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Charge Transfer Dynamics at the α/β Subunit Interface of a Photochemical Ribonucleotide Reductase

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Abstract

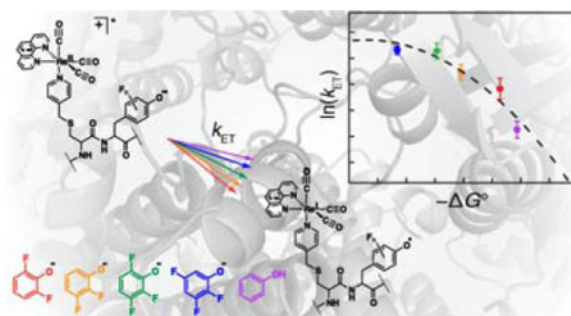
Ribonucleotide reductase (RNR) catalyzes the conversion of ribonucleotides to deoxyribonucleotides to provide the monomeric building blocks for DNA replication and repair. Nucleotide reduction occurs by way of multi-step proton-coupled electron transfer (PCET) over a pathway of redox active amino acids spanning ~ 35 Å and two subunits (α_2 and β_2). Despite the fact that PCET in RNR is rapid, slow conformational changes mask kinetic examination of these steps. As such, we have pioneered methodology in which site-specific incorporation of a [Re^I] photooxidant on the surface of the β_2 subunit (photo β_2) allows photochemical oxidation of the adjacent PCET pathway residue β -Y₃₅₆ and time-resolved spectroscopic observation of the ensuing reactivity. A series of photo β_2 s capable of performing photoinitiated substrate turnover have been prepared in which four different fluorotyrosines (F_nY_s) are incorporated in place of β -Y₃₅₆. The F_nY_s are deprotonated under biological conditions, undergo oxidation by electron transfer (ET) and provide a means by which to vary the ET driving force (G°) with minimal additional perturbations across the series. We have used these features to map the correlation between G° and k_{ET} both with and without the fully assembled photoRNR complex. The photooxidation of F_nY₃₅₆ within the α/β subunit interface occurs within the Marcus inverted region with a reorganization energy of $\lambda \approx 1$ eV. We also observe enhanced electronic coupling between donor and acceptor (H_{DA}) in the presence of an intact PCET pathway. Additionally, we have investigated the dynamics of proton transfer (PT) by a variety of methods including dependencies on solvent isotopic composition, buffer concentration, and pH. We present evidence for the role of α_2 in facilitating PT during β -Y₃₅₆ photooxidation; PT occurs by way of readily exchangeable positions and within a relatively “tight” subunit interface. These findings show that RNR controls ET by lowering λ , raising H_{DA} , and directing PT both within and between individual polypeptide subunits.

Graphical abstract

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ASSOCIATED CONTENT

Experimental methods and instrumentation, expression, purification, radical yields and specific activities of photo β_2 s, table of simulated Marcus parameters as a function of distance, SDS-PAGE purity gels for photo β_2 s, additional controls for photochemical turnover experiments, exemplary emission decay traces, low temperature emission spectrum of Y₃₅₆F-photo β_2 , and differential pulse voltammetry data for *N*-acetyl *C*-amide protected F_nYs. This material is available free of charge via the Internet at <http://pubs.acs.org>.



INTRODUCTION

Enzymatic electron transfer (ET) plays a central role in biological energy transduction. Catalytic cofactors face the formidable challenge of maintaining control over highly reactive species within the milieu of the protein scaffold. This feat is particularly remarkable when ET processes involve the formation of amino acid radical intermediates, the reversibility of which is dictated by the surrounding environment. Such events are usually coupled to the transfer of a proton via proton-coupled ET (PCET).⁻ The coordination of proton and electron is critical to the function of ribonucleotide reductase (RNR), whose catalytic ability to convert ribonucleotides to deoxyribonucleotides relies on reversible long-range PCET spanning two subunits and ~35 Å (Figure 1). The active form of the class Ia RNR from *E. coli* is composed of two obligate homodimers, α_2 and β_2 (Figure 1).⁻ The α_2 subunit contains the active site, as well as additional binding sites for allosteric effectors that control both overall activity, and substrate specificity. The β_2 subunit contains the diferric-tyrosyl radical cofactor, $\text{Fe}^{\text{III}}_2(\mu\text{-O})/\text{Y}^\bullet$, responsible for initiating active site chemistry. Translocation of this stable radical in the β_2 subunit, to the active site in the α_2 subunit occurs by way of multi-site PCET hopping over a pathway of redox active amino acids.[·] The RNR mechanism begins with substrate binding in α_2 ,[·] which triggers a conformational change resulting in proton transfer from a water molecule ligated to the diferric co-factor to the stable $\beta\text{-Y}_{122}^\bullet$. Simultaneous ET results in oxidation of $\beta\text{-Y}_{356}$,[·] exemplifying orthogonal, or bidirectional PCET. From here, PCET across the subunit interface sequentially oxidizes $\alpha\text{-Y}_{731}$, $\alpha\text{-Y}_{730}$,[·] and then $\alpha\text{-C}_{439}$ (Figure 1 inset), via collinear PCET (in which protons and electrons are mutually exchanged between the same donor and acceptor partners). Upon oxidation, $\alpha\text{-C}_{439}^\bullet$ initiates active site chemistry by hydrogen atom abstraction from substrate.⁻ Multi-step substrate-based radical chemistry follows,[·] after which reverse PCET carries the radical “hole” back to its stable resting state at Y_{122} in β via the same PCET pathway.

To measure the kinetics of PCET events in RNR, we have developed phototriggering methods to bypass rate-determining conformational changes. This methodology has enabled detailed studies of photoinitiated substrate turnover, spectroscopic observation of photogenerated radicals, and measurement of both radical injection rates into α_2 , and radical propagation rates through α_2 to the active site. A photochemically competent β_2 subunit (photo β_2) is prepared by installing three mutations (C_{268}S , C_{305}S , and S_{355}C) to render a single cysteine residue surface-exposed, and facilitate site-specific conjugation of a

bromomethylpyridyl rhenium(I) tricarbonyl phenanthroline complex ($[Re^I]$) to position β_{355} via an S_N2 reaction. Recently, we extended this methodology by incorporating 2,3,5- F_3Y at position β_{356} via nonsense codon suppression methodology (Figure 1, inset). This allowed direct observation of photogenerated radicals by transient absorption (TA) spectroscopy and elucidation of the kinetics for radical propagation steps through α .

Little is known about the structure and dynamics occurring at the RNR subunit interface. Not only is a complete crystal structure of the $\alpha_2\beta_2$ complex absent, but structures of the β_2 and α_2 subunits alone all lack structural information regarding the key redox active residue $\beta\text{-}Y_{356}$. Herein, we unveil new aspects of the molecular and intermolecular interactions governing charge transfer within the RNR subunit interface by modulating the driving force through incorporation of unnatural amino acid fluorotyrosines (F_nY_s), whose unique electrochemical and acid/base properties allow systematic correlation of k_{ET} and G° , as well as dependencies on pH, buffer concentration, and solvent isotopic composition. A series of photo β_2 s have been prepared in which four different F_nY_s ($n = 2\text{--}3$) are incorporated at position β_{356} . Each of these photo β_2 s is capable of photochemical substrate turnover, and they demonstrate reactivity that depends on the oligomeric state (dictated by allosteric effectors), highlighting an allegiance to wt-RNR chemistry. The F_nY_s used here display a range of reduction potentials (Figure 2), yet incur only small perturbations relative to each other across the series. These features, coupled with the fact that F_nY_s can exist in their deprotonated forms under biologically compatible conditions, render the series of photo β_2 s ideally suited for examination of the relationship between k_{ET} and G° within the unique dielectric environment of a protein/protein interface.

By examining the relationship between k_{ET} and G° , we have found that the photooxidation of F_nY_{356} occurs within the Marcus inverted region. From the Marcus curve, we estimate that $\lambda \approx 1$ eV within the α/β subunit interface, and that the presence of an intact PCET pathway increases the electronic coupling between donor and acceptor. Through this work we have uncovered a distinct role of the α_2 subunit in facilitating PT during $\beta\text{-}Y_{356}$ oxidation. We have further elaborated our studies by examining the PCET kinetics as a function of pH, solvent isotopic constitution, and buffer concentration all as a function of oligomeric state and in the presence and absence of the next PCET pathway residue, $\alpha\text{-}Y_{731}$. Our data support a model for the PT pathway that exhibits tightly bound, yet solvent exchangeable protons that assist the intersubunit ET.

EXPERIMENTAL SECTION

Materials

Wt- α_2 (2,000 nmol/mg/min) was expressed from pET28a-*nrdA* and purified as previously described. A glycerol stock of $Y_{731}F\text{-}\alpha_2$ was available from a previous study, and was expressed and purified as wt- α_2 . All α_2 proteins were pre-reduced and treated with hydroxyurea (HU, Sigma Aldrich) by incubation with 30 mM dithiothreitol (DTT, Promega) for 30 min at room temperature, then addition of 15 mM more DTT and 15 mM HU, followed by buffer exchange on a Sephadex G-25 or G-50 column. [5- 3H]-cytidine 5'-diphosphate sodium salt hydrate ([5- 3H]-CDP) was purchased from ViTrax (Placentia, CA). 2,3,5-Trifluoroboronic acid, 2,3,6-trifluorophenol, 2,3-difluorophenol, and 3,5-

performed in triplicate on independently prepared samples, with the exception of pH dependence data, in which measurements were performed only in duplicate.

Analysis of kinetics data was performed by fitting each emission decay trace to Eq. 1 over the span of 0.1–4.5 μ s using OriginPro 8.0 software (OriginLab). Acceptability of fitting was determined on the basis of qualitative symmetry of residuals about zero amplitude and the R^2 factor. Exemplary traces and residuals are available in Figure S3.

$$y=y_0+Ae^{-x/t} \quad (1)$$

Under each experimental condition two sets of lifetime (τ) data were obtained in order to calculate k_{ET} or k_{PCET} . These two τ values correspond to the $[Re^I]^*$ lifetime measured (under each experimental condition) with the photo β_2 containing F at position β_{356} (τ_o , green in Scheme 1) and with the photo β_2 containing Y or F_nY at position β_{356} (τ_Y , purple in Scheme 1). Plugging these numbers into Eq. 2 then yields the rate constant for the relevant pathway of $[Re^I]^*$ quenching, namely Y or F_nY oxidation.

$$k_{(PC)ET} = \frac{1}{\tau_Y} - \frac{1}{\tau_o} = k_Y - k_o \quad (2)$$

The absolute uncertainty of each $k_{(PC)ET}$ (δ) was calculated according to Eq. 3 below, where σ represents one standard deviation among each set of lifetime data (τ). As a result of high S/N in these experiments, the error associated with the fitting process was consistently close to three orders of magnitude smaller than the lifetime value itself. As such, this error was insignificant relative to the standard deviation between replicates.

$$\delta = \sqrt{\left(\frac{\sigma_Y}{\tau_Y^2}\right)^2 - \left(\frac{\sigma_o}{\tau_o^2}\right)^2} \quad (3)$$

Dependence of k_{PCET} on oligomeric state was ascertained by conducting emission quenching experiments as outlined above in assay buffer (15 mM $MgSO_4$, 1 mM EDTA, 50 mM HEPES adjusted to pH 7.6) containing either 1 mM CDP and 3 mM ATP to promote the active $\alpha_2\beta_2$ oligomer, or 200 μ M dATP to promote the inactive $\alpha_4\beta_4$ oligomeric state.

Solvent isotope effect samples were prepared by lyophilizing 4 \times concentrations of assay buffer followed by rehydration in D2O. The buffer pL was adjusted to 7.6 for all samples by addition of NaOD or NaOH according to Eq. 4.

$$pH=0.929pD+0.41 \quad (4)$$

All protein samples used were exchanged into $[^2H]$ -assay buffer by 5 cycles of 5-fold concentration followed by resuspension using pre-soaked 30 kDa MWCO centrifugal filters

(Millipore) to ensure that <3% H₂O remained in D₂O samples. Samples prepared in [¹H]-assay buffer were treated identically to ensure that any effect of this treatment was identical for all samples under study.

pH dependence of $k_{(PC)ET}$ was performed by measuring emission lifetimes as outlined above with samples containing 1 mM CDP, 3 mM ATP in assay buffer containing 50 mM of one of the following: MES: pH 5.4–6.8; HEPES: pH 6.8–8.0; TAPS: pH 8.2–9.0, adjusted to the appropriate pH. In these experiments the emission is quenched much more rapidly (where the forward and reverse rates occur simultaneously to set up an equilibrium) when F_nY is deprotonated, thus revealing the pK_a by a sharp reduction in the total emission intensity.

RESULTS

In order to assess the influence of the protein environment on ET kinetics, a series of photoβ₂s were prepared in which four different F_nYs (Figure 2) are incorporated at position β₃₅₆. In this way, the driving force for electron transfer from F_nY⁻ to [Re^I]* was varied over ~150 mV while incurring minimal additional perturbations across the series. The dissociation constant for photoβ₂ binding to α₂ is 0.7 ± 0.1 μM. This value reveals that incorporation of the [Re^I] complex incurs only a small disruption to the subunit interaction, as the K_d for wt-RNR is 0.2 μM. Of the six photoβ₂s under study, only that with 3,5-F₂Y displayed native enzymatic activity in its holo-state (still containing the Y₁₂₂• cofactor, prior to reduction with HU). The specific activities measured before and after labeling with [Re^I] were 1000 and 300 nmol(min-mg)⁻¹, respectively for this photoβ₂ (Table S1). For reference wt-β₂ has specific activity of 6000 nmol(min-mg)⁻¹. The photoβ₂ with Y at position 356 exhibits specific activities of 2300 and 200 nmol(min-mg)⁻¹, before and after labeling with [Re^I], respectively. In both of these cases (3,5-F₂Y and Y), the activity prior to labeling exhibits nonlinear behavior due to the oxidation of S₃₅₅C. The activity is linear after labeling with [Re^I] because the thiol is converted to a thioether, which is more difficult to oxidize. The fact that Y and 3,5-F₂Y are the easiest Y-derivatives in the series to oxidize coincides with the fact that these are the only two photoβ₂s exhibiting non-photochemical activity.

Photochemical turnover

We sought to confirm that these photoβ₂s are photochemically competent for turnover. To this end, we performed single turnover experiments under white light illumination in the presence of radiolabelled substrate (Figure 3). We note that in all cases the native Y₁₂₂• cofactor within the photoβ₂s has been quantitatively reduced by treatment with HU, rendering them incapable of undergoing turnover via the non-photochemical mechanism. Residual reactivity can be seen in the amount of turnover observed in the dark (green bars, Figure 3). Similarly, control experiments with variants containing redox-inactive pathway substitutions exhibit very low background levels of product formation under illumination (Figure S2).

Two methods for the photochemical generation of F_nY• have been implemented in this work. In the direct method, (Scheme 1, purple) the excited state ³[Re^I]* complex oxidizes the adjacent Y-derivative directly. A second method, utilizes flash-quench methodology where, in the presence of excess Ru(NH₃)₆Cl₃, the excited state ³[Re^I]* is oxidized via

bimolecular reaction with $\text{Ru}(\text{NH}_3)_6^{3+}$, yielding $\text{Ru}(\text{NH}_3)_6^{2+}$ and $[\text{Re}^{\text{II}}]$. This $[\text{Re}^{\text{II}}]$ species then oxidizes F_nY (Scheme 1, blue). These two photochemical generation methods were implemented under single turnover conditions for all photo β_2 s. Wt-RNR can generate a theoretical maximum of 4 dCDP products per α_2 subunit, one for each α monomer, and a second per monomer arising from the re-reduction of active site cysteines (α -C₂₂₅ and C₄₆₂) by disulfide exchange with two C-terminal cysteines (α -C₇₅₄ and C₇₅₉); this maximum is diminished for wt- β_2 under flash-quench conditions (Figure 3, right). Notwithstanding, all photo β_2 s exhibited the ability to produce products via photochemical activation (Figure 3), lending confidence that our method retains authenticity to RNR.

Oligomeric state

RNR exists naturally as a dimer, but can enter a complex equilibria involving oligomeric forms (α_2 , β_2 , $\alpha_2\beta_2$, and $\alpha_4\beta_4$) as dictated in vivo by the presence of allosteric effectors and substrates. The formation of an inhibited complex, the $\alpha_4\beta_4$ oligomeric state, occurs at the high protein concentrations often required for in vitro biophysical measurements. Because the total protein concentration (35 μM) of our spectroscopic experiments may engender the inactive $\alpha_4\beta_4$ state, we needed to establish the presence of $\alpha_2\beta_2$ dimer in our experiments. The inhibited $\alpha_4\beta_4$ state is favored for dATP concentrations $>100 \mu\text{M}$, whereas the active form is known to exist (at least transiently) in the case where ATP and CDP are present. We used these facts to investigate which oligomeric state(s) predominate under our experimental conditions. Under the conditions of ATP and CDP, we observe a 34% enhancement in k_{PCET} (with the photo β_2 containing Y at position 356) in the presence of wt- α_2 relative to Y₇₃₁F- α_2 (Table 1). We have previously ascribed this enhancement to a dependence on the presence of an intact PCET pathway. On the contrary, in the presence of 200 μM dATP (promoting the $\alpha_4\beta_4$ state), identical k_{PCET} values were obtained in the presence of wt- α_2 and Y₇₃₁F- α_2 (Table 1). These results are consistent with the structure of the $\alpha_4\beta_4$ oligomer, which unlike the globular form of the active $\alpha_2\beta_2$ complex, forms a donut-shaped oligomer. The C-termini of β_2 remain bound to α_2 in this state, giving rise to exposure of Y₃₅₆ to bulk solution. Thus Y₇₃₁ of the α_2 subunit is not adjacent to Y₃₅₆ in the $\alpha_4\beta_4$ oligomer and the PCET pathway from β_2 to the neighboring α_2 subunit is disrupted. Hence the photooxidation of Y₃₅₆ within the inhibited oligomeric form should be unaffected by the presence or absence of α -Y₇₃₁; the data in Table 1 show this to be the case. The results of Table 1 therefore establish that the photo $\beta_2\alpha_2$ state exists under normal experimental conditions in which CDP and ATP are present and that we are examining photooxidation events within the subunit interface of the active $\alpha_2\beta_2$ complex.

Energetics

The overall free energy change accompanying the generation of the photo β_2 charge-separated state is schematically represented in Figure 2 and given by,

$$-\Delta G^\circ = E^\circ([\text{Re}^{\text{I}*}] / [\text{Re}^{\text{0}}]) - E^\circ(\text{F}_n\text{Y}^\bullet / \text{F}_n\text{Y}^-) \quad (5)$$

$[\text{Re}^{\text{I}}]$ and F_nY^- initially have equal and opposite charges that are neutralized upon ET; the additional contribution to G° from an electrostatic work term is negligible. The excited

state emission energy was determined from the low temperature (77 K) emission spectrum of [Re]-Y₃₅₆F-β₂ as previously described to yield G_{MLCT} of 2.73 eV (Figure S4). A shift in this value is not observed when the measurement is performed in the presence of the α₂ subunit. The ground state reduction potential of [Re^I(phen)(CO)₃EtPy]-PF₆ has previously been reported to be -0.79 V vs NHE. As summarized on the Latimer diagram of Figure 4, from these values, the excited state reduction potential is calculated to be $E^{\circ}([Re^I]^*/[Re^0]) = 1.94$ V vs NHE.

The F_nY reduction potentials were measured by differential pulse voltammetry (DPV) with the *N*-acetyl *C*-amide protected amino acid derivatives (Figure S5). The propensity of tyrosine to undergo fast bimolecular chemical reactions following its one-electron oxidation has precluded accurate determination of the thermodynamic (reversible) reduction potentials. However, evidence from de novo protein model systems in which redox active amino acids are sequestered from solution, the reversible reduction potentials for both Y and 3,5-F₂Y have been determined. In each case, the absolute values of E° differ by ~150 mV from those determined by DPV. However, the *relative* potentials of Y and 3,5-F₂Y (E° = 30 mV at pH 5.7), remain very close to the relative potentials measured by DPV (E_p = 50 mV, pH 5.7). Thus, in the current work we apply the assumption that all F_nYs within the subunit interface will be subject to similar perturbations, prompting the caveat that all G° s may be shifted and that values are to be interpreted as relative rather than absolute. We note that rapid freeze quench electron paramagnetic resonance (RFQ-EPR) spectroscopy of radical equilibration along the PCET pathway reveals that β-Y₃₅₆ is about 100 mV easier to oxidize than the adjacent α-Y₇₃₁ residue but we have not included any correction factors in our calculations of G° herein and simply reiterate the aforementioned caveat.

The p*K*_as of the F_nYs within the photoβ₂α₂ complex (Table 2) are obtained by measuring the intensity of steady-state [Re^I]^{*} emission as a function of pH. Figure 5 shows the normalized integrated emission intensity (*I*) as a function of pH. The p*K*_a is afforded from fitting these data to the following,

$$10^{(pH-pK_a)} = (I - I_{min}) / (I_{max} - I_{min}) \quad (6)$$

Only slight differences are found between the p*K*_as for F_nYs in the protein and the corresponding *N*-acetyl *C*-amide protected F_nYs. As previously observed for RNR, there is little perturbation to the phenolic p*K*_as at position β₃₅₆ within the complex.

Based on these data, experiments to examine the kinetics of ET for the photoβ₂s were conducted at pH 8.2 such that the F_nYs are nearly fully deprotonated. Accordingly, the reduction potentials to be used in Eq. 5 were taken from the pH-independent regions of DPV Pourbaix diagrams (Figure S5). For the cases of Y and 2,3-F₂Y (which is only ~80% deprotonated at pH 8.2), the Nernst equation (Eq. 7) was applied to calculate the reduction potentials under the experimental conditions.

$$E^{\circ} = E_p(F_2Y^{\bullet}/F_2Y^{-}) + \frac{2.3RT}{nF} \log \left\{ 1 + \frac{10^{-pH}}{10^{-pK_a}} \right\} \quad (7)$$

With the excited state reduction potential of $^3[Re^I]^*$ and each F_nY^{-} in hand, the driving force for photogeneration of each charge-separated state may be determined from Eq. 5.

Electron Transfer Kinetics

The excited state lifetime of $^3[Re^I]^*$ (τ) for each photo β_2 was measured by nanosecond laser flash photolysis (in the absence of flash quencher). The rate constant for the formation of the charge-separated state resulting from $^3[Re^I]^*$ reduction via F_nY oxidation may be determined from Eq. 2, where τ for photo β_2 s containing Y or F_nY at position β_{356} is referenced to τ_0 for $Y_{356}F$ -photo β_2 (purple versus green in Scheme 1). In this way, all pathways for $^3[Re^I]^*$ decay besides that proceeding via F_nY oxidation are accounted for and k_{ET} represents the rate constant of only the relevant reaction. The rate constants summarized in Figure 6 were determined from measurements in triplicate for each photo β_2 alone, in the presence of wt- α_2 , and in the presence of $Y_{731}F$ - α_2 . The latter experiment was performed as a control wherein the $\alpha_2\beta_2$ complex is intact but charge injection into α_2 is prevented; thus it is a measure of the decay of the photogenerated radical within the β_2 subunit of a fully assembled system. Experiments with α_2 variants were conducted at protein concentrations such that >95% of the photo β_2 moiety was in complex with α_2 , based on the K_d measured for photo $\beta_2\alpha_2$ dissociation of $0.7 \pm 0.1 \mu M$. All samples contained saturating concentrations of substrate (CDP) and effector (ATP) in order to ensure binding in an active conformer (Table 1). A detailed account of our interpretation and analysis of these data is included in the Discussion section, wherein the data are replotted and overlaid with simulations of the semiclassical Marcus relation (Eq. 9).

The effects of solvent isotopic composition, buffer concentration, and pH, on the kinetics of charge transfer at the interface were examined, both in the presence and absence of α_2 and the $Y_{731}F$ - α_2 variant. We note that where Y, or a protonated F_nY is positioned at β_{356} we have labeled the corresponding rate constants with k_{PCET} rather than k_{ET} . Solvent kinetic isotope measurements (SKIE) of the photooxidation of Y_{356} occurs with SKIE ~ 1.5 under all three conditions listed in Table 3. Such isotope effects are observed⁻ and theoretically predicted when the PCET involves electron transfer through a hydrogen bond. This observation of a small isotope effect suggests that proton acceptor(s) reside in positions that are hydrogen bonded and are readily exchangeable with solvent. This holds both in the presence and absence of α_2 , and regardless of whether or not the PCET pathway is intact. Alternatively, a modest to negligible value may be the result of viscosity and/or pK_a differences resulting from the different properties of the two buffers.

Figure 7 displays the results of the dependence of k_{PCET} on buffer concentration. The effects are small when compared to similar studies performed with small molecule model systems in solution, in which k_{PCET} may change by an order of magnitude over similar ranges of buffer concentration. Unlike studies with small molecules, we also find that a dependence

on buffer concentration is manifest only at low concentrations. These data suggest that the subunit interface is relatively protected from solvent species. The dependence that we do observe is slightly larger in the presence of wt- α_2 relative to Y₇₃₁F- α_2 . This may implicate a role for α -Y₇₃₁ in facilitating PT from β -Y₃₅₆, although it has been shown that this residue does not interact directly with β -Y₃₅₆.

To further probe the extent that the α_2 subunit facilitates PT during photooxidation of β -Y₃₅₆ (either by specific residues or by dictating conformation), the pH dependence of k_{PCET} and k_{ET} below and above the phenolic pK_a of 2,3-F₂Y-photo β_2 was examined. This F_nY was selected because it has a pK_a of 7.8 (Figure 5, Table 2), in the middle of the pH range that is accessible for the class Ia RNR. Figure 8 summarizes the pH dependence of photooxidation of this residue with the photo β_2 alone, or in the presence of wt- or Y₇₃₁F- α_2 . In the absence of α_2 , k_{PCET} increases linearly with pH up to the phenolic pK_a , after which a sharp break occurs. Conversely, in the presence of α_2 continuous behavior is observed over the entire pH range examined. This result supports the contention that α_2 facilitates PT during β -Y₃₅₆ photooxidation.

DISCUSSION

RNR maintains control over long-range PCET events by way of well-choreographed conformational dynamics.³³ Kinetics measurements suggest that the PCET pathway is finely tuned and that it is the target of the conformational gating.³⁴ Crystallographic elucidation of the active $\alpha_2\beta_2$ structure, however, has not been achieved so the structural details of conformational gating and its relation to radical transport via PCET remain to be elucidated. Some structural insight is provided from shape and charge complementarity studies of the separately crystallized subunits, furnishing the so-called $\alpha_2\beta_2$ docking model (Figure 1), as supported by cryogenic electron microscopy (cryoEM), small angle x-ray scattering (SAXS), and pulsed electron-electron double resonance (PELDOR) spectroscopy with metastable trapped states of the $\alpha_2\beta_2$ complex.³⁵ Nonetheless, the last 25–30 amino acids remain disordered in the available structures of the β_2 subunit. It is this C-terminal tail of β , specifically the last 15 residues, that is largely responsible for binding interactions with the α_2 subunit and importantly the structurally disordered portion includes the key redox-active Y₃₅₆, which mediates charge transfer at the interface of the α_2 and β_2 subunits.

Owing to this disorder, the specific molecular interactions within the $\alpha_2\beta_2$ subunit interface remain largely unknown. More generally, insights have emerged from high-field EPR and [²H]-electron nuclear double resonance (ENDOR) spectroscopies in which radicals are trapped en route to the active site by a genetically encoded NH₂Y residue to furnish a spectroscopic handle for examining hydrogen bonding (H-bonding) interactions along the PCET pathway. This work has revealed strong collinear H-bonding between α -Y₇₃₁ and α -Y₇₃₀, and that no such strong and/or static H-bonding interactions exist at position Y₃₅₆. However, large shifts in g_x values for β -Y₃₅₆ do persist, suggesting that the dielectric medium of the interface is strongly perturbed within the active complex despite the absence of distinct H-bonds.

To directly investigate charge (electron and proton) dynamics at the interface, we have examined radical formation at the β -Y₃₅₆ position within the $\alpha_2\beta_2$ subunit interface by using fluorinated tyrosines. These unnatural amino acids allow charge transfer to be examined over a sizable driving force range according to the Marcus relation,

$$k_{ET} = \frac{2\pi H_{AD}^2}{\hbar \sqrt{4\pi k_B T \lambda}} \exp\left[-\frac{(\Delta G^\circ + \lambda)^2}{4\lambda k_B T}\right] \quad (8)$$

where the electronic coupling H_{AD} exhibits an exponential dependence on the distance between donor and acceptor (r) relative to the van der Waals contact distance (r_0),

$$H_{AD}(r) = H_{AD}(r_0) \exp\left[-\frac{1}{2}\beta(r - r_0)\right] \quad (9)$$

k_{ET} is related to G° via reorganization energy λ and electronic coupling matrix element H_{AD} . For the homologous series of F_nYs, λ and H_{AD} may be assumed to be similar across the series and hence the correlation between $\ln(k_{ET})$ and G° is straightforward. In this case, the parabolic relationship between $\ln(k_{ET})$ and G° predicts that an increase in G° results in faster rates when $\lambda > |G^\circ|$ and slower rates when $\lambda < |G^\circ|$, known as the Marcus inverted region.

For the study reported herein, over the pH regime examined, the F_nYs are deprotonated and radical generation in β -F_nY₃₅₆ is consequently well described by a pure ET. As Figure 6 shows, k_{ET} decreases with increasing driving force, a clear signature of Marcus inverted behavior. Entry into the Marcus inverted regime is largely a result of the high excited state reduction potential of ³[Re^I]* (1.94 V vs NHE).

Figure 9 shows a simulation of Eq. 8. The Marcus simulations were performed using a full range of possible distances ($r = 4\text{--}16$ Å, Table S2) accounting for all reasonable conformations of the [Re]–C–F_nY fragment (Figure S6). Regardless of which value of r is selected, the relative ratios of H_{AD} values under the different conditions remain the same (Table S2), facilitating qualitative comparisons to be made. The range of G° spanned by the F_nY-photo β_2 s (~150 mV) did not permit accurate fitting of Eq. 8. As such, the dashed lines of Figures 9 and 10 represent simulations of Eq. 8. The parameters used in these simulations represent the best agreement with the measured values. The data in Figure 9 were simulated with $\lambda = 0.98$ eV, a value that is approximately half that found for model [Re]–F_nY systems ($\lambda = 1.9$ eV). The lower λ for the RNR complex indicates that the subunit interface has evolved to facilitate inter-protein charge transfer, shielding reactive radical intermediates from the bulk solvent by lowering λ , yet all without precluding the required concomitant PT (vide infra). This result is in line with the general observation that λ within the protein environments is depressed as a result of a combination of the constraints imposed on dipole motions due to their confinement within the polypeptide matrix, as well as the low dielectric permittivity within the protein. Conversely, ET across weakly bound protein-

protein complexes, or involving solvent-exposed cofactors incur higher reorganization energy penalties.⁷

We note that the curve in Figure 9 for $\beta\text{-F}_n\text{Y}_{356}$ in the presence of wt- α_2 is higher than that for $\beta\text{-F}_n\text{Y}_{356}$ in the presence of $\text{Y}_{731}\text{F-}\alpha_2$ indicating greater electronic coupling for charge transfer in the former complex. The simulation of data for the $\beta\text{-F}_n\text{Y}_{356}\text{:wt-}\alpha_2$ complex furnished $H_{\text{AD}} = 0.051 \text{ cm}^{-1}$ vs. $H_{\text{AD}} = 0.044 \text{ cm}^{-1}$ for the $\beta\text{-F}_n\text{Y}_{356}\text{:Y}_{731}\text{F-}\alpha_2$ complex. This result is consistent with our previous results that show charge transfer is facilitated by delocalization of the radical over the $\text{Y}_{731}\text{-Y}_{730}$ dyad of α_2 . When the Y-Y dyad is disrupted by a $\text{Y} \rightarrow \text{F}$ substitution in α_2 , radical propagation by charge transfer is attenuated.⁷ Accordingly, we ascribe the higher H_{AD} in the wt complex to delocalization of the hole across the adjacent Y_{731} residue, a process that is precluded when Y_{731}F is present. From the work of Beratan et al., the amplitude of H_{AD} determined from our data ($4 \times 10^{-11} \text{ eV}$) falls within a regime that implicates an ordered network of water molecules and H-bonding interactions within the subunit interface. This result is in line with the small SKIE (Table 3) that we observe, further supporting the presence of water molecules within the α/β interface that facilitate rapid proton exchange at $\beta\text{-Y}_{356}$. The presence of a network of water molecules is also consistent with the lack of observed signals for strong and static H-bonds at position Y_{356} by ENDOR spectroscopy.

In the case of Y, tyrosine largely resides in its protonated state at pH 8.2. However, within the $\alpha_2\beta_2$ complex, the protonated $\beta\text{-Y}_{356}$ follows the trend of the deprotonated $\beta\text{-F}_n\text{Y}_{356}$ s. The rate constant for the former is smallest, as the system exhibits the largest driving force and hence falls deepest into the inverted region. In contrast, when uncomplexed, the protonated $\beta\text{-Y}_{356}$ exhibits anomalous behavior from the deprotonated F_nY s. Because charge transfer is occurring in the inverted region, the faster rate for the photo $\beta\text{-Y}_{356}$ implies a greater activation barrier for charge transfer. Indeed a simulation of the ET reaction of the photo $\beta\text{-F}_n\text{Y}_{356}$ yields $\lambda = 0.74 \text{ eV}$ whereas for photo $\beta\text{-Y}_{356}$ a value of $\lambda = 0.90 \text{ eV}$ is obtained (Figure 10). This simulated curve is presented only to illustrate the graphical repercussions of changing only the value of λ in the simulated curve and does not represent robust support. The difference of 160 mV is entirely consistent with the requirement of PT accompanying ET for the oxidation of tyrosine.

Calculations comparing reorganization energies for concerted PCET versus stepwise ET/PT charge transfer mechanisms in model systems find differences in λ of 134 mV; moreover experimental studies show that the PCET versus ET pathways in model systems incurs higher reorganization energies of 60–500 mV.⁷ The data in Figures 7 and 8 are also consistent with tyrosine oxidation by PCET in photo $\beta\text{-Y}_{356}$. Exposed to solution, the rate of reaction is accelerated by the presence of buffer (Figure 7), which provides a facile acceptor for the proton in the PCET reaction. Moreover, pH dependence of the rate of 2,3- F_3Y oxidation in the 2,3- F_2Y -photo β shows a discontinuity at the $\text{p}K_a$ of the 2,3- F_2Y (Figure 8), also consistent with a PCET process. It is noteworthy that the PCET kinetics are accelerated when the PCET pathway is fully assembled within an $\alpha_2\beta_2$ complex. Whether via organization of the C-terminal tail of β , or through specific interactions with amino acid residues in α , or both, the PCET process for Y oxidation in the presence of α_2 appears to

behave kinetically like an ET process. These data highlight the exquisite control that RNR maintains over reactivity, engendered in part by managing PCET at the protein interface.

CONCLUSION

Radical transport in RNR occurs across two subunits along a PCET pathway that is emerging as the target of conformational gating within the enzyme. A critical step along the PCET pathway occurs at the subunit interface where PCET is proposed to transition from a bidirectional to a collinear PCET pathway. To probe charge transport among the critical amino residues across the interface, we have introduced a series of fluorotyrosines in photo β_2 , thus allowing for the modulation of the free energy driving force and protonation state of residue β_{356} . With this method, few additional perturbations between variants are incurred, allowing for the extraction of the energetics and electronic coupling of charge transport at the RNR interface. We have shown that each photo β_2 is photochemically competent and that rate constants for photooxidation depend on maintaining the active oligomeric state via the presence of allosteric effectors. Analysis of the correlation between k_{ET} and G° reflects the ability of the protein to minimize the reorganization energy and increase electronic coupling at the protein-protein interface in the presence of the intact PCET pathway. Additionally, we present evidence that the α_2 subunit facilitates PT from β -Y $_{356}$, and that this process occurs via solvent exchangeable protons, within a tightly bound subunit interface. Our data add to mounting evidence that PCET through RNR is controlled by way of macromolecular conformational changes targeting the precise alignment of the PCET pathway.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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References

1. Winkler JR, Gray HB. *Annu Rev Biochem.* 1996; 65:537–561. [PubMed: 8811189]
2. Stubbe J, van der Donk WA. *Chem Rev.* 1998; 98:705–762. [PubMed: 11848913]
3. Cukier RI, Nocera DG. *Annu Rev Phys Chem.* 1998; 49:337–369. [PubMed: 9933908]
4. Reece SY, Nocera DG. *Annu Rev Biochem.* 2009; 78:673–699. [PubMed: 19344235]
5. Migliore A, Polizzi NF, Therien MJ, Beratan DN. *Chem Rev.* 2014; 114:5832–3645.
6. Weinberg DR, Gagliardi CJ, Hull JF, Murphy CF, Kent CA, Westlake BC, Paul A, Ess DH, McCafferty DG, Meyer TJ. *Chem Rev.* 2012; 112:4016–4093. [PubMed: 22702235]
7. Hammes-Schiffer S, Stuchebrukhov AA. *Chem Rev.* 2010; 110:6939–6960. [PubMed: 21049940]
8. Stubbe J, Nocera DG, Yee CS, Chang MCY. *Chem Rev.* 2003; 103:2167–2202. [PubMed: 12797828]
9. Brown NC, Reichard P. *J Mol Biol.* 1969:25–38. [PubMed: 4902211]
10. Uhlin U, Eklund H. *Nature.* 1994; 370:533–539. [PubMed: 8052308]
11. Bennati M, Robblee JH, Mugnaini V, Stubbe J, Freed JH, Borbat P. *J Am Chem Soc.* 2005; 127:15014–15015. [PubMed: 16248626]

12. Seyedsayamdost MR, Chan CTY, Mugnaini V, Stubbe J, Bennati M. *J Am Chem Soc.* 2007; 129:15748–15749. [PubMed: 18047343]
13. Minnihan EC, Ando N, Brignole EJ, Olshansky L, Chittuluru J, Asturias FJ, Drennan CL, Nocera DG, Stubbe J. *Proc Natl Acad Sci U S A.* 2013; 110:3835–3840. [PubMed: 23431160]
14. Minnihan EC, Nocera DG, Stubbe J. *Acc Chem Res.* 2013; 46:2524–2535. [PubMed: 23730940]
15. Ge J, Yu G, Ator MA, Stubbe J. *Biochemistry.* 2003; 42:10071–10083. [PubMed: 12939135]
16. Seyedsayamdost MR, Stubbe J. *J Am Chem Soc.* 2006; 128:2522–2523. [PubMed: 16492021]
17. Wörsdörfer B, Conner DA, Yokoyama K, Livada J, Seyed-seyamdost MR, Jiang W, Silakov A, Stubbe J, Bollinger JMJr, Krebs C. *J Am Chem Soc.* 2013; 135:8585–8593. [PubMed: 23676140]
18. Minnihan EC, Seyedsayamdost MR, Uhlin U, Stubbe J. *J Am Chem Soc.* 2011; 133:9430–9440. [PubMed: 21612216]
19. Seyedsayamdost MR, Xie J, Cham CTY, Schultz PG, Stubbe J. *J Am Chem Soc.* 2007; 129:15060–15071. [PubMed: 17990884]
20. Yokoyama K, Smith AA, Corzilius B, Griffin RG, Stubbe J. *J Am Chem Soc.* 2011; 133:18420–18432. [PubMed: 21967342]
21. Nick TU, Lee W, Koßmann S, Neese F, Stubbe J, Bennati M. *J Am Chem Soc.* 2015; 137:289–298. [PubMed: 25516424]
22. Stubbe J, Ackles D. *J Biol Chem.* 1980; 255:8027–8030. [PubMed: 6997288]
23. Stubbe J, Ator M, Krenitsky T. *J Biol Chem.* 1983; 258:1625–1630. [PubMed: 6337142]
24. Olshansky L, Pizano AA, Wei Y, Stubbe J, Nocera DG. *J Am Chem Soc.* 2014; 136:16210–16216. [PubMed: 25353063]
25. Stubbe J, van der Donk WA. *Chem Rev.* 1998; 98:705–762. [PubMed: 11848913]
26. Licht S, Stubbe J. *Compr Nat Prod Chem.* 1999; 5:163–203.
27. Seyedsayamdost MR, Stubbe J. *J Am Chem Soc.* 2007; 129:2226–2227. [PubMed: 17279757]
28. Chang MCY, Yee CS, Stubbe J, Nocera DG. *Proc Natl Acad Sci U S A.* 2004; 101:6882–6887. [PubMed: 15123822]
29. Reece SY, Seyedsayamdost MR, Stubbe J, Nocera DG. *J Am Chem Soc.* 2007; 129:8500–8509. [PubMed: 17567129]
30. Reece SY, Seyedsayamdost MR, Stubbe J, Nocera DG. *J Am Chem Soc.* 2007; 129:13828–13830. [PubMed: 17944464]
31. Holder PG, Pizano AA, Anderson BL, Stubbe J, Nocera DG. *J Am Chem Soc.* 2012; 134:1172–1180. [PubMed: 22121977]
32. Pizano AA, Lutterman DA, Holder PG, Teets TS, Stubbe J, Nocera DG. *Proc Natl Acad Sci U S A.* 2012; 109:39–43. [PubMed: 22171005]
33. Pizano AA, Olshansky L, Holder PG, Stubbe J, Nocera DG. *J Am Chem Soc.* 2013; 135:13250–13253. [PubMed: 23927429]
34. Minnihan EC, Young DD, Schultz PG, Stubbe J. *J Am Chem Soc.* 2011; 133:15942–15945. [PubMed: 21913683]
35. Eriksson M, Uhlin U, Ramaswamy S, Ekberg M, Regnström K, Sjöberg B-M, Eklund H. *Structure.* 1997; 5:1077–1092. [PubMed: 9309223]
36. Högbom M, Galander M, Andersson M, Kolberg M, Hofbauer W, Lassmann G, Nordlund P, Lenzian F. *Proc Natl Acad Sci U S A.* 2003; 100:3209–3214. [PubMed: 12624184]
37. Seyedsayamdost MR, Reece SY, Nocera DG, Stubbe J. *J Am Chem Soc.* 2006; 128:1569–1579. [PubMed: 16448128]
38. Reece SY, Seyedsayamdost MR, Stubbe J, Nocera DG. *J Am Chem Soc.* 2006; 128:13654–13655. [PubMed: 17044670]
39. Pigiet VP, Conley RR. *J Biol Chem.* 1977; 252:6367–6372. [PubMed: 330529]
40. Lunn CA, Kathju S, Wallace BJ, Kushner SR, Pigiet V. *J Biol Chem.* 1984; 259:10469–10474. [PubMed: 6381486]
41. Chen H, Gollnick P, Phillips RS. *Eur J Biochem.* 1995; 229:540–549. [PubMed: 7744078]
42. Simon J, Salzbrunn S, Prakash GKS, Petasis NA, Olah GA. *J Org Chem.* 2001; 66:633–634. [PubMed: 11429844]

43. Bollinger JM Jr, Tong WH, Ravi N, Huynh BH, Edmondson DE, Stubbe J. *Meth Enzymol.* 1995; 258:278–303. [PubMed: 8524156]
44. Kr̄el A, Bal W. *J Inorg Biochem.* 2004; 98:161–166. [PubMed: 14659645]
45. Climent I, Sjöberg BM, Huang CY. *Biochemistry.* 1991; 30:5164–5171. [PubMed: 2036382]
46. Reece SY, Lutterman DA, Seyedsayamdost MR, Stubbe J, Nocera DG. *Biochemistry.* 2009; 48:5832–5838. [PubMed: 19402704]
47. Brown NC, Reichard P. *J Mol Biol.* 1969:39–55. [PubMed: 4902212]
48. Ando N, Brignole EJ, Zimanyi CM, Funk MA, Yokoyama K, Asturias FJ, Stubbe J, Drennan CL. *Proc Natl Acad Sci U S A.* 2011; 108:21046–21051. [PubMed: 22160671]
49. Reece SY, Nocera DG. *J Am Chem Soc.* 2005; 127:9448–9458. [PubMed: 15984872]
50. Kober EM, Caspar JV, Lumpkin RS, Meyer TJ. *J Phys Chem.* 1996; 90:3722–3734.
51. Berry BW, Martínez-Rivera MC, Tommos C. *Proc Natl Acad Sci U S A.* 2012; 109:9739–9743.
52. Ravichandran KR, Liang L, Stubbe J, Tommos C. *Biochemistry.* 2013; 52:8907–8915. [PubMed: 24228716]
53. Yokoyama K, Uhlin U, Stubbe J. *J Am Chem Soc.* 2010; 132:8385–8397. [PubMed: 20518462]
54. Turro C, Chang CK, Leroi GE, Cukier RI, Nocera DG. *J Am Chem Soc.* 1992; 114:4013–4015.
55. Young ER, Rosenthal J, Hodgkiss JM, Nocera DG. *J Am Chem Soc.* 2009; 131:7678–7684. [PubMed: 19489645]
56. DeRege PJF, Williams SA, Therien MJ. *Science.* 1995; 269:1409–1413. [PubMed: 7660123]
57. Cukier RI. *J Phys Chem.* 1994; 98:2377–2381.
58. Irebo T, Reece SY, Sjödin M, Nocera DG, Hammarström L. *J Am Chem Soc.* 2007; 129:15462–15464. [PubMed: 18027937]
59. Bonin J, Constantin C, Louault C, Robert M, Savéant J-M. *J Am Chem Soc.* 2011; 133:6668–6674. [PubMed: 21476550]
60. Offenbacher AR, Watson RA, Pagba CV, Barry BA. *J Phys Chem B.* 2014; 118:2993–3004. [PubMed: 24606240]
61. Yokoyama K, Uhlin U, Stubbe J. *J Am Chem Soc.* 2010; 132:15368–15379. [PubMed: 20929229]
62. Climent I, Sjöberg BM, Huang CY. *Biochemistry.* 1992; 31:4801–4807. [PubMed: 1591241]
63. Argirevi T, Riplinger C, Stubbe J, Neese F, Bennati M. *J Am Chem Soc.* 2012; 134:17661–17670. [PubMed: 23072506]
64. Marcus RA, Sutin N. *Biochim Biophys Acta.* 1985; 811:265–322.
65. Gray HB, Winkler JR. *Annu Rev Biochem.* 1996; 65:537–561. [PubMed: 8811189]
66. Krishtalik LI. *Biochim Biophys Acta.* 2011; 1807:1444–1456. [PubMed: 21777562]
67. Meade TJ, Gray HB, Winkler JR. *J Am Chem Soc.* 1989; 111:4353–4356.
68. Crane BR, Di Biolio AJ, Winkler JR, Gray HB. *J Am Chem Soc.* 2001; 123:11623–11631. [PubMed: 11716717]
69. McLendon G, Miller JR. *J Am Chem Soc.* 1985; 107:7811–7816.
70. Jiang N, Kuznetsov A, Nocek JM, Hoffman BM, Crane BR, Hu X, Beratan DN. *J Phys Chem B.* 2013; 117:9129–9141. [PubMed: 23895339]
71. Moser CC, Keske JM, Warnke K, Farid RS, Dutton LP. *Nature.* 1992; 355:796–802. [PubMed: 1311417]
72. Co NP, Young RM, Smeigh AL, Wasielewski MR, Hoffman BM. *J Am Chem Soc.* 2014; 136:12730–12736. [PubMed: 25133915]
73. Davidson VL. *Acc Chem Res.* 2000; 33:87–93. [PubMed: 10673316]
74. Song DY, Pizano AA, Holder PG, Stubbe J, Nocera DG. *Chem Sci.* 2015; 6:4519–4524. [PubMed: 26504513]
75. Lin J, Balabin IA, Beratan DN. *Science.* 2005; 310:1311–1313. [PubMed: 16311331]
76. Carra C, Iordanova N, Hammes-Schiffer S. *J Am Chem Soc.* 2003; 125:10429–10436. [PubMed: 12926968]
77. Young ER, Rosenthal J, Nocera DG. *Chem Commun.* 2008:2322–2324.

78. Sjödin M, Irebo T, Utas JE, Lind J, Merényi G, Åkermark B, Hammarström L. *J Am Chem Soc.* 2006; 128:13076–13083. [PubMed: 17017787]
79. Constantin C, Robert M, Savéant J-M. *J Am Chem Soc.* 2007; 127:9953–9963.

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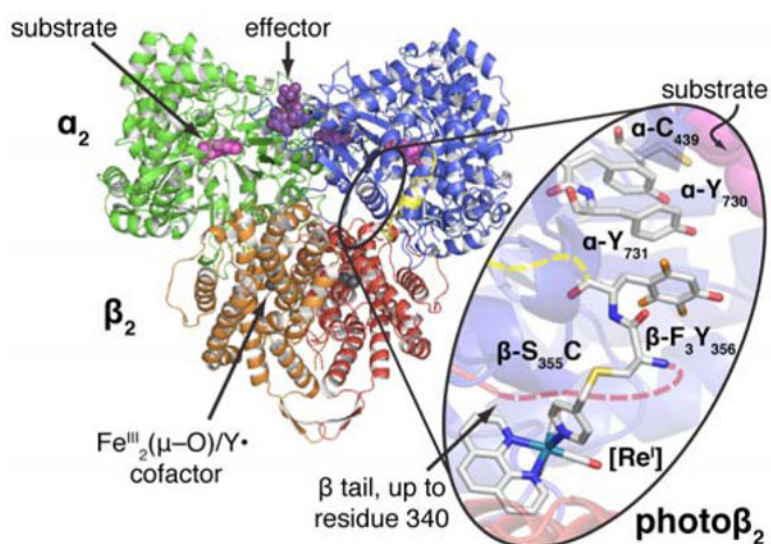


Figure 1.

Docking model of the active *E. coli* class Ia RNR, an $\alpha_2\beta_2$ complex. α_2 (blue and green) co-crystallizes with a peptide corresponding to the 15 C-terminal residues of β (yellow) (ref 35). β_2 crystallizes up to residue 340 (red and orange) (ref 36). Photo β_2 s are prepared via cysteine ligation of a [Re^I] complex at position 355, and incorporation of a fluorotyrosine at position 356 (inset). Due to the absence of structural information for residues 341–359 of β , attachment of the [Re^I]-S₃₅₅C-F₃Y₃₅₆ fragment is illustrated by dashed lines (red leading from the crystalline β subunit, yellow leading to the C-terminal peptide bound to α).

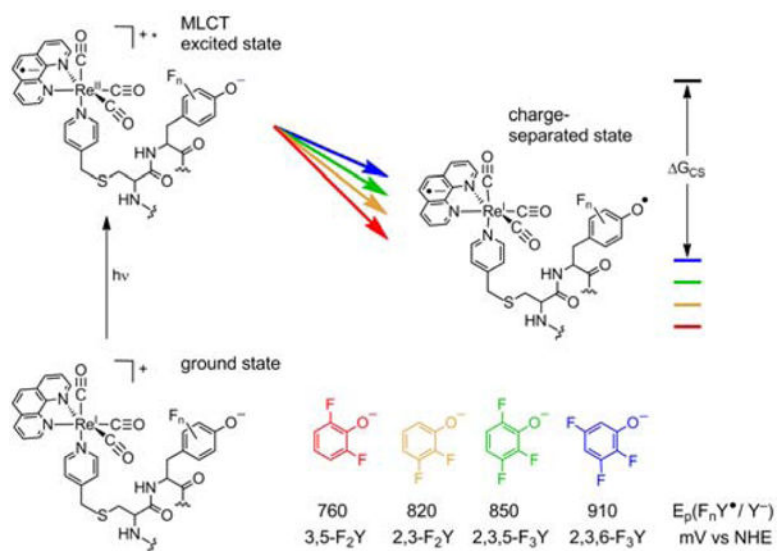


Figure 2. Photooxidation of β -F_nY₃₅₆ by the metal-to-ligand charge transfer (MLCT) excited state of [Re^I]^{*}. Variation of the driving force for this process is achieved by incorporation of various F_nYs at position β_{356} . Peak potentials are taken from differential pulse voltammetry performed on the *N*-acetyl *C*-amide-protected amino acid derivatives.

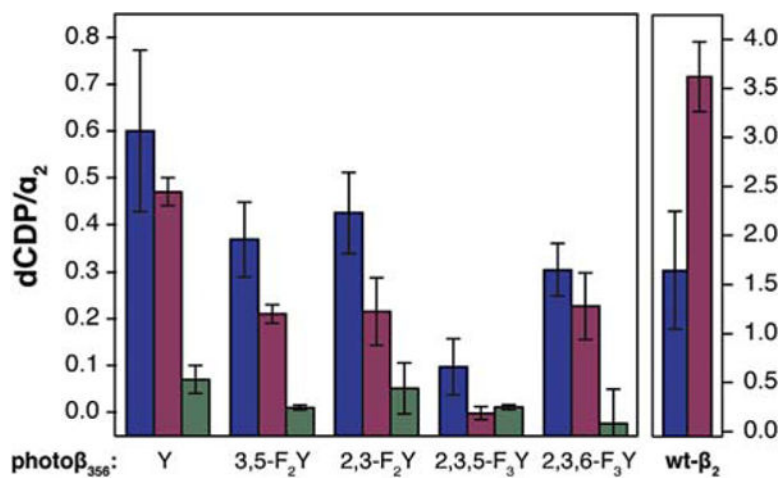


Figure 3. Photochemical single turnover experiments in the presence (blue) and absence (purple) of 10 mM Ru(NH₃)₆Cl₃. Control experiments (green) were performed in the dark. Samples contained 10 μ M (blue) or 20 μ M (purple and green) of photo- or wt- β_2 as indicated along the x-axis, 10 μ M wt- α_2 , 200 μ M [5-³H]-CDP with 27,000 cpm/nmol radioactivity, 3 mM ATP, in assay buffer at pH 7.6. Error bars represent 1 s.d. for 3 independent trials.

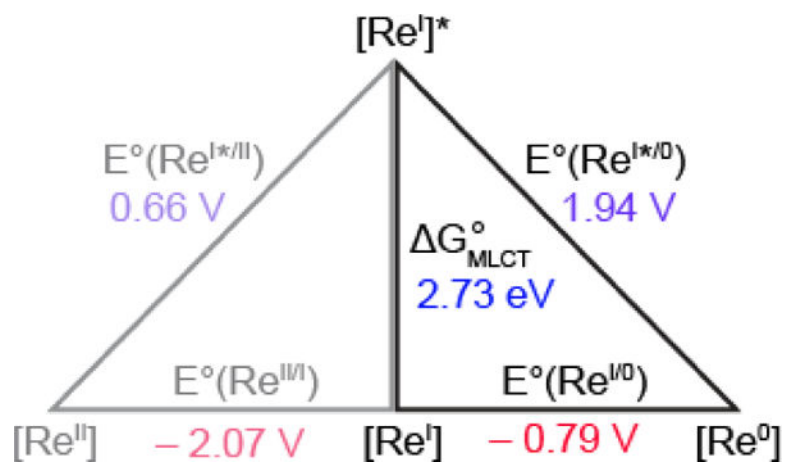


Figure 4. Latimer diagram describing excited state reduction potentials of $[\text{Re}^{\text{I}}]$. G_{MLCT} determined from the $E_{0/0}$ emission of $[\text{Re}]-\text{Y}_{356}\text{F}-\beta_2$ frozen in assay buffer at 77 K (Figure S4). Ground state reduction potentials are from ref 50. Values along the diagonal are the calculated excited state reduction potentials.

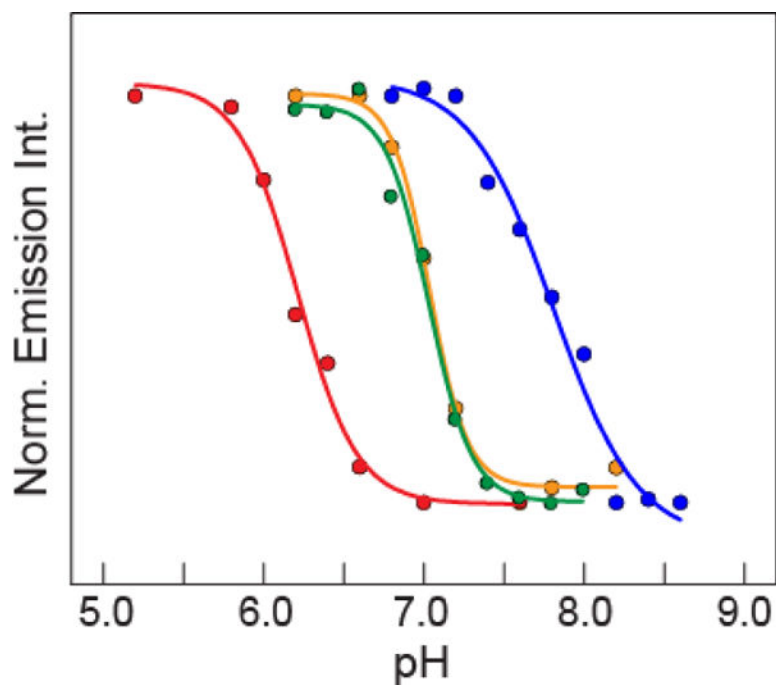


Figure 5. Steady state emission titrations of F_nY₃₅₆ within photoβ₂α₂ complexes are plotted as normalized integrated emission intensity measured over 450–650 nm for 2,3,5-F₃Y₃₅₆ (●); 3,5-F₂Y₃₅₆ (●); 2,3,6-F₃Y₃₅₆ (●); 2,3-F₂Y₃₅₆ (●) and fit to Eq. 6 (lines). Samples contained 5 μM photoβ₂, 20 μM wt-α₂, 1 mM CDP, 3 mM ATP, in assay buffers described in the Methods section, held at 25 °C. The solid lines are the result of fitting the data to Eq. 6.

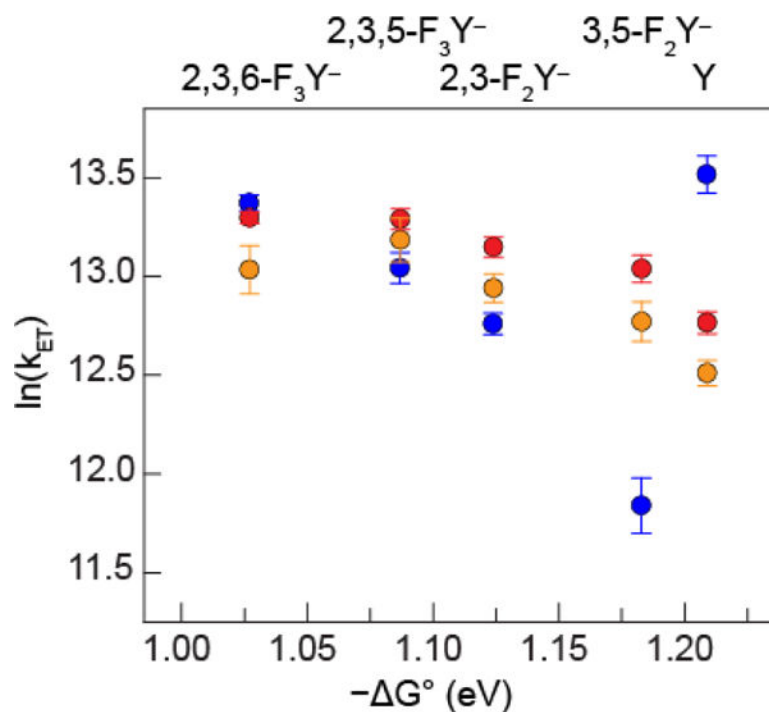


Figure 6.

Correlation of the natural log of k_{ET} and G° for photo β_2 s alone (\bullet), or in the presence of wt- α_2 (\bullet), or Y₇₃₁F- α_2 (\bullet). k_{ET} and G° were calculated according to Eq.s 2 and 5, respectively. Triplicate sets of independently prepared samples contained 10 μ M F_nY₃₅₆-photo β_2 ($n = 0-3$) or Y₃₅₆F-photo β_2 , 25 μ M wt- α_2 or Y₇₃₁F- α_2 , 1 mM CDP, 3 mM ATP in assay buffer at pH 8.2.

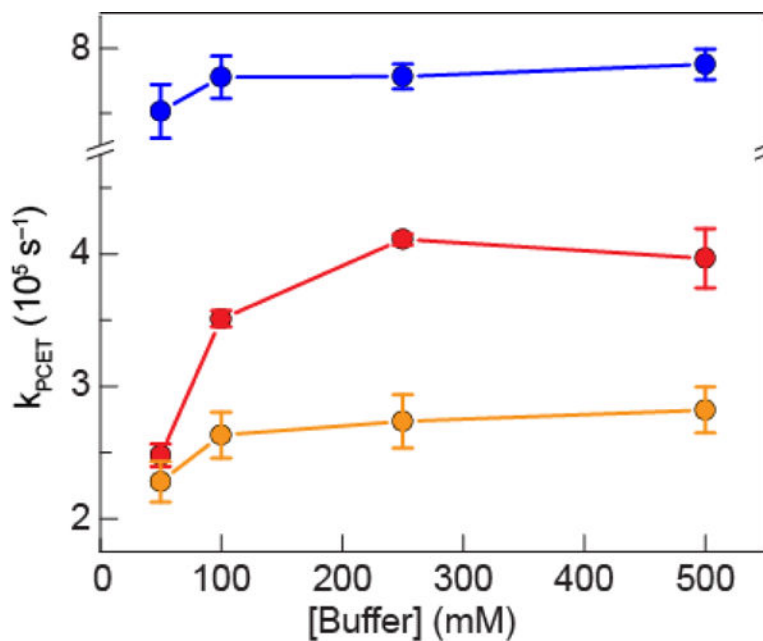


Figure 7. Dependence of k_{PCET} on buffer concentration for photo β_2 s alone (●), or in the presence of wt- α_2 (●), or Y₇₃₁F- α_2 (●). k_{PCET} was calculated according to Eq. 2. Triplicate sets of independently prepared samples contained 10 μM Y₃₅₆- or Y₃₅₆F-photo β_2 , 25 μM wt- α_2 or Y₇₃₁F- α_2 , 1 mM CDP, 3 mM ATP in assay buffer containing 15 mM MgSO₄, 1 mM EDTA, and 50, 100, 150, or 500 mM HEPES buffer adjusted to pH 7.6.

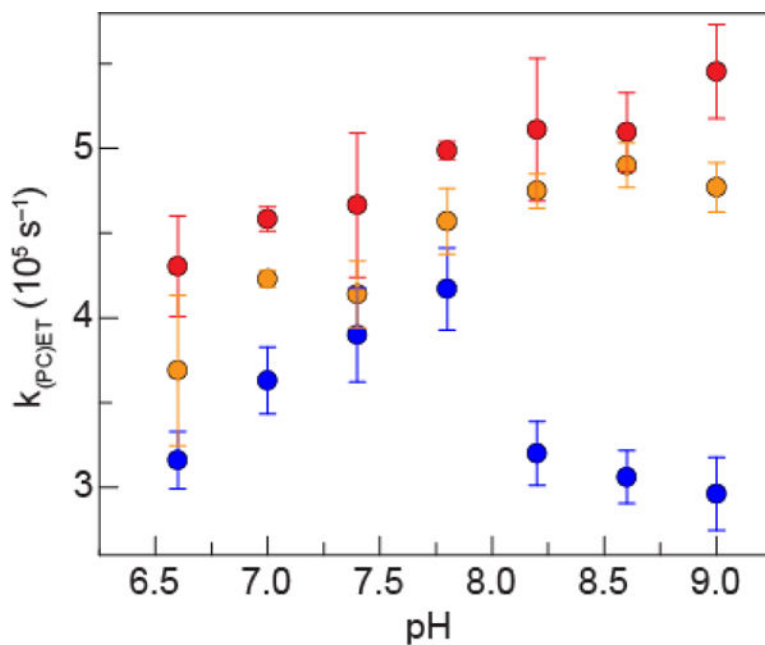


Figure 8. Dependence of $k_{(PC)ET}$ on pH for 2,3-F₂Y-photoβ₂ alone (●), or in the presence of wt-α₂ (●), or Y₇₃₁F-α₂ (●). $k_{(PC)ET}$ was calculated according to Eq. 2. Duplicate sets of independently prepared samples contained 10 μM 2,3-F₂Y₃₅₆-photoβ₂ or Y₃₅₆F-photoβ₂, 25 μM wt-α₂ or Y₇₃₁F-α₂, 1 mM CDP, 3 mM ATP in assay buffer.

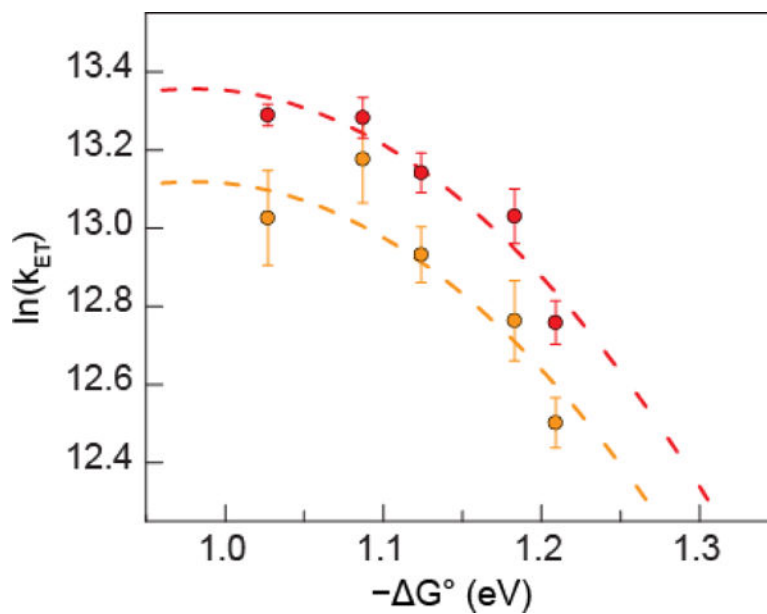


Figure 9. Correlation of the natural $\ln(k_{ET})$ with $-\Delta G^\circ$ for photo β_2 in the presence of wt- α_2 (●), or Y731F- α_2 (●). Dashed lines represent simulations of Eq. 8 for β -FnY356S in the presence of wt- α_2 (---) and Y731F- α_2 (---), with $r = 12.5 \text{ \AA}$, $\lambda = 0.98 \text{ eV}$ and $H_{AD} = 0.051$ and 0.044 cm^{-1} in the presence of wt- and Y731F- α_2 , respectively.

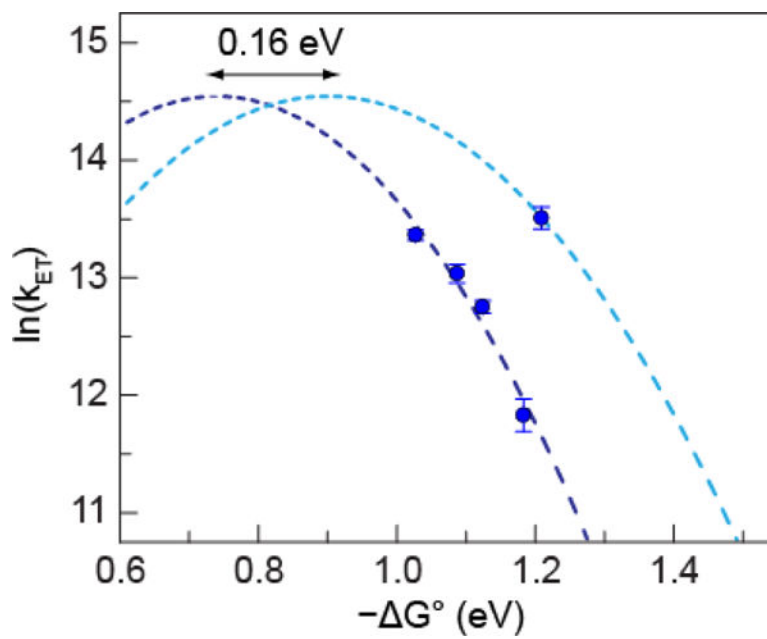
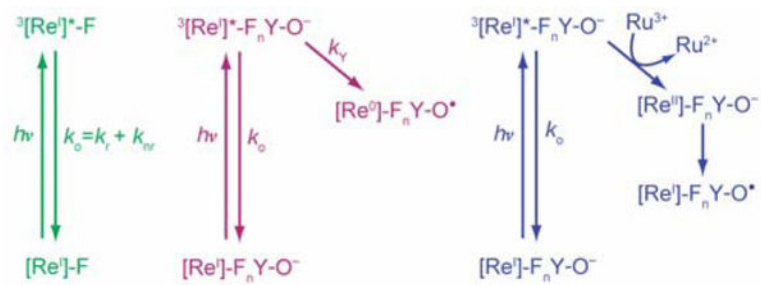


Figure 10.

Correlation of the $\ln(k_{ET})$ with $-\Delta G^\circ$ for $\text{photo}\beta_2$ in the absence of α_2 (\bullet). Dashed lines represent simulations of Eq. 8 for $\text{photo}\beta\text{-F}_n\text{Y}_{356}$ (---) and $\text{photo}\beta\text{-Y}_{356}$ (---) with $\lambda = 0.74$ eV and 0.90 eV ($r = 12.5$ Å, $H_{AD} = 0.083$ cm $^{-1}$), respectively.



Scheme 1.
Photochemical generation of $\text{F}_n\text{Y}\bullet$

Table 1

Dependence of k_{PCET} on oligomeric state

Effector	Interface Residues		τ^a (ns)	k_{PCET}^b (10^5 s^{-1})	Pathway Enhancement
	β_{356}	α_{731}			
ATP	Y	Y	522 (8)	4.8 (3)	34(6)%
	F	Y	696 (3)		
	Y	F	578 (9)	3.6 (3)	
	F	F	728 (5)		
dATP	Y	Y	505 (10)	5.5 (4)	statistically identical
	F	Y	698 (2)		
	Y	F	511 (14)	5.3 (6)	
	F	F	699 (8)		

^a Triplicate sets of independently prepared samples contained 10 μM Y356- or Y356F-photocp2, 25 μM wt- or Y731F- α_2 , and either 1 mM CDP or 0.2 mM dATP, in assay buffer at pH 7.6.

^b Calculated according to Eq. 2.

Table 2p*K*_a values of F_nYs within the photoRNR complex

β -Y ₃₅₆ Z	p <i>K</i> _a , complex	p <i>K</i> _a , free F _n Y ^b
2,3,5-F ₃ Y	6.2 ^a	6.4
3,5-F ₂ Y	7.0	7.2
2,3,6-F ₃ Y	7.0	7.0
2,3-F ₂ Y	7.6	7.8

^aData reproduced from reference 24.^bData reproduced from reference 37, determined for *N*-acetyl *C*-amide protected fluortyrosine derivatives.

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Table 3

Solvent kinetic isotope effect on k_{PCET}

Interface residues	τ (ns) ^a	k_{PCET} (10^5 s^{-1}) ^b				SKIE
		¹ H	² H	¹ H	² H	
β_{356}						
α_{731}						
Y	410 (20)	568 (5)	8 (1)	5.5 (2)	1.5 (2)	
F	615 (5)	827 (9)				
Y	522 (8)	713 (3)	4.8 (3)	3.2 (1)	1.4 (1)	
F	696 (3)	920 (2)				
Y	578 (9)	778 (6)	3.6 (3)	2.4 (1)	1.5 (1)	
F	728 (5)	952 (8)				

^a Triplicate sets of independently prepared samples contained 10 μM Y356 or Y356F-photo β 2, 25 μM α 2 or Y731F- α 2, 1 mM CDP, 3 mM ATP in assay buffer at pH 7.6.^b Calculated according to Eq. 2.