieves (3 Å, 8–12 mesh, Sigma-Aldrich) and later filtered using Acrodisc 2 mm syringe filters (0.45 μm, WWPTEF membrane). The temperature was controlled using an Optistat DN1704 cryostat (Oxford Instruments NanoScience) with an ITC 501 controller. The cryostat was equipped with a 0.8 mm-thick heated chamber (Newport TAS). To avoid major effects of temperature and solvent polarity on the photoinduced CPET reactivity of triads 1 and 2, using UV–vis femtosecond transient absorption (TA) spectroscopy, (Triad 3 also shows CR of the CSS with inverted region kinetics; however, it was excluded from the present studies due to its negligible solubility in nitrile solvents and toluene.) By using mixtures of similar solvents n-butylformamide (PrCN) and acetonitrile (MeCN), the solvent polarity was systematically varied to study the effects of modulating \(\Delta G^\circ\) and \(\lambda\) of the reactions. Temperature-dependent experiments in PrCN probed the influences of \(\Delta G^\circ\), \(\lambda\), Marcus barriers, and thermal state populations. Experiments in toluene (Tol) were conducted to extend the investigation to a nonpolar solvent, which should maximize the inverted region effect for CR by minimizing the reorganization energy and maximizing the driving force.

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### RESULTS

Overview of Triad Properties. Substituent effects on anthracene and phenyl groups of triads 1–8 vary the driving forces for CPET by ∼0.9 eV for CS and by ∼1.1 eV for CR in DCM. Specific structural features of the triads include a strong intramolecular hydrogen bond between phenol and pyridine as well as a methylene spacer to keep the anthracene and phenol–pyridine moieties electronically distinct. The X-ray structures of 1, 3, 5, and 6 confirm that the anthracene and phenol–pyridine units lie out-of-plane relative to each other,
and that the structure is sufficiently rigid to prevent intramolecular π–π stacking.\textsuperscript{31,43}

Figure 1 shows the steady-state absorption and fluorescence spectra of 1 measured in MeCN,\textsuperscript{31} which is representative of those of triad 2. The reference spectra of the individual subunits in the same solvent are also included, specifically the absorption spectra of 2,4-di-tert-butyl-6-(pyridin-2-yl)phenol (PhOH-py) and the absorption and fluorescence spectra of 9-cyano-10-methylnanthracene (9-CN-10-Me-An). The absorption spectrum of 1 indicates that the phenol–pyridine and anthracene subunits are weakly coupled as it agrees well with the sum of the spectra of the two subunits. The fluorescence spectra of 1 and those of 9-CN-10-Me-An are similar, but for the triad, the fluorescence yield is more than 1000 times smaller due to emission quenching by the phenol–pyridine unit.\textsuperscript{43} The absorption and emission of the 0 → 0 transition in anthracene overlap significantly in the spectra. Hence, the excited-state energy ($E_{0}−E_{\text{GS}}$ which approximates $\Delta G^{0}$ of the locally excited state (LES) relative to the GS) can be estimated as 2.97 eV from the average wavenumber of the two 0 → 0 transition maxima. For 1 in PrCN, it was estimated that $\Delta G^{0}\text{CS} \sim −0.49$ eV and $\Delta G^{0}\text{CR} \sim −2.48$ eV, with the corresponding values for 2 being $\sim −0.54$ eV and $\sim −2.43$ eV, respectively.\textsuperscript{31}

TA in PrCN at Room Temperature. Figure 2A–D shows UV–vis TA data of 1 and 2 in PrCN at 298 K after fs excitation at 400 nm and results of global fits (E,F). (A,B): TA spectra at different times after excitation (gray) and fitted spectra (in color); (C,D): TA time traces at selected wavelengths (gray circles) and fitted traces (in color); and (E,F): evolution-associated spectra (EAS) from global analysis with a sum of three exponential components. The resulting time constants and assignment of the process are listed in Table 2. Residual plots from the fits are shown in the Supporting Information.

UV–vis and mid-IR range.\textsuperscript{31} The CSS features disappear on a time scale of $\sim 100$ ps, leaving no significant TA signal after 1 ns.

A global fit of the TA data to a sum of exponential decays requires three time components to obtain satisfactory fits. For 1, the time constants are $\tau_{1} = 2.5$ ps, $\tau_{2} = 6.1$ ps, and $\tau_{3} = 88$ ps, while for 2, the time constants are slightly smaller (Table 1). The resulting EAS for each component represent the absorption spectrum of the species corresponding to that time constant (note that the anthracene ground-state bleach is outside the spectral window shown in Figure 2). The EAS of the first two components each show SE (negative bands at

Table 1. Time Constants$^{a,b}$ and Assignments$^{c}$ from Global Fits to the TA Data in PrCN between 180 and 298 K

| time constants$^{a,b}$ | triad 1 | triad 2 |
|------------------------|---------|---------|
| $\tau_{1}$ (ps)        | 2.5     | 2.5     |
| $\tau_{2}$ (ps)        | 6.1     | 6.1     |
| $\tau_{3}$ (ps)        | 88      | 88      |

$^{a}$Rate constants correspond to $k_{\text{CS}} = 1/\tau_{1}$ and $k_{\text{CR}} = 1/\tau_{3}$ (see text).

$^{b}$Standard deviations of the time constants are estimated at ±5%. VR = relaxation of LES$_{\text{hot}}$ CS$_{\text{hot}}$ = CS from LES$_{\text{hot}}$ CS = charge separation from a relaxed LES; and CR = charge recombination.
450–500 nm) and ESA (a positive band with maxima around 570 nm). Both spectral features are attributed to \(^1\text{An}\).\(^{31}\) The SE is red-shifted while the ESA is blue-shifted in the second component relative to the first one (Figure 2E,F). However, the second component additionally contains characteristic features of the CSS: broad absorption with peaks around 620 and 680 nm attributed to the \(\text{An}^*\) radical and a narrower band around 425 nm attributed to the phenoxy radical.\(^{31}\) Thus, we conclude that the first component corresponds to the “hot” local \(^1\text{An}\) state (LES\(_{\text{hot}}\)) undergoing CPET to the CSS with \(\tau_1 = 2.5\) ps in parallel with thermal relaxation. Thermal relaxation of “hot” excited states in solution is nonexponential,\(^{44}\) but given the competing CS reaction, the data do not justify a more complicated kinetic model. The relaxed LES then undergoes CS to form the CSS with \(\tau_2 = 6.1\) ps. This explains the slight shift of SE and ESA between the first and second components and the biexponential generation of CSS. The EAS of the third component shows only the spectroscopic features of the CSS, which decays monotonically by CR to reform the GS with \(\tau_3 = 88\) ps. The EAS \(\text{An}^*\) features above 600 nm blueshift slightly from the second to third component indicating that the initial CPET from the LES\(_{\text{hot}}\) generates hot CSS (CSS\(_{\text{hot}}\)) to some extent. In contrast, the narrow band around 425 nm, distinctive of \(\text{PhO}^*\), remains unshifted indicating that the excess thermal energy resides on the \(\text{An}^*\) moiety. Relaxation of CSS\(_{\text{hot}}\) could not be resolved, but its expected TA changes are very small compared to those resulting from the CEPT from the relaxed LES. Therefore, we believe that the effect of the CSS\(_{\text{hot}}\) relaxation on \(\tau_3\) is within the experimental and analytical errors. Triads 1 and 2 show qualitatively the same behavior (Figure 2). All time constants and their assignments are listed in Table 1.

The rate constants are calculated as \(k_{\text{CS}} = 1/\tau_3\) and \(k_{\text{CR}} = 1/\tau_1\) because in PrCN, there are no apparent contributions of other decay pathways to the time constants for thermalized CS and CR (as opposed to the behavior in Tol, vide infra). This is further justified by the observation that CPET from the relaxed LES is about three orders of magnitude faster than excited-state decay in 9-CN-10-Me-An (\(\tau = 17\) ns).\(^{31}\) The above results are very close for 1 and 2 in PrCN in our previous study (\(\tau_{\text{CS}} = 5.2\) ps and 4.6 ps for 1 and 2, respectively)\(^{31}\) but the “hot” species are resolved much better in the present data. Note that the data in the previous study were fitted with two components, and their re-evaluation using three component shows no change to the reported values in DCM or DMF. The small underestimation of \(\tau_{\text{CS}}\) in PrCN in our previous study, however, does not affect its discussion and conclusions.

**Temperature-Dependent Experiments.** The temperature dependence of CPET CS and CR for 1 and 2 was studied between 298 and 180 K in PrCN, thus avoiding glass formation. Going from 298 to 180 K, the spectral features remained the same. We observe a weak temperature dependence for CS and CR over the 120 K interval examined with ca. sixfold decrease in \(k_{\text{CS}}\) and a mere twofold decrease in \(k_{\text{CR}}\) as the temperature decreases. For both 1 and 2, the Arrhenius plot (eq 4) shows a good linear relationship for CS (Figure 3A) with an apparent activation energy, \(E_a\) of \(\sim 70\) meV (\(\sim 1.6\) kcal mol\(^{-1}\)) for both triads. In contrast, the Arrhenius plot for CR is substantially curved (Figure 3B), and linear fits give a crude \(E_a\) estimate of \(\sim 17\) meV for 1 (\(\sim 0.4\) kcal mol\(^{-1}\)) and \(\sim 23\) meV for 2 (\(\sim 0.5\) kcal mol\(^{-1}\)). \(k_{\text{CR}}\) is essentially constant in the higher end of the studied temperature range, from 240 to 298 K. This is opposite to the expected behavior for a reaction with tunneling activation, where temperature-independent tunneling dominates at low temperatures and the reaction is activated only at higher temperatures. This suggests that the Arrhenius model is, perhaps unsurprisingly, inappropriate to analyze these reactions. Alternative analyses are presented in the Discussion section.

\[
\ln k = \ln A - \frac{E_a}{RT} \tag{4}
\]

**TA Experiments in Nitrile Mixtures.** TA experiments were conducted in MeCN/PrCN solvent mixtures to investigate the effect of solvent polarity on the free-energy barriers (\(\Delta G^*\)). The use of MeCN/PrCN solvent mixtures allowed us to systematically vary the static dielectric constant (\(\varepsilon\)) and refractive index (\(n\)) from those of pure MeCN (\(\varepsilon\,(293\text{K}) = 36.64\) and \(n\,(293\text{K}) = 1.3414\)) to those of pure PrCN (\(\varepsilon\,(293\text{K}) = 24.83\) and \(n\,(293\text{K}) = 1.3842\)).\(^{32}\) In addition to having similar specific solute–solvent interactions, nitrile mixtures were selected due to the linear correlation between the solvent mole fraction and dielectric properties as evidenced by the linear shift in absorption \(\lambda_{\text{max}}\) of the betaine \(E_2(30)\) probe (see the Supporting Information).

All the TA spectral features for 1 and 2 in nitrile mixtures are the same as described above at 298 K. For both 1 and 2, \(k_{\text{CS}}\) and \(k_{\text{CR}}\) systematically increase as the solvent polarity is increased from that of neat PrCN to MeCN (Figure 4, Table 2, and Figures S15–S26). This is illustrated in Figure 4A–D where time traces at a selected wavelength have been chosen to emphasize the kinetic differences in neat solvents for CS and CR. Traces at 455 nm show an initially negative TA signal due to SE from LES\(_{\text{hot}}\) that is converted to a positive absorption from the CSS and finally followed by decay of the latter to restore the GS. Traces at 540 nm are at an isosbestic point between the LES and CSS and thereby monitor only the CR process. \(k_{\text{CS}}\) shows a modest increase by 30–60% from neat PrCN to neat MeCN. In contrast, \(k_{\text{CR}}\) shows a larger effect with 2.5–3 times larger rates in MeCN than in PrCN for both triads.

The observed trend of the dependence of \(k_{\text{CS}}\) and \(k_{\text{CR}}\) on \(\varepsilon\) and \(n\) is consistent with expectations based on the Marcus outer reorganization energies and driving forces (eqs 5–8). In Marcus theory,\(^{11,45}\) \(\Delta G^*\) depends on \(\Delta G^0\) and \(\lambda\) according to eq 1. The same dependence is found in CPET theory (eq 3), where \(\Delta G^0\) for each vibronic transition is given. The total reorganization energy for ET and CPET is the sum of inner-sphere (\(\lambda_{\text{in}}\)) and outer-sphere (\(\lambda_{\text{out}}\)) contributions (eq 5).

\[
\lambda = \lambda_{\text{in}} + \lambda_{\text{out}} \tag{5}
\]

where \(\lambda_{\text{in}}\) is related to changes in bond lengths and angles, while \(\lambda_{\text{out}}\) is due to solvent polarization changes between the...
reactant and product states. By applying a dielectric continuum model and assuming spherical reactants, the following approximation can be made:11,45

$$\lambda_{\text{out}} = \frac{e^2}{4\pi\varepsilon_0} \left( \frac{1}{2a} + \frac{1}{2b} - \frac{1}{R} \right) \left( \frac{1}{n^2} - \frac{1}{\varepsilon_i} \right)$$

where $e$ is the elementary charge, $\varepsilon_0$ is the vacuum permittivity, $a$ and $b$ are the radii of the donor and the acceptor, respectively, and $R$ is the distance between their centers. The last parenthetical term in eq 6 describes the dependence of $\lambda_{\text{out}}$ on the solvent $\varepsilon_i$ and $n$. $\lambda_{\text{out}}$ is larger in MeCN than in PrCN due to the lower $n$ and higher $\varepsilon_i$ of the former (eq 6). At the same time, the zwitterionic CSS is energetically stabilized as predicted by eqs 7 and 8:

$$\Delta G^0_{\text{CS}} = \Delta G^0_{\text{CS,ref}} + \frac{e^2}{4\pi\varepsilon_0} \left( \frac{1}{2a} + \frac{1}{2b} - \frac{1}{R} \right) \left( \frac{1}{\varepsilon_i} - \frac{1}{\varepsilon_{i,\text{ref}}} \right)$$

$$\Delta G^0_{\text{CR}} = \Delta G^0_{\text{CR,ref}} - \frac{e^2}{4\pi\varepsilon_0} \left( \frac{1}{2a} + \frac{1}{2b} - \frac{1}{R} \right) \left( \frac{1}{\varepsilon_i} - \frac{1}{\varepsilon_{i,\text{ref}}} \right)$$

where the subscript "ref" refers to any reference solvent used for comparison of $\Delta G^0$. For CPET reactions like in 1 and 2, the driving force $-\Delta G^0_{\text{CS}}$ and $\lambda$ will then increase, thereby having a counter-balancing effect on $\Delta G^*$, which is consistent with the observed small changes in $k_{\text{CS}}$ for both triads.

For CR on the other hand, increasing the MeCN mole fraction in the mixtures makes $\Delta G^0_{\text{CR}}$ less negative (eq 8) while $\lambda$ increases. This decreases $\Delta G^*$ and makes the reaction less inverted. Therefore, the large solvent effect observed for CR is consistent with the inverted region behavior since changes in $\Delta G^*$ and $\lambda$ act in the same direction. A quantitative analysis based on this continuum model is used to investigate the variation of $k_{\text{CS}}$ and $k_{\text{CR}}$ in the MeCN/PrCN mixtures (see the Discussion section).

**TA Experiments in Tol.** The CSS should be more destabilized in the very low-polarity solvent Tol ($\varepsilon_{i,\text{(298K)}} = 2.38$) compared to PrCN. Despite its higher energy, the CSS is observed for 1 and 2, and its assignment can be done as previously described based on the absorption bands at 425 nm (PhO•) and at 625 and 670 nm (An•+) (Figure 5). Notably, the CSS is more long-lived in Tol than in more polar solvents: $\tau_{\text{CS}} = 140$ ps for 1 and 2.5 ns for 2 at 298 K. The destabilization of the CSS should result in CR driving forces that are higher than those in PrCN ($\Delta G^0_{\text{CR,Tol}} < -2.48$ eV for 1 and $<-2.54$ eV for 2), making the observation of a 2.5 ns lifetime remarkable. The vibronic coupling is strong enough to give $k_{\text{CR}} = (27 ps)^{-1}$ in the polar solvent MeCN. This clear

Table 2. Time Constants $a, b$ and Assignments $c$ from Global Fits to the TA Data in MeCN/PrCN Solvent Mixtures and Their Respective Solvent Parameters

| $X_{\text{MeCN}}$ | $\varepsilon_i$ | $n^*$ | $t_1$ (ps) | $t_2$ (ps) | $t_3$ (ps) | $t_4$ (ps) | $t_5$ (ps) | $t_6$ (ps) |
|-------------------|-----------------|-------|------------|------------|------------|------------|------------|------------|
| 1.00              | 36.64           | 1.3456| 1.6        | 3.7        | 32         | 0.3        | 3.1        | 27         |
| 0.87              | 35.10           | 1.3506| 2.3        | 4.1        | 39         | 0.4        | 3.4        | 31         |
| 0.71              | 33.27           | 1.3566| 1.8        | 4.2        | 45         | 0.2        | 3.4        | 38         |
| 0.53              | 31.05           | 1.3639| 1.2        | 4.4        | 56         | 0.7        | 4.1        | 48         |
| 0.29              | 28.30           | 1.3728| 1.5        | 4.1        | 75         | 2.6        | 5.4        | 60         |
| 0.00              | 24.83           | 1.3842| 1.0        | 4.8        | 100        | 2.0        | 4.9        | 70         |

$^a$Rate constants correspond to $k_{\text{CS}} = 1/t_1$ and $k_{\text{CR}} = 1/t_4$ (see text). $^b$Standard deviations of the time constants are estimated at ±5%. $^c$VR = relaxation of LES$_{\text{relaxed}}$, CS$_{\text{hot}}$ = CS from LES$_{\text{relaxed}}$, CS = charge separation from a relaxed LES, and CR = charge recombination. $^d$Mole fraction of MeCN in MeCN/PrCN mixtures. $^e$Values of $\varepsilon_i$ and $n$ were assumed to vary linearly with the mole fraction.
and 2 component 2 τ τ small and spectrally indistinct TA changes, making the EAS of the TA experiments (the decay of the CSS and does not decay on the time scale of absorption band at 435 nm, which appears simultaneously with information.

Normalized EAS from the fitting of the TA data in Tol at 298 K. The second component, with a better agreement with that of triplet anthracene (3*An). The time constants resulting from the global fit and respective assignments are listed in Table 3. Except for the formation of 3*An upon CR, 1 and 2 follow the same reactions described using PrCN. One differentiating factor between 1 and 2 in Tol is that while τ1 for 1 represents VR of the initial LES state, CS in 2 seems to be completed during the first component τ1 = 12 ps, representing CS hot as determined from the EAS (Figure S30). Therefore, CS is assigned to τ2 for 1 and τ1 for 2, while CR is assigned to τ1 for both triads. The EAS for both τ2 and τ1 in 2 show the CSS (bands at 425, 625, and 670 mm). The second component, with τ2 = 103 ps, shows only small and spectrally indistinct TA changes, making the EAS of the second and third components very similar. The second component is tentatively assigned to CSS relaxation (although τ2 appears to be quite long to support the assignment).

Formation yields of the CSS (ΦCSS) and 3*An (ΦT) were estimated relative to the LES, using the EAS amplitudes and extinction coefficients of the species involved (Table 4 and details in the Supporting Information). The ΦCSS values were estimated at ~26% for 1 and ~90% for 2. Assuming similar extinction coefficients in PrCN and Tol, the ΦCSS value for 2 was as high in Tol as in PrCN and within the experimental uncertainty (cf. Figures 4B and 2B). This is in striking contrast to the much lower yield for 1 in Tol versus PrCN (cf. Figures 4A and 2A). The ΦT values were estimated at ~0.5% for 1 and ~6.8% for 2. Thus, the CSS mostly undergoes CR to reform the GS (as in nitrile solvents) with only a minor fraction forming 3*An, presumably via intersystem crossing in the CSS state followed by CR to 3*An (see the Discussion section).

■ DISCUSSION

Our previous study31 allowed the assignment of the LES → CSS conversion in triads 1–3 to a concerted PCET mechanism (CPET) on the basis of the observed kinetic evolution in the TA and thermochemical and kinetic estimates for the different PCET mechanisms. Thus, in a single kinetic step, one e− transfers from phenol to the excited anthracene moiety, while H+ transfers from phenol to pyridine. This assignment was first based on the simultaneous formation of the spectral features of PhO• and An• as well as the observed KIE (1.7 ± 0.2), both of which were in agreement with the CPET mechanism. Second, stepwise proton transfer followed by electron transfer (PTET) and electron transfer followed by proton transfer (ETPT) were excluded by considering that the PT step of PTET should be significantly uphill (ΔG° > 10, for phenol and pyridinium in MeCN2), which would not allow for the observed rate constant of ~1011 s−1. The initial ET seemed inefficient, emphasized by the use of a reference triad where the phenolic hydroxyl group was replaced by a methoxy group, which did not show any quenching of 3*An.43 Finally, the free-energy dependence of the observed kCSS for the CSS formation in 1−8 varied with ΔG°CPET, in agreement with theoretical predictions for CPET (eq 3) but not with ETPT or PTET.

In PrCN, the CSS spectroscopic features of PhO• and An• showed simultaneous and monotonic decay to the GS, thus closing a simple three-state photochemical cycle.14 An analysis of this CR based on thermochemical and kinetic estimates for the different PCET mechanisms, analogous to that for CS, allowed us to assign the mechanism to CPET. As described in the Introduction, the CPET CR displayed an inverted region behavior, both in a comparison of 1−3 in the same solvent and for each compound when the solvent polarity was varied. The prior computational modeling suggested that vibronic transitions around 0 → 3 dominate this reaction for triad 1 in DCM with an activation barrier ΔG°0,3 = 90 meV. A table of the relative contribution of the most important vibronic transitions and their parameters is provided in the Supporting Information (Table S2).

In this section, we will analyze and discuss the reaction barriers for CS and CR, using the data for the MeCN/PrCN
mixtures and the temperature-dependent data in PrCN, and provide further experimental evidence of the inverted region character of CR. In particular, we explore whether the explanation for the inverted region behavior, such as being dominated by transitions to higher vibrational product states (the $0 \rightarrow 3$ vibronic transition when in DCM), is supported by experimental estimates of the effective $\Delta G^*$ and $-\Delta G^\circ$ values. The latter values should in turn reflect a weighted average of the vibronic transitions contributing to each reaction. Therefore, we will discuss the results in the nonpolar solvent TOL, which show that the CR can indeed be pushed further into the inverted region to give nanosecond lifetimes of the CSS.

Marcus-Type Modeling—Motivation for Fits with One Averaged Transition. As a first approximation to model the solvent polarity and temperature dependencies of CPET, we use a Marcus-type fit (eq 2) for the nitrile mixture and temperature-dependent data in PrCN. An analysis using eq 3 is not feasible here as it contains too many unknown parameters to allow for a fit to the kinetic data. The Marcus-type fit represents a gross simplification for a CPET reaction as eq 3 includes additional temperature-dependent factors beyond the classical barrier $\Delta G^*$ for the $0 \rightarrow 0$ transition. First, the Boltzmann population of proton vibrational states of the reactant ($P_\nu$) can give a temperature dependence. For the present CR reactions, however, computational results showed that only transitions from $\mu = 0$ were important because of the relatively high energy of $\mu = 1$ and above. Second, the distribution of PT distances ($R_{PT}$) increases in width with increasing temperature, and this is expected to add to the experimentally observed activation energy. Calculations indicate that the dominant proton donor–acceptor distance is close to the equilibrium distance, thereby suggesting a minimal temperature dependence of $R_{PT}$. Third, because multiple vibronic transitions are predicted to contribute to the rate, each with a different $\Delta G^\circ_{\mu \nu}$, the observed effective $\Delta G^*$ is a weighted average of these transitions. In an effort to assess the latter effect, we have adapted the previous computational results for CR in DCM (Table S2) to PrCN by adding 60 meV to $\Delta G^\circ_{\mu \nu}$ in the more polar PrCN and using a value of $\lambda = 1.3$ eV in PrCN (Table S, see reasoning below and the Supporting Information for details on the value for $\lambda$). In Table S, it is assumed that all CSS energies are 60 meV lower than those in the less polar solvent DCM, relative to the GS, but that the proton vibrational wavefunction overlaps ($S^\nu_{\mu \nu}$) remain unchanged. Table S therefore shows which vibronic transitions contribute most to $k_{CR}$ and gives the values of their respective energetic and overlap factors.

The following sections show how a qualitative and even semi-quantitative agreement with the previous theoretical modeling can be obtained from a semi-classical Marcus-type fit of the temperature dependence of $k_{CR}$ in which the manifold of vibronic transitions in eq 3 is represented by a weighted average according to eq 2, yielding an averaged value of $\Delta G^\circ_{\mu \nu}$ from the fits. This value is compared with $\Delta G^\circ_{\mu \nu}$ of the calculated vibronic transitions that are predicted to give the largest contribution to $k_{CR}$ (Table S). The effective $\Delta G^*$ and $\Delta G^\circ$ values obtained from the fits to the temperature-dependent and MeCN/PrCN data are in good agreement with each other and correspond to the computationally predicted most important transitions based on the values in Table S.

### Table 5. Computational Data for the Vibronic Transitions That Are Predicted To Give the Main Contributions to the CR Rate Constant for 1 in PrCN at 298 K, cf. Eq 3

| $(\nu, \lambda)$ | $P_\nu$ | $\Delta G^\circ_{\mu \nu}$ | $\Delta G^\circ_{\mu \nu}$ | $S^\nu_{\mu \nu}$ | $\exp(-\Delta G^\circ_{\mu \nu}/RT)$ | % contrib |
|-----------------|--------|-----------------|-----------------|-------------|-------------------------------|---------|
| 0–0             | 1.00   | -2.48 0.27      | $1.03 \times 10^{-3}$ | 3.18 $10^{-1}$ | 0.00                          |         |
| 0–1             | 1.00   | -2.26 0.18      | $4.53 \times 10^{-1}$ | 9.49 $10^{-4}$ | 9.63                          |         |
| 0–2             | 1.00   | -2.20 0.16      | $2.42 \times 10^{-1}$ | 2.23 $10^{-1}$ | 21.19                         |         |
| 0–3             | 1.00   | -2.06 0.11      | $1.03 \times 10^{-1}$ | 1.39 $10^{-1}$ | 31.95                         |         |
| 0–4             | 1.00   | -1.90 0.07      | $1.73 \times 10^{-2}$ | 7.02 $10^{-2}$ | 27.21                         |         |
| 0–5             | 1.00   | -1.71 0.03      | $1.45 \times 10^{-3}$ | 2.83 $10^{-3}$ | 9.20                          |         |
| 0–6             | 1.00   | -1.51 0.01      | $4.99 \times 10^{-5}$ | 7.28 $10^{-5}$ | 0.81                          |         |
| 0–7             | 1.00   | -1.28 0.00      | $4.78 \times 10^{-8}$ | 9.98 $10^{-8}$ | 0.00                          |         |
| 0–8             | 1.00   | -1.05 0.01      | $1.41 \times 10^{-7}$ | 6.24 $10^{-7}$ | 0.00                          |         |

The data of Table 5 were therefore used to calculate $\Delta G^*$ and gives the values of their respective energetic and overlap factors. The values of $\Delta G^*$ for each solvent composition were calculated from the values of $\Delta G^\circ$ and $\lambda$ in PrCN and their estimated variation with the solvent. The variation in $\lambda$ was estimated using the expressions for charged spheres in a continuum (eqs 5 and 6), assuming that $\epsilon_s, \epsilon_r$ and $n$ vary linearly with the mole fraction of the solvents. The driving forces for CS and CR were also assumed to vary linearly with the solvent mole fraction. The difference in $\Delta G^\circ$ for CS and CR between the neat solvents was previously calculated to be 60 meV, which is in fair agreement with predictions from eqs 7 and 8 (30 meV). In ref 31, $-\Delta G^\circ_{CS}$ in PrCN was calculated to be $\sim 0.49$ eV and $\approx 0.54$ eV for 1 and 2, respectively. A value of $\lambda$, similarly estimated by calculations, was used; see next paragraph. The CS data were fitted by using these predetermined $\Delta G^\circ_{CS}$ and $\lambda$ values in PrCN as input parameters for the analysis. Eqs 5 and 6 were then used to calculate $\lambda$ in MeCN, as described above, and the value of $\Delta G^\circ_{CS}$ in MeCN was varied until a good fit with a slope equal to $-\lambda (RT)^{-1}$ was obtained. The CR data were fitted independently following the same procedure, with the same value of $\lambda$ as for CS. The resulting fits according to eq 3 are shown in Fig 5.
9a were good, as shown in Figure 6. Note that CR and CS are expected to have very similar λ values considering the small structural rearrangements between the GS and LES of An, as inferred by the small Stokes shift of the 0→0 lines in the absorption and fluorescence spectra (Figure 2).

For the choice of λ in the fits, we used the value of λ = 1.4 eV previously calculated in DCM for λ as a reference value for which λ = 0.38 eV (cf. eq 5). From eqs 5 and 6, we can expect a value of λ CS = 1.5 eV in the more polar solvent PrCN. The results for CS with λ = 1.5 eV however gave unphysically large pre-exponential factors (∼1 × 10^-14 s^-1; Table 6), whereas a value of λ = 1.3 eV gave quite reasonable results. The latter value also agreed with the results of the temperature-dependent analysis, see below, and was therefore preferred. The difference in ΔG° CS between the neat solvents for 1 (70 meV) was in good agreement with the previous calculations above, while the difference was slightly larger (110 meV) for 2. The more negative ΔG° CS in MeCN was counterbalanced by a larger λ, such that there was ≤30% variation in ΔG° CS between the two neat solvents. The pre-exponential factor obtained (C in eq 9a) was reasonable and corresponds to a Marcus pre-exponential factor in eq 2 of B = (2 ± 1) × 10^13 s^-1 for 1 and 2 (Table 6).

In PrCN, the ΔG° CR value for the 0→0 transition was calculated to −2.48 and −2.43 eV for 1 and 2, respectively. A fit according to eq 9a required even more negative values in MeCN: ΔG° CR = −2.51 and −2.53 eV for 1 and 2. These results were unreasonable as the more polar MeCN should further stabilize the CSS compared to PrCN. In addition, this would imply that the combined equilibrium driving forces for CS and CR (−ΔG° CS − ΔG° CR) would exceed the excited-state energy (E° ex = 2.97 eV) of the LES. Hence, the results seem to indicate an overestimation of −ΔG° CR for the CR and its most contributing vibronic transitions. Instead, Table 5 suggests that the transitions giving the main contributions to the rate constant in PrCN are centered around the 0→3 transition, which has a much lower driving force than the 0,0 transition: ΔG° 0,3 = −2.06 eV. Acceptable linear fits for CR with λ = 1.3 eV, resulting in a slope of (−RT)−1, were in fact obtained with ΔG° CR/PrCN ≈ −2.06 eV for 1 and −2.01 for 2 (Figure 6 and see the Supporting Information for discussion of fits). This supports our previous computational results, suggesting CR to higher vibrational states with a CR thermal barrier (ΔG°*). It should be noted that the averaged ΔG° CR is smaller for 2 than 1, which can be explained by the somewhat lower energy of the CSS in 2, making the CR less inverted. The difference in ΔG° CR between the dyads is, however, still smaller than the difference in ΔG° CS, which is consistent with the general notion of a shallower driving force dependence of the rate constant in the inverted region.

The difference in effective ΔG° CR between neat PrCN and MeCN is negligible, which can be understood from the fact that as the CSS is stabilized in the more polar solvent, transitions to lower vibrational states become more important. Table S4, constructed on the basis of Table S5, highlights the different vibronic contributions in the nitrile mixtures, where ΔG° CR/MeCN is near that of the 0→2 transition (see the Supporting Information for description). It is evident however that the reason for the considerable change in ΔG° CR and observed kCR between solvents (ranging from DCM to DMF in ref 31) cannot be traced back to solely changes in ΔG° CR but rather the significant change in λ.31

To conclude, the nitrile mixture rate constants can be satisfactorily modeled with eq 9a with values of λ and ΔG° that are close to those calculated for λ and ΔG° CR of the dominating vibronic transitions. For CR, ΔG° CR is clearly less negative than for the 0→0 transition, and by comparison with calculations from ref 31, it represents an effective average around the 0→3 transitions in the nitrile mixtures. The much stronger increase in the CR rates with increased mole fraction of MeCN, compared to the CS, is consistent with a combined effect of an increase in λ and a less negative ΔG°, making the reaction less inverted. This observation and the fit results support the assignment of the CR occurring in the inverted region with a significant effective barrier of ΔG°* ~1.0 eV in PrCN, as computed in ref 31.

CEPT Temperature Dependence in PrCN. Rate constants for CS and CR are smaller for 1 than 2 over the temperature interval examined (Table 1). This is consistent with the ca. 50 meV higher energy of the CSS for 1 that results in a smaller driving force in the normal region CS and a larger driving force in the inverted region CR.

### Table 6. Resulting Parameters for the Neat Solvents from a Fit According to Eq 9a to the Kinetic Data in MeCN/PrCN Mixtures

| triad | λ/eV | solvent | ΔG° CS/eV | ΔG° CS/eV | B CR/s^-1 | ΔG° CS/eV | ΔG° CS/eV | B CR/s^-1 |
|-------|------|---------|-----------|-----------|-----------|-----------|-----------|-----------|
| 1     | 1.3  | PrCN    | −0.49     | 0.13      | 3.1 × 10^14 | −2.06     | 0.11      | 8.0 × 10^11 |
|       |      | MeCN    | −0.56     | 0.12      | 3.1 × 10^14 | −2.06     | 0.08      | 8.0 × 10^11 |
| 1     | 1.5  | PrCN    | −0.49     | 0.17      | 1.7 × 10^14 | −2.17     | 0.17      | 1.8 × 10^11 |
|       |      | MeCN    | −0.57     | 0.16      | 1.7 × 10^14 | −2.14     | 0.05      | 1.8 × 10^11 |
| 2     | 1.3  | PrCN    | −0.54     | 0.11      | 1.5 × 10^14 | −2.01     | 0.07      | 1.6 × 10^11 |
|       |      | MeCN    | −0.65     | 0.10      | 1.5 × 10^14 | −2.01     | 0.07      | 1.6 × 10^11 |
| 1     | 1.5  | PrCN    | −0.54     | 0.15      | 8.0 × 10^13 | −2.12     | 0.06      | 1.6 × 10^11 |
|       |      | MeCN    | −0.66     | 0.14      | 8.0 × 10^13 | −2.09     | 0.04      | 1.6 × 10^11 |

Figure 6. Kinetic data and fits according to eq 9a for CS (A) and CR (B) in 1 and 2 in MeCN/PrCN mixtures. λ = 1.3 eV for both reactions (see Table 6).

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Despite a temperature change by almost 120 K, from 180 to 298 K, the variation in the CS and CR rate constants for both 1 and 2 in PrCN is small (Table 1). A previous study of the CS reaction for the parent triad 6, using fluorescence measurements, found a very small temperature dependence for CS between 145 and 350 K in 2-methyltetrahydrofuran (Me-THF) with an Arrhenius activation barrier of only ∼14 meV (0.33 kcal mol⁻¹). A similar result may be expected for CR in the inverted region, based on prior observations for ET systems. ET reactions in the inverted region are strongly affected by nuclear tunneling to excited vibrational states of the products. This not only decreases the classically expected falloff of rate with further increase of $-\Delta G^\circ$ but it may also be expected to result in a weak temperature dependence, as observed experimentally.

Classical and semi-classical Marcus expressions provide a foundation to analyze the temperature dependence of ET reactions (cf. eq 2). Typically, a plot of $k_{ET}$ versus $1/T$, or $\ln k_{ET}$ versus $1/T$ (to account for the weak temperature dependence of the pre-exponential factor), is assumed to yield a straight line with a slope equal to $-\Delta G^\circ/R$. In a corresponding analysis of our CPET data, we find that $\ln k_{CS}$ correlates linearly with $1/T$ (cf. Figure 3). In contrast, neither $\ln k_{CR}$ nor $\ln k_{ET}/T^{1/2}$ versus $1/T$ yields linear correlations, and a clear downward curvature is observed instead (Figure 7).

The above approach may be justified over a small temperature range, which is suitable when $\Delta G^\circ$ is large. For large ranges of temperature, the variation of $\Delta G^\circ$ and $\lambda$ with temperature may be significant, which is attributed to the reaction entropy and reorganization entropy, respectively. The CSS is a zwitterionic species that polarizes the solvent, resulting in a smaller entropy than for the charge neutral LES and GS. Corrections for the temperature dependence of $\Delta G^\circ$ and $\lambda$ in ET reactions have in many cases been made by using the temperature dependence of $n$ and $\varepsilon_i$ and the continuum models of eqs 5–8.

The dielectric properties of PrCN change substantially upon cooling from 298 to 180 K: $n$ increases from 1.38 to 1.44 and $\varepsilon_i$ increases from 24.3 to 43.6 (see the Supporting Information for details). The much larger solvent polarity at lower $T$ stabilizes the CSS. However, according to eq 6, $\lambda$ also decreases as $T$ decreases because of the larger effect from the concomitant increase in $n$. The net result is that $\Delta G^\circ$ for CR is predicted to increase as $T$ is lowered. This is in contrast to the results from the studies of, for example, Liang et al. and Serpa et al. in nonpolar solvents Me-THF and isopropyl ether, respectively. In these solvents, eq 6 predicts that $\lambda$ increases as $T$ decreases because the increase of $\varepsilon_i$ is more important than that of $n$.

Here, the predicted changes of $\Delta G^\circ$ with temperature were calculated from eqs 5–8 and the temperature-dependent data for $\varepsilon_i$ and $n$ (see the Supporting Information). To obtain a linear plot of $\ln k$ versus $1/T$, a term was added in the fit, representing the change in barrier $\lambda/RT$ as the temperature was lowered from 298 to 180 K. In order to also correct for the small changes due to $\lambda/RT$ in the pre-exponential factor, the kinetic data were therefore plotted versus $1/T$ according to eq 10:

$$\ln(k \cdot \sqrt{T} \cdot \lambda) + \frac{\Delta(\Delta G^\circ)}{RT} = \ln D - \frac{\Delta G^\circ_{298 K}}{R} \frac{1}{T}$$  (10)

where $D$ encompasses the terms of the pre-exponential factor from eq 2 ($B = D \cdot (T \cdot \lambda)^{-1}$). The initial input values of $\lambda$ and $\Delta G^\circ$ at 298 K for an iterative optimization, guided by the calculations of Table S, were used to find a satisfactory linear fit to the data, with a resulting slope $(-\Delta G^\circ_{298 K}/R)$. Here, the iterative process of fitting according to eq 10 aimed to find input values of $\lambda_{298 K}$ and $\Delta G^\circ_{298 K}$ that gave a value of $\Delta G^\circ_{298 K}$ according to eq 1 that was identical to the value of $\Delta G^\circ_{298 K}$ returned by the fit.
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Table 7. Fit Results for 1 and 2 in PrCN at 298 K Using Eq 10 and MSM$^{a}$

| triad | T/K | $\lambda$/eV | $\Delta G^\circ_{CS}$/eV | $\Delta G^\circ_{CR}$/eV | $B_{CS}$/s$^{-1}$ | $\Delta G^\circ_{CS}$/eV | $\Delta G^\circ_{CR}$/eV | $B_{CR}$/s$^{-1}$ |
|-------|-----|-------------|-----------------|-----------------|-------------|-----------------|-----------------|-------------|
| 1     | 298 | 1.30        | −0.49           | 0.13            | 2.4 $\times 10^{15}$ | −2.15          | 0.14            | 5.1 $\times 10^{12}$ |
| 2     | 298 | 1.30        | −0.52           | 0.12            | 2.0 $\times 10^{15}$ | −2.19          | 0.15            | 8.8 $\times 10^{12}$ |

$^{a}$The range of values for acceptable fits (see text) are: ±0.05 eV ($\Delta G^\circ_{CS}$), ±0.2 eV ($\Delta G^\circ_{CR}$), ±0.01 eV ($\Delta G^\circ_{CS}$), ±0.04 eV ($\Delta G^\circ_{CR}$), ±0.1 ($\lambda$), ±1.0 ($B_{CS}$), and ±2.0 ($B_{CR}$), respectively; see the Supporting Information for details.

Satisfactory fits of the CS data were obtained with eq 10, but the CR data have a clear downward curvature (Figure 7). The downward curvature cannot be attributed to the failure of the implicit assumption of a single, averaged vibronic transition. Parameters of a full CPET model, which were neglected when assuming a single transition (cf. eqs 3 and 2; see discussion above), become more important with increasing temperature. In turn this would rather cause an upward curvature of the plots. This is verified by a simple summation of the rate contributions from the main vibronic transitions shown in Table 5 at different temperatures (Table S7). Instead, the downward curvature of the data plotted according to eq 10 (Figure 7, blue data points) suggests that the continuum solvent expressions in eqs 5–8 fail to properly correct the temperature dependence of $\Delta G^\circ_{CR}$ and $\lambda$.

Shortcomings of the continuum model (eqs 7 and 8) to accurately describe the temperature dependence of $\Delta G^\circ$ and $\lambda$ have been discussed by several authors.56–65 In the molecular solvent models (MSMs) developed by Matyushov et al.56 and Newton et al.,63 solvent density fluctuations are important components of $\lambda_{s0}$ but are obviously neglected in dielectric continuum solvent models (CSMs). In a study of particular relevance for the present work, Matyushov et al. applied MSM to interpret the temperature dependence of the CR of a donor–acceptor molecule in MeCN.56 The study found that the predicted decrease of $\lambda$ with increasing temperature reproduced the experimental temperature dependence, while the continuum model prediction of increase of $\lambda$ with increasing temperature failed to do so.

We believe that the situation is similar in the present system in which we study the CR of the CSS back to a charge neutral GS in a nitrile solvent. A full theoretical calculation according to the molecular model is far beyond the scope of the present study. Instead, we attempted a fit according to eq 10 but now calculating $\Delta(\Delta G^\circ)$ with the assumption that $\lambda$ decreased linearly with increasing temperature in a similar way as in ref 56. The magnitude of the temperature dependencies (slopes) $\Delta \lambda/\Delta T$ and $\Delta(\Delta G^\circ)/\Delta T$ will obviously depend on the details of the donor–acceptor molecule studied and will presumably be somewhat different for MeCN used in ref 56 and PrCN used here. We employed several different slopes of $\Delta G^\circ$ and $\lambda$, using those of ref 56 as the starting point. Ultimately, we only considered slopes that yielded values of $\Delta G^\circ_{CS}$, $\Delta G^\circ_{CR}$, $\lambda$ and the pre-exponential factor ($B$) at 298 K, which are physically reasonable. Thus, $\Delta G^\circ_{CS}$ and $\lambda$ were selected to be within 50 meV and 0.3 eV, respectively, of the calculated values from ref 31, with $-(\Delta G^\circ_{CS} + \Delta G^\circ_{CR}) \leq E_{S0}$ and the pre-exponential factor $B \leq 3 \times 10^{13}$ s$^{-1}$ (Table S9).

Remarkably, this procedure yields good linear correlations ($R^2 \geq 0.99$) with a combination of physically reasonable parameters for only a rather small range of slopes. The best fits have the same relative slopes of $\Delta G^\circ$ and $\lambda$ versus $T$ as those obtained in ref 56, with $\Delta G^\circ_{CR}$ and $\lambda$ decreasing by $\sim 100$ and $\sim 50$ meV, respectively, from 180 to 298 K, and $\Delta G^\circ_{CS}$ increasing (becoming less negative) by $\sim 100$ meV. These variations with temperature can be compared with the predictions from eqs 7 and 8, where $\Delta G^\circ_{CR}$ and $\lambda$ both change by $\sim 50$ meV but in opposite directions. The room-temperature parameter values obtained from the best fits for 1 and 2 are given in Table 7.

The results are in good agreement with the fits to the data from MeCN/PrCN mixtures at 298 K (Table 6) and are consistent with the calculations shown in Table 5. In particular, the CR evidently occurs predominantly in higher vibronic states of the electronic GS. The effective $\Delta G^\circ_{CR}$ obtained is very similar to that of the calculated 0 → 3 transition ($\sim 2.06$ eV) as shown in Table 5, which is the same effective $\Delta G^\circ_{CR}$ used in the fit of the MeCN/PrCN mixtures. The value of $\Delta G^\circ_{CR} = 0.14 \pm 0.04$ eV for 1 is in good agreement with the value of 0.11 eV from the fit to the MeCN/PrCN mixture data. Thus, we believe that the fitting model (eq 10 with a MSM correction), although simplified, is semi-quantitatively correct. The narrow parameter range of acceptable fits to the temperature-dependent rate constants is in good agreement with the fits of data from the MeCN/PrCN mixtures, and the corresponding calculated data shown in Table 5 support that notion.

The differences between the MSM and CSM models are less significant for the MeCN/PrCN mixture data. Although we used a mole fraction-averaged Pekar factor $(\frac{1}{\varepsilon_n} - \frac{1}{\varepsilon_s})$ in eq 6 to calculate $\lambda$ and $\Delta G^\circ$, the errors are presumably similar for the two nitrile solvents. It is also little doubtful that MeCN, with a larger $n$ and smaller $\varepsilon_s$ than PrCN, gives both a larger $\lambda$ and a more stabilized CSS. Therefore, we believe that the actual variation in $\Delta(\Delta G^\circ)$ in these solvent mixtures is similar to the one predicted in our fit to eq 9a.

The fits for the MSM give a much stronger variation in $\ln(k_{CS} \sqrt{T\lambda}) + \Delta(\Delta G^\circ)/RT$ with temperature than the CSM (Figure 7B,D). This is because $\Delta G^\circ_{CS}$ is predicted to be much larger with the MSM than with the CSM (e.g., for 1: $\sim 0.14$ vs $0.023$ eV at 298 K). On the other hand, the variation of $\ln(k_{CS} \sqrt{T\lambda}) + \Delta(\Delta G^\circ)/RT$ for the CS step is very similar for the MSM and CSM fits, which may seem surprising given the fact that $\lambda$ changes with temperature in opposite directions for the two models. The reason for the similar result is that $\Delta G^\circ_{CS}$ is predicted to be very similar and shows quite parallel changes with temperature for the two models. With decreasing temperature, $\lambda$ increases by ca. 40 meV in the MSM instead of decreasing by a similar amount in the CSM, but this is compensated by the fact that $\Delta G^\circ_{CS}$ decreases twice as much in the MSM. Thus, the term $\Delta G^\circ_{CS} + \lambda$ of the numerator in eq 1 is the same within $\sim 30$ meV in the two models. The greater difference between MSM and CSM for CR than for CS is not because the former is in the inverted region but because of the opposite dependencies of $\Delta G^\circ_{CS}$ and $\Delta G^\circ_{CR}$ on temperature.

To conclude, variable temperature experiments in PrCN confirm our previous observation of inverted CR and are consistent with calculations using the PCET theory. The
effective barrier for the CR, determined from the fits, is \( \Delta G^\ddagger_{9298K} = 0.14 \pm 0.04 \) eV, which is similar to the barrier previously suggested by calculations of the dominating vibronic transitions (Table 5).\(^{31}\) The value is also similar to that obtained in the fits to the data in MeCN/PrCN mixtures at 298 K, \( \Delta G^\ddagger_{CR} = 0.11 \) eV for 1. This supports our claim that CR occurs in the inverted region and that it has a distinct barrier for the classical coordinates, instead of occurring entirely via activationless nuclear tunneling.

**CEP in Toluene.** The differences in the interplay between \( \Delta G^\ddagger \) and \( \lambda \) in the normal and inverted regions become larger in nonpolar solvents. In analogy to previous discussion, in Tol, we observe the expected qualitative trends in CS and CR, that is, in Tol, a decrease of both \( -\Delta G^\ddagger \) and \( \lambda \) is expected for CS (maintaining a counter-balancing effect on \( \Delta G^\ddagger \)), whereas an increase in \( -\Delta G^\ddagger \) and a decrease in \( \lambda \) is expected for CR, thereby pushing the reaction further into the inverted region. In agreement with these trends, there are larger differences in \( k_{CS} \) versus \( k_{CR} \) in Tol compared to the more polar solvents (cf. Tables 3 and 2, respectively). \( (k_{CR})^{-1} \) is on the order of 10–30 ps in Tol, similar to the values observed in PrCN and MeCN, while CR becomes significantly slower, so that the CSS decays back to the GS with \( \tau \sim 2.5 \) ns for 2.

Reliable estimates of \( \Delta G^\ddagger \) and \( \lambda \) for CS and CR in nonpolar solvents are typically challenging.\(^{66-70}\) Estimates of \( \lambda_{out} \) become negligibly small because \( 1/\varepsilon \) and \( 1/\nu^2 \) become equally large (eq 6). Estimates of \( \Delta G^\ddagger \) using eqs 7 and 8 predict that the CSS is destabilized by \( \sim 0.9 \) eV in Tol compared to PrCN; this is obviously exaggerated as it would make the CS endergonic by \( \sim 0.4 \) eV. Similar cases of rapid ET (ps time scale) in Tol have been reported before but where continuum models predict endergonic CS.\(^{66-70}\) Here, we can only conclude that \( \Delta G^\ddagger_{CS} \) in Tol must be somewhere between \( -0.06 \) eV (because the conversion of LES to CSS is at least 90% complete) and \( -0.49 \) eV in the stabilizing polar solvent PrCN. Estimates of \( \lambda_{out} \) using eqs 5 and 6 would predict a value of \( \sim 0 \) eV and therefore \( \lambda \approx \lambda_{in} \approx 0.38 \) eV (see above). The MSR predict a larger \( \lambda_{out} \) in nonpolar solvents; however, particularly \( \lambda_{out} \approx 0.1-0.3 \) eV,\(^{76} \) suggesting a value of \( \lambda = 0.5-0.7 \) eV. Thus, it seems that \( \lambda > -\Delta G^\ddagger_{CS} \) in our case, that is, CS is in the normal region. Assuming that the pre-exponential factor for CS in 1, where the \( 0 \rightarrow 0 \) transition would dominate, it would be similar in Tol and PrCN (\( B = 3 \times 10^{13} \) s\(^{-1} \)), and the observed \( k_{CS,Tol} = 1/\tau_2 \approx 3 \times 10^{10} \) s\(^{-1} \) (Table 3) would be consistent with a barrier of \( \Delta G^\ddagger_{CS} \approx 0.15 \) eV in Tol at room temperature. This estimate is in agreement with the suggested range of values above, for example, \( \Delta G^\ddagger_{CS,Tol} = -0.06 \) eV and \( \lambda = 0.7 \) eV, which would produce a barrier of ca. 0.15 eV (eq 1). These estimates point to a CSS energy for 1 in Tol around 0.1–0.2 eV below the singlet LES (2.97 eV), that is, \( \sim 2.8 \) eV relative to the GS.

Destabilization of the CSS, and the resulting long lifetime of the CSS, allows for the formation of a long-lived \( ^3 \)An state as the CSS recombines. \( ^3 \)An lies ca. 1.75 eV above the GS,\(^{72} \) such that \( \Delta G^\ddagger_{CR} \approx 3 \)An should be \( \sim -1.0 \) eV. With the estimated value of \( \lambda = 0.6-0.9 \) eV, this reaction should be in the inverted region, but less inverted than the singlet CR (\( \Delta G^\ddagger_{CR} \approx -2.8 \) eV). Nevertheless, the quantum yield for \( ^3 \)An formation is low, \( \sim 7\% \) for 2 and even smaller for 1, \( \sim 0.5\% \). For the latter, the \( ^3 \)An extinction coefficient at the absorption maximum (435 nm) is an order of magnitude larger than that for the CSS allowing for its observation in the TA spectra (Figure S30). Formation of \( ^3 \)An during CR requires that the CSS first undergoes intersystem crossing (ISC), that is, singlet–triplet conversion, most likely via hyperfine interactions as described for related radical ion pairs.\(^{47,74-77}\)

ISC in radical ion pairs of organic molecules typically occurs on the time scale of \( >1 \) ns,\(^{47} \) such that it can only compete with CR of the CSS when its lifetime approaches this time scale like in Tol for 1 and 2. A similar feature with a narrow 435 nm band at long delay times was observed in the low-polarity solvent DCM employed in our previous study where \( k_{CR} = (755 \) ps)\(^{-1} \).\(^{31} \) ISC should be slow on the time scale of CR and limit the yield of \( ^3 \)An formation. Indeed, the low yield of \( ^3 \)An shows that direct CR to the GS dominates the observed CR process, with the triplet pathway giving only a minor contribution to the rate.

The CSS formation yield of 2 in Tol, and of both triads in PrCN, is \( \Phi_{CSS} \geq 90\% \) based on literature extinction coefficients (see the Results section and the Supporting Information). The much lower \( \Phi_{CSS} \) for 1 in Tol (\( \sim 26\% \)) cannot be explained by direct deactivation to the GS as the LES lifetime of cyanoanthracene is three orders of magnitude longer than that in 1. This suggests that there is another parallel deactivation process in Tol, which is not active in 2 to the same extent and cannot be identified by the present TA experiments. We note that Sayfutyarova et al.\(^{75} \) have suggested, based on a computational study, direct formation of a local electron–proton transfer state, with a phenoxyl–pyridiniumyl biradicaloid subunit (An–PhO\(^{-}\)–pyH\(^{+}\)). Its potential experimental verification would require different experiments that are beyond the scope of the present study.

### CONCLUSIONS

In this study, the photoinduced PCET CS and CR rates of triads 1 and 2 were studied as a function of temperature and solvent polarity. Corrections for the temperature dependence of the reorganization energy and driving force, and the consequent change in the activation barrier with temperature, were attempted with both CSM and MSM.\(^{56} \) Satisfactory fits to the data could be obtained using a simplified model with a single vibronic transition (cf. eq 2) that represents a weighted average of the contributing vibronic transitions (cf. eq 3) and using MSM to correct for the temperature dependence of the activation barrier. The fit results suggested that CR had an effective (averaged) barrier of \( \Delta G^\ddagger_{9298K} = 0.14 \pm 0.04 \) eV and a reaction free energy of \( \Delta G^\ddagger_{CR,298K} \approx -2.1 \) eV for 1. The latter value is much less negative than that for CR between the lowest proton vibrational states of CSS and GS (\( \mu = 0 \)). The results match the predictions based on previous calculations\(^{31} \) that transitions to proton vibrationally excited states of the electronic GS (\( 0 \rightarrow 2 \) and \( 0 \rightarrow 3 \) in PrCN) dominate the CR reactivity. Consistent fits were obtained for the room-temperature data in the PrCN/MeCN mixtures. Here, the contributing vibronic transitions are slightly lower (\( 0 \rightarrow 1, 0 \rightarrow 2, \) and \( 0 \rightarrow 3 \)) because the more polar MeCN stabilizes the CSS. Nonetheless, the CR still forms a hot GS with a similar effective \( \Delta G^\ddagger \) as in the temperature study.

Both analyses of the temperature and nitrile mixture experiments resulted in pre-exponential factors for CS close to the theoretical limit of \( \leq 1 \times 10^{13} \) s\(^{-1} \) (from transition-state theory or dynamic fluorescence Stokes shift). The value for \( \Delta G^\ddagger_{CS} \) from the fit was equal to that for the \( 0 \rightarrow 0 \) transition. This is consistent with a reaction in the normal region, with a moderate barrier of \( \Delta G^\ddagger \approx 0.1 \) eV, for which the proton wavefunction overlap is good (\( S_0^2 \) not much smaller than
unity) and does not strongly limit the pre-exponential factor. For CR, the pre-exponential factor is 1–1.5 orders of magnitude smaller than that for CS. This is consistent with the computational results shown in Table 5 that transitions to higher proton vibrational states, for which $\Delta G^*_{\mu \nu}$ is small, are limited by a small wavefunction overlap and the transitions with the best wavefunction overlap have a higher $\Delta G^*_{\mu \nu}$ value. Thus, the effective average corresponds to a transition with a moderate barrier of $\Delta G^* \sim 0.14$ eV and a wavefunction overlap around $S^2_{\mu \nu} \sim 0.1$, which are approximately the values for the 0 $\rightarrow$ 3 transition. We note that the actual barrier is thus larger than that suggested by the weak temperature dependence of the experimental CR rate constant, which is a consequence of the concomitant variation in barrier with temperature.

The results support previous assignment of CR to a concerted PCET reaction in the inverted region.1 The model explains why an inverted region behavior is possible, thanks to a poor proton wavefunction overlap for the barrierless transitions. Computational studies have suggested that conditions for the inverted region behavior of CPET are dependent on asymmetric, double well proton potentials, while more symmetric potentials would not give an inverted region behavior.26,33 It is interesting to note that most computational studies of CPET oxidations of small-molecule phenol–base systems suggest highly asymmetric proton potentials.26,33 Nevertheless, an inverted region behavior has only been reported for the present series of triads. As discussed before,7 there are very few studies that report even a curvature in the rate versus free-energy correlation, as is suggested by eqs 2 and 3.7,17 It seems that the relation between proton potentials and rate versus free-energy dependence is not yet fully understood on a combined theoretical and experimental level.

In the nonpolar solvent Tol, CR occurs much more slowly with a time constant of $\sim 2.5$ ns for 2 at 298 K. This is a clear manifestation of the inverted region, where the expected decrease in $\lambda$ and a more negative $\Delta G^*_{CR}$ lead to an even larger reaction barrier. The very slow CR in Tol leads to distinctly different excited-state dynamics with other deactivation pathways competing with the inverted CR. These pathways include spin conversion/PCET to form the spectrally identified long-lived triplet anthracene state.

Overall, this report confirms and enriches our understanding of the PCET reactivity of two triads that undergo CR in the MIR. The results of this study have implications for future developments of reactions relying on the utilization of high-energy, proton-coupled CS states such as photocatalysis and solar fuel technologies.

ASSOCIATED CONTENT

1 Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acjpa.1c05764.

Additional TA data and fit results and details of the analysis and fits to temperature- and solvent polarity-dependent data (PDF)

AUTHOR INFORMATION

Corresponding Authors
James M. Mayer — Department of Chemistry, Yale University, New Haven, Connecticut 06520, United States; orcid.org/0000-0002-3943-5250; Email: james.mayer@yale.edu

Leif Hammarström — Department of Chemistry — Ångström Laboratory, Uppsala University, SE75120 Uppsala, Sweden; orcid.org/0000-0002-9933-9084; Email: leif.hammarstrom@kemi.uu.se

Authors
Laura F. Cotter — Department of Chemistry, Yale University, New Haven, Connecticut 06520, United States
Belinda Petersson Ringard — Department of Chemistry — Ångström Laboratory, Uppsala University, SE75120 Uppsala, Sweden
Giovan A. Parada — Department of Chemistry, Yale University, New Haven, Connecticut 06520, United States; Present Address: The College of New Jersey, Department of Chemistry, Ewing, New Jersey 08628, United States

Complete contact information is available at: https://pubs.acs.org/doi/10.1021/acjpa.1c05764

Author Contributions
§ L. F. C., B. P. R., and G. A. P. contributed equally to this work.

Notes
The authors declare no competing financial interest.

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REFERENCES

1. Huynh, M. H. V.; Meyer, T. J. Proton-Coupled Electron Transfer. Chem. Rev. 2007, 107, 5004–5064.
2. Warren, J. J.; Tronic, T. A.; Mayer, J. M. Thermochemistry of Proton-Coupled Electron Transfer Reagents and its Implications. Chem. Rev. 2010, 110, 6961–7001.
3. Migliore, A.; Polizzi, N. F.; Therien, M. J.; Beratan, D. N. Biochemistry and Theory of Proton-Coupled Electron Transfer. Chem. Rev. 2014, 114, 3381–3465.
4. Tryburski, R.; Liu, T.; Glover, S. D.; Hammarström, L. Proton-Coupled Electron Transfer Guidelines, Fair and Square. J. Am. Chem. Soc. 2021, 143, 560–576.
5. Hames-Schiffer, S. Proton-Coupled Electron Transfer: Moving Together and Charging Forward. J. Am. Chem. Soc. 2015, 137, 8860–8871.
6. Reece, S. Y.; Nocera, D. G. Proton-Coupled Electron Transfer in Biology: Results from Synergistic Studies in Natural and Model Systems. Annu. Rev. Biochem. 2009, 78, 673–699.
7. Costentin, C.; Savéant, J.-M. Theoretical and mechanistic aspects of proton-coupled electron transfer in electrochemistry. Curr. Opin. Electrochem. 2017, 1, 104–109.
8. Wengen, O. S. Proton-Coupled Electron Transfer with Photoexcited Metal Complexes. Acc. Chem. Res. 2013, 46, 1517–1526.
9. Lennox, J. C.; Kurtz, D. A.; Huang, T.; Dempsey, J. L. Excited-State Proton-Coupled Electron Transfer: Different Avenues for Promoting Proton/Electron Movement with Solar Photons. ACS Energy Lett. 2017, 2, 1246–1256.
10. Mora, S. J.; Odella, E.; Moore, G. F.; Gust, D.; Moore, T. A.; Moore, A. L. Proton-Coupled Electron Transfer in Artificial Photosynthetic Systems. Acc. Chem. Res. 2018, 51, 445–453.
11. Marcus, R.; Sutin, N. Electron transfers in chemistry and biology. Biochim. Biophys. Acta, Rev. Bioenerg. 1985, 811, 265–322.
12. Levich, V. Present state of the theory of oxidation-reduction in solution (bulk and electrode reactions). Adv. Electrochem. Electrochem. Eng. 1966, 4, 249–371.
1968.
752.
barriers, and Brønsted slopes of chemical reactions. J. Phys. Chem. 1968, 72, 891–899.
(16) Levich, V. G.; Dogonadze, R. R.; German, E. D.; Kuznetsov, A. M.; Kharksats, Y. I. Theory of homogeneous reactions involving proton transfer. Electrochim. Acta 1970, 15, 353–367.
(17) Borgis, D.; Hynes, J. T. Curve Crossing Formulation for Proton Transfer Reactions in Solution. J. Phys. Chem. 1996, 100, 1118–1128.
(18) Cukier, R. I. Proton-Coupled Electron Transfer through an Asymmetric Hydrogen-Bonded Interface. J. Phys. Chem. 1995, 99, 16101–16115.
(19) Soudackov, A.; Hammes-Schiffer, S. Multistate continuum theory for multiple charge transfer reactions in solution. J. Chem. Phys. 1999, 111, 4672–4687.
(20) Soudackov, A.; Hammes-Schiffer, S. Derivation of rate expressions for nonadiabatic proton-coupled electron transfer reactions in solution. J. Chem. Phys. 2000, 113, 2385–2396.
(21) Hammes-Schiffer, S.; Stuchebrukhov, A. A. Theory of Coupled Electron and Proton Transfer Reactions. Chem. Rev. 2010, 110, 6959–6960.
(22) Edwards, S. J.; Soudackov, A. V.; Hammes-Schiffer, S. Analysis of Kinetic Isotope Effects for Proton-Coupled Electron Transfer Reactions. J. Phys. Chem. A 2009, 113, 2117–2126.
(23) KrishTalk, L. I. The mechanism of the proton transfer: an outline. Biochim. Biophys. Acta. Bioenerg. 2000, 1458, 6–27.
(24) Kiefer, P. M.; Hynes, J. T. Kinetic Isotope Effects for Nonadiabatic Proton Transfer Reactions in a Polar Environment. I. Interpretation of Tunneling Kinetic Isotope Effects. J. Phys. Chem. A 2004, 108, 11793–11808.
(25) Markle, T. F.; Rhile, I. J.; Mayer, J. M. Kinetic Effects of Increased Proton Transfer Distance on Proton-Coupled Oxidations of Phenol-Amines. J. Am. Chem. Soc. 2011, 133, 17341–17352.
(26) Glover, S. D.; Parada, G. A.; Markle, T. F.; Ott, S.; Hammarström, L. Isolating the Effects of the Proton Tunneling Distance on Proton-Coupled Electron Transfer in a Series of Homologous Tyrosine-Base Model Compounds. J. Am. Chem. Soc. 2017, 139, 2090–2101.
(27) Miller, J. R.; Calcaterra, L. T.; Closs, G. L. Intramolecular long-distance electron transfer in radical anions. The effects of free energy and solvent on the reaction rates. J. Am. Chem. Soc. 1984, 106, 3047–3049.
(28) Closs, G. L.; Miller, J. R. Intramolecular Long-Distance Electron Transfer in Organic Molecules. Science 1988, 240, 440.
(29) Wasielewski, M. R.; Niemczyk, M. P.; Svec, W. A.; Pawiitwong, P.; Tsuchida, A.; Pettiworsh, E. B. Dependence of rate constants for photoinduced charge separation and dark charge recombination on the free energy of reaction in restricted-distance porphyrin-quinoxine molecules. J. Am. Chem. Soc. 1985, 107, 1080–1082.
(30) Hammes-Schiffer, S. Theory of Proton-Coupled Electron Transfer in Energy Conversion Processes. Acc. Chem. Res. 2009, 42, 1881–1889.
(31) Parada, G. A.; Goldsmith, Z. K.; Kolmar, S.; Pettersson Ringard, B.; Mercado, B. Q.; Hammarström, L.; Hammes-Schiffer, S.; Mayer, J. M. Converged proton-electron transfer reactions in the Marcus inverted region. Science 2019, 364, 471.
(32) CRC Handbook of Chemistry and Physics, Internet Version 2005. Lide, D. R., Ed. CRC Press: Boca Raton, FL, 2005.
(33) Goldsmith, Z. K.; Alexander, V. S.; Hammes-Schiffer, S. Theoretical analysis of the inverted region in photoinduced proton-coupled electron transfer. Faraday Discuss. 2019, 216, 363–378.
(34) Liang, N.; Miller, J. R.; Closs, G. L. Temperature-independent long-range electron transfer reactions in the Marcus inverted region. J. Am. Chem. Soc. 1990, 112, 5353–5354.
(35) Kroon, J.; Oevering, H.; Verhoeven, J. W.; Warman, J. M.; Oliver, A. M.; Paddon-Row, M. N. Temperature effects on intramolecular electron transfer kinetics under “normal”, “inverted”, and “nearly optimal” conditions. J. Phys. Chem. 1993, 97, 5065–5069.
(36) Smit, K. J.; Warman, J. M.; De Haas, M. P.; Paddon-Row, M. N.; Oliver, A. M. Charge recombination kinetics of giant dipoles in saturated hydrocarbon solvents. Chem. Phys. Lett. 1988, 152, 177–182.
(37) Marcus, R. A. Electron Transfer Reactions in Chemistry: Theory and Experiment (Nobel Lecture). Angew. Chem., Int. Ed. Engl. 1993, 32, 1111–1121.
(38) Makita, H.; Hastings, G. Inverted-region electron transfer as a mechanism for enhancing photosynthetic solar energy conversion efficiency. Proc. Natl. Acad. Sci. U. S. A. 2017, 114, 9267.
(39) Mulliken, R. K.; van Stokkum, I. H. M. TIMP: An R Package for Modeling Multi-way Spectroscopic Measurements. J. Stat. Softw. 2007, 18.
(40) Snellenburg, J. J.; Laptenok, S.; Seger, R.; Mulliken, K. M.; van Stokkum, I. H. M. Glotaran: A Java-Based Graphical User Interface for the R Package TIMP. J. Stat. Softw. 2012, 49.
(41) Petersson, J. Ultrafast, Non-Equilibrium Electron Transfer Reactions of Molecular Complexes in Solution. PhD Dissertation, Uppsala University, 2014.
(42) Föhlinger, J. Shining Light on Molecules: Electron Transfer Processes in Model Systems for Solar Energy Conversion Investigated by Transient Absorption Spectroscopy. Ph.D. Dissertation, Uppsala University, Uppsala, Sweden, 2018.
(43) Bowring, M. A.; Bradshaw, L. R.; Parada, G. A.; Pollock, T. P.; Fernández-Terán, R. J.; Kolmar, S. S.; Mercado, B. Q.; Schlenker, C. W.; Gamelin, D. R.; Mayer, J. M. Activationless Multiple-Site Concerted Proton–Electron Tunneling. J. Am. Chem. Soc. 2018, 140, 7449–7452.
(44) Sension, R. J.; Repinec, S. T.; Hochstrasser, R. M. Femtosecond laser study of energy disposal in the solution phase isomerization of stilbene. J. Chem. Phys. 1990, 93, 9185–9188.
(45) Marcus, R. A. On the Theory of Oxidation-Reduction Reactions Involving Electron Transfer. I. J. Chem. Phys. 1956, 24, 966–978.
(46) Weller, A. Photoinduced Electron Transfer in Solution: Exciplex and Radical Ion Pair Formation Free Enthalpies and their Solvent Dependence. Z. Phys. Chem. 1982, 133, 93–98.
(47) Feskov, S. V.; Rogozina, M. V.; Ivanov, A. I.; Astor, A.; Koch, M.; Vauthey, E. Magnetic field effect on ion pair dynamics upon bimolecular photoinduced electron transfer in solution. J. Chem. Phys. 2019, 150, No. 024501.
(48) Hasharoni, K.; Levanon, H.; Greenfield, S. R.; Gosztola, D. J.; Svec, W. A.; Wasielewski, M. R. Mimicry of the Radical Pair and Triplet States in Photosynthetic Reaction Centers with a Synthetic Model. J. Am. Chem. Soc. 1995, 117, 8055–8056.
(49) Yee, E. L.; Cave, R. J.; Guyer, K. L.; Tyma, P. D.; Weaver, M. J. A survey of ligand effects upon the reaction entropies of some transition metal redox couples. J. Am. Chem. Soc. 1979, 101, 1131–1137.
(50) Hupp, J. T.; Weaver, M. J. Solvent, ligand, and ionic charge effects on reaction entropies for simple transition-metal redox couples. Inorg. Chem. 1984, 23, 3639–3644.
(51) Hupp, J. T.; Weaver, M. J. Influence of specific reactant-solvent interactions on intrinsic activation energies for outer-sphere electron-transfer reactions. J. Phys. Chem. 1984, 88, 1860–1864.
(52) Closs, G. L.; Johnson, M. D.; Miller, J. R.; Pietrowski, P. A connection between intramolecular long-range electron, hole, and triplet energy transfers. J. Am. Chem. Soc. 1989, 111, 3751–3753.
(53) Finckh, P.; Heitze, H.; Volk, M.; Michel-Beyerle, M. E. Electron donor/acceptor interaction and reorganization parameters from temperature-dependent intramolecular electron-transfer rates. J. Phys. Chem. 1988, 92, 6584–6590.
(54) Liu, J. Y.; Bolton, J. R. Intramolecular photochemical electron transfer. 7. Temperature dependence of electron-transfer rates in...
covalently linked porphyrin-amide-quinone molecules. J. Phys. Chem. 1992, 96, 1718−1725.

(55) Serpa, C.; Gomes, P. J. S.; Arnaut, L. G.; de Melo, J. S.; Formosinho, S. J. Temperature Dependence of Ultra-Exothermic Charge Recombinations. ChemPhysChem 2006, 7, 2533−2539.

(56) Vath, P.; Zimmt, M. B.; Matyushov, D. V.; Voth, G. A. A Failure of Continuum Theory: Temperature Dependence of the Solvent Reorganization Energy of Electron Transfer in Highly Polar Solvents. J. Phys. Chem. B 1999, 103, 9130−9140.

(57) Parson, W. W. Temperature Dependence of the Rate of Intramolecular Electron Transfer. J. Phys. Chem. B 2018, 122, 8824−8833.

(58) Mentel, K. K.; Serra, A.; Abreu, P. E.; Arnaut, L. G. Higher activation barriers can lift exothermic rate restrictions in electron transfer and enable faster reactions. Nat. Commun. 2018, 9, 2903.

(59) Li, X.-Y. An overview of continuum models for nonequilibrium solvation: Popular theories and new challenge. Int. J. Quantum Chem. 2015, 115, 700−721.

(60) Newton, M. D.; Basilevsky, M. V.; Rostov, I. V. A frequency-resolved cavity model (FRCM) for treating equilibrium and nonequilibrium solvation energies: 2. Evaluation of solvent reorganization energies. Chem. Phys. 1998, 232, 201−210.

(61) Leontyev, I. V.; Vener, M. V.; Rostov, I. V.; Basilevsky, M. V.; Newton, M. D. Continuum level treatment of electronic polarization in the framework of molecular simulations of solvation effects. J. Chem. Phys. 2003, 119, 8024−8037.

(62) Small, D. W.; Matyushov, D. V.; Voth, G. A. The Theory of Electron Transfer Reactions: What May Be Missing? J. Am. Chem. Soc. 2003, 125, 7470−7478.

(63) Leontyev, I. V.; Basilevsky, M. V.; Newton, M. D. Theory and computation of electron transfer reorganization energies with continuum and molecular solvent models. Theor. Chem. Acc. 2004, 111, 110−121.

(64) Barzykin, A. V.; Frantsuzov, P. A.; Seki, K.; Tachiya, M. Solvent Effects in Nonadiabatic Electron-Transfer Reactions: Theoretical Aspects. Adv. Chem. Phys. 2002, 111, 611−616.

(65) Vath, P.; Zimmt, M. B. A Spectroscopic Study of Solvent Reorganization Energy: Dependence on Temperature, Charge Transfer Distance, and the Type of Solute−Solvent Interactions. J. Phys. Chem. A 2000, 104, 2626−2633.

(66) Schmidt, J. A.; Liu, J.-Y.; Bolton, J. R.; Archer, M. D.; Gadžekepo, V. P. Y. Intramolecular photochemical electron transfer. Part 5—Solvent dependence of electron transfer in a porphyrin−amide−quinone molecule. J. Chem. Soc., Faraday Trans. 1 1989, 85, 1027−1041.

(67) Mataga, N.; Chosrowjan, H.; Taniguchi, S. Ultrafast charge transfer in excited electronic states and investigations into fundamental problems of exciplex chemistry: Our early studies and recent developments. J. Photochem. Photobiol., C 2005, 6, 37−79.

(68) Mataga, N.; Taniguchi, S.; Chosrowjan, H.; Osuka, A.; Yoshida, N. Ultrafast charge transfer and radiationless relaxations from higher excited state (S2) of directly linked Zn-porphyrin (ZP)-acceptor dyads: investigations into fundamental problems of exciplex chemistry. Chem. Phys. 2003, 295, 215−228.

(69) Brunschwig, B. S.; Ehrenson, S.; Sutin, N. Solvent reorganization in optical and thermal electron-transfer processes. J. Phys. Chem. 1986, 90, 3657−3668.

(70) Wallin, S.; Monneron, C.; Blart, E.; Gankou, J.-R.; Odobel, F.; Hammarström, L. State-Selective Electron Transfer in an Unsymmetric Acceptor−Zn[II]porphyrin−Acceptor Triad: Toward a Controlled Directionality of Electron Transfer from the Porphyrin S2 and S1 States as a Basis for a Molecular Switch. J. Phys. Chem. A 2010, 114, 1709−1721.

(71) Matyushov, D. V. Electron transfer in nonpolar media. Phys. Chem. Chem. Phys. 2020, 22, 10653−10665.

(72) Wilkinson, F.; McGarvey, D. J.; Olea, A. F. Factors governing the efficiency of singlet oxygen production during oxygen quenching of singlet and triplet states of anthracene derivatives in cyclohexane solution. J. Am. Chem. Soc. 1993, 115, 12144−12151.