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Re(bpy)(CO)$_3$CN as a Probe of Conformational Flexibility in a Photochemical Ribonucleotide Reductase†

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Abstract

Photochemical ribonucleotide reductases (photoRNRs) have been developed to study the proton-coupled electron transfer (PCET) mechanism of radical transport in class I E. coli ribonucleotide reductase (RNR). The transport of the effective radical occurs along several conserved aromatic residues across two subunits: β2(•Y122→W48→Y356)→α2(Y731→Y730→C439). The current model for RNR activity suggests that radical transport is strongly controlled by conformational gating. The C-terminal tail peptide (Y-βC19) of β2 is the binding determinant of β2 to α2 and contains the redox active Y356 residue. A photoRNR has been generated synthetically by appending a Re(bpy)(CO)$_3$CN ([Re]) photo-oxidant next to Y356 of the 20-mer peptide. Emission from the [Re] center dramatically increases upon peptide binding, serving as a probe for conformational dynamics and Y356 protonation state. The diffusion coefficient of [Re]-Y-βC19 has been measured ($k_d$ = 6.1 × 10$^{-7}$ cm$^{-1}$ s$^{-1}$), along with the dissociation rate constant for the [Re]-Y-βC19:α2 complex ($> k_{off} > 400$ s$^{-1}$). Results from detailed time-resolved emission and absorption spectroscopies reveal biexponential kinetics, suggesting a large degree of conformational flexibility in the α2:[Re]-Y-βC19 complex that partitions the N-terminus of the peptide into both bound and solvent-exposed fractions.

Amino acid radicals often serve as cofactors for substrate activation and vehicles for charge transport in enzymes (1). Free amino acid radicals in solution typically have a micro- to milli-second lifetime. Yet Nature has evolved enzymes to manage both proton and electron equivalents so that the oxidative power of amino acid radical intermediates may be harnessed for chemical transformations over disparate timescales (2). Oxidation of amino acids at physiological pH usually involves the loss of both a proton and electron, implicating proton-coupled electron transfer (PCET) as the redox mechanism (3-7). In a PCET event, proton tunneling is limited to short distances, usually within hydrogen bonds, whereas the electron, as the lighter particle, may tunnel over tens of ångströms (8-10). Enzymes have evolved to manage the disparate distance requirements of proton and electron tunneling to achieve efficient radical transport and catalysis (11).

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Supporting Information Available. Transient absorption spectra of the [Re]-F-βC19 and [Re]-Y-βC19 peptides in aqueous solution, SF traces, and MP-FRaP trace. This material is available free of charge via the Internet at http://pubs.acs.org.
Ribonucleotide reductase (RNRs) serves as an exemplar for enzymes that derive their function from amino acid radicals (12). Class I *E. coli* RNR contributes to DNA replication and repair by catalyzing the reduction of nucleoside diphosphates (NDPs) to deoxynucleoside diphosphates (dNDPs) (1, 13). The enzyme comprises two homodimeric subunits designated α2 and β2, and a complex between the two is required for enzyme activity (14). α2 houses the NDP/NTP substrate/effector binding sites that control the specificity and rate of nucleotide reduction (15). β2 harbors a diferric tyrosyl radical (α-Y122) cofactor proposed to initiate nucleotide reduction by oxidizing a cysteine residue (C439) in the active site of α2 (16). Radical transport from α-Y122 to C439 occurs over 35 Å (17) via a PCET amino acid radical transport pathway involving conserved amino acid residues β2(α-Y122→W48→Y356)→α2(Y731→Y730→C439) (12, 18).

Photochemical RNRs (photoRNRs) have been designed to probe the intrinsic mechanistic steps attendant to radical transport between the tyrosines of the β2 to α2 subunits (Y356→Y731) and along the α2(Y731→Y730→C439) pathway (19). In this construct, α2 subunits are bound to the 20-mer, C-terminal peptide tail of β2 (Y-βC19) containing a chemically appended photo-oxidant (20-22). Y-βC19 is known to contain both the binding determinant of β2 to α2 (23, 24) and the redox active Y356 residue that facilitates radical transport between the two subunits (25-27). Figure 1 shows the X-ray crystal structure of α2 subunits highlighting the putative residues that comprise the radical transport pathway in this subunit, along with the bound Y-βC19 peptide (18). The N-terminal residues of Y-βC19, including the redox-active “Y356” residue, are not located in the structure. Excitation of the photo-oxidant appended proximal to Y356 on the peptide produces α-Y356 (21, 22). When bound to the α2 subunit in the presence of RNR substrate and effector, this photo-triggered radical can be transported into the α2 active site to initiate deoxynucleotide production. PhotoRNRs designed to date have been based on different photo-oxidants of tyrosine including Trp (28), the benzophenone-containing unnatural amino acid (BPA) (21, 29), anthraquinone (Anq) (21), and the inorganic complex Re3(bpy)(CO)3CN ([Re]) (30, 31). Of these various photoRNRs, the [Re]-based RNR affords the highest single-turnover yield (22). In model systems with tyrosine appended to the bpy ligand of [Re], the MLCT excited state of [Re] is only a competent oxidant of the deprotonated, tyrosinate anion (Y−, pH >10) forming the [Re3-Y•] charge separated state (30). Fluorotyrosine unnatural amino acids may be used to lower the pKa of the phenolic proton (29) in order to obtain the tyrosinate form of the amino acid and thus enable photoactivity within the pH window for RNR activity (6 < pH < 8.5) (31). We have directly observed a 3,5-F2Y• radical on a [Re]-3,5-F2Y-βC19 peptide and showed that the peptide bound to the Y731F-α2 protein is competent for initiating RNR turnover at pH 7.5 and 8.2 (22).

With radical generation and competency established at the Y356 position, we have turned our attention toward a thorough characterization of Y356 on the peptide. We now report the utilization of the emission of the [Re]* excited state within the [Re]-Y-βC19:α2 complex to probe conformational dynamics of the N-terminus of the Y-βC19 peptide bound to α2. We show that the [Re]* emission blue-shifts and increases in intensity upon binding to α2.

![Biochemistry. Author manuscript; available in PMC 2012 April 30.](image)
modulation of the [Re]* emission intensity, as the deprotonated 3,5-F₂Y⁻ quenches the
[Re]* excited state via electron transfer. Plots of the [Re]* emission intensity vs. pH allow
for determination of the pKₐ of 3,5-F₂Y on the peptide bound to α2. Stopped-flow and
photobleaching experiments provide information on diffusion and binding rate constants
between the peptide and the protein. Time-resolved emission spectroscopy reveals multiple
populations of the [Re]* excited state. These data, in combination with the X-ray data shown
in Figure 1, support a model for a conformationally flexible N-terminus of the Y-βC19
peptide bound to α2.

Materials and Methods

Materials

The peptides [Re]-F-βC19, [Re]-3,5-F₂Y-βC19 and [Re]-Y-βC19 were available from a
previous study (22), and the synthesis of Ac-Y-βC19 has been described (20). The
concentration of each [Re]-peptide stock solution was estimated by UV-vis absorption
spectra using the known ε₃₅₅ = 5300 M⁻¹ cm⁻¹ for the [Re]-3,5-F₂Y-OMe dipeptide (31)
and the concentration of Ac-Y-βC19 was estimated using ε₂₈₀ = 1197 M⁻¹ cm⁻¹ for
tyrosine (32). E. coli thioredoxin (TR, SA of 40 U/mg), E. coli thioredoxin reductase (TRR,
SA of 1800 U/mg) and E. coli β₂ (SA of 6800 nmol/min mg) were isolated as previously
described (33). E. coli α₂ was isolated, purified, and pre-reduced according to an established
procedure (21).

Steady-State Emission Spectroscopy

Steady state emission spectra were recorded on an automated Photon Technology
International (PTI) QM 4 fluorimeter equipped with a 150-W Xe arc lamp and a Hamamatsu
R928 photomultiplier tube. Solutions (200 μL) containing [Re]-peptides and
α₂ were placed
in a 2 × 10 mm quartz micro-cuvette and excited with 315 nm light focused along the 10
mm pathlength. Emission was collected at 90° to the excitation source and passed through a
450 nm long-pass filter prior to entering the detector housing. Emission spectra were
recorded as a function of pH for solutions containing 20 μM α₂, 5.5 μM [Re]-3,5-F₂Y-
PC19, 50 mM buffer (either KPi or Tris), 15 mM Mg²⁺, 3 mM ATP, and 1 mM CDP.
Phosphate and Tris buffers were used in the pH range 6.3-7.0 and 7.2-8.9, respectively.

Stopped-Flow Kinetics

Stopped-flow (SF) kinetic experiments were performed on an Applied Photophysics DX.
17MV equipped with an emission PMT. Samples were excited with λ = 355 nm and
emission detection was limited to λ > 455 nm using a long-pass cut-off filter from Applied
Photophysics LTD. The temperature was maintained at 9 °C with a Lauda RE106 circulating
water bath. All experiments were performed in assay buffer (50 mM Tris, 15 mM MgSO₄, 3
mM ATP, 1 mM CDP, pH = 7.6). Prior to each experiment, the instrument was flushed with
copious amounts of water and then with the assay buffer. [Re]-F-βC19 in one syringe was
mixed with assay buffer in another syringe in equal volumes to yield a final concentration of
45 μM [Re]-F-βC19. This experiment provided the baseline [Re] emission of the [Re]-F-
βC19 peptide. [Re]-F-BC19 and α₂ in one syringe were mixed with assay buffer in another
syringe in equal volumes to yield a final concentration of 45 μM [Re]-F-βC19 and 1 μM α₂.
The peptide emits more intensely when bound to the protein (vide infra); hence this
experiment provided the maximum emission expected from [Re] when the peptide was
bound to α₂. Finally, [Re]-F-βC19 and α₂ in one syringe were mixed with Ac-Y-βC19 in
another syringe in equal volumes to yield a final concentration of 45 μM [Re]-F-βC19 and 1
μM α₂, 450 μM Ac-Y-βC19. 2000 evenly spaced data points were collected over 50 ms or
20 s.
Analysis of [Re]-Y-βC19 binding to α2

Binding of the [Re]-Y-βC19 peptide to α2 was monitored by steady state emission spectroscopy. Solutions of 5 μM peptide were titrated with α2 from concentrations of 0 → 65 μM. Emission from [Re] did not significantly change at higher concentrations of α2; thus at 65 μM we concluded that each molecule of α2 was bound to two [Re]-Y-βC19 peptide.

The fraction of bound peptide as a function of α concentration, \( \chi_{\text{bound}}(\alpha) \), was then calculated from the equation:

\[
\chi_{\text{bound}}(\alpha) = \frac{I_{600\text{nm}}(\alpha) - I_{600\text{nm}}(0)}{I_{600\text{nm}}(130\mu\text{M}) - I_{600\text{nm}}(0)}
\]

where \( I_{600\text{nm}} \) is the emission intensity at 600 nm. Data were fit to the following simplified binding model:

\[
αP \xrightarrow{K_D} α + P
\]

where P = [Re]-Y-βC19. From this model we can derive an expression for \( \chi_{\text{bound}} \) (34):

\[
\chi_{\text{bound}} = \frac{1}{1 + \frac{K_D}{\alpha}}
\]

assuming that \( K_D \) is the same for binding of the first and second peptide to α2, as previously discussed (23). \( K_D \) is derived from a plot of \( \chi_{\text{bound}} \) vs. [α] and fitting the data to eq. 3.

Diffusion Coefficients

The diffusion coefficient of [Re]-Y-βC19 (3.65 mM concentration) was determined using Multi-Photon Fluorescence Recovery after Photobleaching (MP-FRaP) (35). The excitation source was a tunable mode-locked Ti:sapphire laser (Mai-Tai Broadband, Spectra-Physics, Mountain View, CA) at 800 nm. The beam was fed into a custom-built multiphoton microscope containing an Axioskop upright microscope (Carl Zeiss, Inc., Thornwood, NY) and Bio-Rad MRC-600 confocal laser scanner (Carl Zeiss, Inc., Thornwood, NY). For each measurement, the laser was focused with a 40× (0.75NA, Zeiss) water-immersion objective to a fixed spot (non-scanning) in the sample at low power (\( \sim 5-10 \) mW at the sample) to monitor fluorescence in the absence of photobleaching. Photons were detected with a photon-counting photomultiplier tube (H7421-40, Hamamatsu, Inc., Bridgewater, NJ) and binned using a multichannel scaler and averaged (SR430, Stanford Research Systems, Sunnyvale, CA). An electro-optic modulator (KD*P Pockels cell, ConOptics, Danbury, CT) and digital delay generator (DG535, Stanford Research Systems, Sunnyvale, CA) were used to rapidly modulate the laser power using custom-built hardware and a user interface developed in LabView (National Instruments, Austin, TX). In a single bleaching pulse, the laser power was increased briefly (<100 μs) to produce a more intense pulse of light (\( \sim 40-60 \) mW at the sample), which caused irreversible photobleaching in the focal volume. The laser power was then returned to its low original value and fluorescence recovery was monitored over several milliseconds. Due to the low concentration of fluorophores within the focal volume, the bleach and recovery process was repeated hundreds of times and fluorescence counts were summed into a single measurement to improve signal to noise. Diffusion coefficients were obtained by fitting a model one-component recovery curve to the change in amplitude of fluorescence over time with non-linear least squares fitting incorporating a trust-region algorithm (35).
The diffusion coefficient of α2 (8 μM concentration) was determined by dynamic light scattering (DLS) performed on a DynaPro Dynamic Light Scatterer. The laser power and temperature were set to 37% and 25 °C, respectively. Each autocorrelation function (ACF) was acquired for 10 s and averaged over 10 measurements. The resulting ACF was fit using Dynamics V6 (Version 6.7.7.9) software from Wyatt Technology Corp. employing a non-negative least-squares fitting algorithm. Hydrodynamic radii were obtained from a mass-weighted size distribution analysis and the reported value is the average of three measurements. The diffusion coefficient of α2, \( k_{d2} \), was calculated with the Einstein-Stokes relationship,

\[
k_{d2} = \frac{k_B T}{6\pi \eta r}
\]

where \( k_B \) is Boltzmann’s constant, \( T \) is temperature (25 °C), \( \eta \) (~1 cP) is the viscosity of the solution, and \( r \) is the radius of α2. The radius obtained from DLS experiments was compared to the radius equivalent to a sphere with equal volume to α2,

\[
r = \frac{\sqrt[3]{d_1 d_2 d_3}}{2}
\]

where \( d_1, d_2, \) and \( d_3 \) are the diameters of α2 in the x, y, and z directions (36). Eq. (5) is obtained from equating the volume of a sphere with the volume of an ellipsoid and solving for \( r \).

**Analysis of [Re]-Y-βC19 Binding to α2**

The diffusion limited rate constant, \( k_D \), was determined from (37),

\[
k_D = 4\pi (k_{d1} + k_{d2}) R
\]

where \( k_{d1} \) and \( k_{d2} \) are the diffusion coefficients of the [Re]-Y-βC19 and α2, respectively, and \( R \) is the estimated radius of the binding site on α2. The binding site of the [Re]-Y-βC19:α2 complex was estimated from the crystal structure of α2 with Ac-Y-19mer (18). Only the 16 residues observed were used and an effective radius (8.97 Å) was calculated from the x, y, and z dimensions of the observed binding site (36.4 Å, 15.4 Å, and 10.3 Å) using eq 5.

**Time-Resolved Emission and Transient Absorption**

Time resolved emission measurements on the <20 ns timescale were made with the frequency doubled (400 nm) pump light provided by a Ti:sapphire laser system (100 fs pulsewidth) and collected on a Hamamatsu C4334 Streak Scope streak camera (38). Time resolved emission measurements on the >20 ns timescale were made with pump light provided by the third harmonic (355 nm) of an Infinity Nd:YAG laser (Coherent) running at 20 Hz (39). Nanosecond transient absorption (TA) measurements were made with the same laser running at 10 Hz. The sample preparation and experimental setup were identical to that previously described (22).

To produce a transient absorption (TA) spectrum, the series of four spectra were taken: \( I_F \) (pump on/probe off), \( I \) (pump on/probe on), \( I_B \) (pump off/probe off), and \( I_0 \) (pump off/probe on). Transient spectra were corrected for fluorescence and background light using these spectra by the calculation: \( \Delta OD = \log ([I_0 - I_B]/(I - I_F)) \). For experiments involving α2, the
spectra reported are an average of 125 of the four-spectra sequences. Sample sizes were typically 200 μL in a 2 x 10 mm cuvette containing a Teflon-coated mini-stirbar. Both the white light and pump beams were focused and overlapped to pass through the 2mm-wide window of the cuvette, providing a total pathlength of 1 cm. To provide optimal beam overlap, the pump beam was reflected off a small mirror in front of the collimating lens for the probe beam after the sample.

Results

Photophysics and Photochemistry of [Re]-X-βC19 Peptides

The quenching of the 3[Re]* excited state in the 3[Re]-3,5-F₂Y-βC19 peptide can be investigated with steady state emission spectroscopy. Figure 2 plots the emission spectra of the [Re] center as the pH of a solution of [Re]-3,5-F₂Y-βC19 peptide is varied from 4.5 to 11.5. Between pH 4.5-5.0 the emission red-shifts and decreases in intensity. We ascribe these spectral changes to an alteration of the peptide conformation, as the carboxylic acid groups of Asp and Glu residues are typically deprotonated in this pH regime. Between pH 5 and 11.5, the emission energy and band shape do not change, however, the overall intensity of the band decreases. This emission quenching is consistent with deprotonation of the 3,5-F₂Y phenolic proton, which induces electron transfer from 3,5-F₂Y− to the proximate 3[Re]* excited state (31). By titrating the 3[Re]* emission intensity with pH, we obtain a pKₐ value of 3,5-F₂Y when incorporated into the peptide of 7.3 ± 0.1, which is comparable to the pKₐ = 7.2 of the Ac-3,5-F₂Y-NH₂ amino acid in solution (29).

The emission of the [Re] excited state on [Re]-(F/Y/3,5-F₂Y)-βC19 can be time-resolved to provide insight into the conformational homogeneity of the peptide. The emission decay of [Re]-3,5-F₂Y-βC19 at pH 10 fits to a bi-exponential decay function with τ₁ = 1.9 ns (66%) and τ₂ = 10.7 ns (34%) as does the emission decay of [Re]-Y-βC19 at pH 12 (τ₁ = 0.5 ns (86%) and τ₂ = 3.0 ns (14%). Both lifetime components for each peptide are faster compared to the emission lifetime of the [Re]-F-βC19 control peptide at all pHs or [Re]-Y-βC19 at pH < 10 (τ = 50 ± 2 ns or 62 ± 2 ns).

Binding of the [Re]-Y-βC19 Peptide to α2

Peptide binding to α2 may be probed by monitoring the emission energy of the 3[Re]* excited state. Figure 3 plots the emission spectra of a 5 μM solution of the [Re]-Y-βC19 peptide as the α2 concentration is increased from 0→65 μM. Upon binding to α2, the emission band blue shifts and increases in intensity. This is consistent with partial desolvation of the [Re] chromophore upon binding to the protein (40, 41). Because there are two binding sites on each α2 subunit (one on each α in the dimer) as shown in the crystal structure (18), the inset of Figure 3 plots the emission maxima as a function of α concentration. The fit of these data to the binding model described by eq 3 yields a dissociation constant, K_D, of 9 μM. This value of K_D is comparable to the value obtained from competitive inhibition assay (23, 24) measurements of the [Re]-Y-βC19 peptide to α2 (IC₅₀ = 8 μM ≈ K_D (22)).

pKₐ of 3,5-F₂Y356 Bound to α2

The pKₐ of the 3,5-F₂Y on the peptide bound to α2 may be ascertained by monitoring the emission intensity and lifetime of the 3[Re]* excited state. Figure 4 plots the emission intensity of a 5.5 μM solution of [Re]-3,5,-F₂Y-βC19 in the presence of 20 μM of α2 as the pH is increased from 6.3 to 9.0. The grey spectra in the figure correspond to the emission of 5 μM [Re]-3,5,-F₂Y-βC19 in the absence of α2 at pH 6.3 (lower spectrum) and 9.0 (upper spectrum) and re-emphasize the effect of peptide binding on the [Re] emission band both at pH 6.3 and pH 9.0. As the pH is titrated from 9.0 to 6.3 for the solution containing the
bound peptide, the $^3[\text{Re}]^*$ emission intensity decreases with an apparent pK$_a$ of 7.3 ± 0.1. The same pK$_a$ was obtained whether the integrated emission or emission maximum (at 600 or 650 nm) was plotted vs. pH. As was observed in Figure 2, deprotonation of the phenol of 3,5-F$_2$Y allows for electron transfer to occur between 3,5-F$_2$Y$^-$ and $^3[\text{Re}]^*$ resulting in a decrease in emission intensity. We thus attribute the emission quenching in Figure 4 to electron transfer between $^3[\text{Re}]^*$ and 3,5-F$_2$Y$^-$ on the peptide bound to α2. Thus the pK$_a$ of the bound 3,5-F$_2$Y does not shift significantly from that of the peptide in free solution.

**Kinetic Analysis of [Re]$^*$ Decay in [Re]-(F/Y)-βC19:α2 Complexes**

Time-resolved emission monitoring the decay kinetics from the [Re] excited state ([Re]$^*$) was used to further probe the homogeneity of the peptide:α2 binding interaction. For this experiment, 5 μM peptide and 20 μM α2 (with 3mM ATP and at pH 7.5) were used to ensure that most of the peptide was bound. All experiments were performed in buffered solutions containing 20% glycerol, as this was previously shown to increase the yield of photochemical RNR turnover (22). Figure 5 plots the emission decay trace obtained at 600 nm for 5 μM [Re]-Y-βC19 and 20 μM α2. The same trace was obtained in the presence of 1mM CDP substrate. Under all conditions investigated, the data were best fit with a bi-exponential decay function. One emission decay component was set equal to that of the free peptide while the other component was allowed to vary. We obtained decay components of 60 ns (63%) and 150 ns (37%), which were unaffected by the addition of 1 mM CDP substrate. These data reveal that the conformation of the peptide bound to α2 is heterogeneous, with multiple binding modes. For some binding modes, the [Re] unit may be solvated similar to that in free solution, while for others the [Re] chromophore is partially desolvated resulting in a blue-shifted emission energy with longer emission lifetime.

**Dissociation Kinetics of [Re]-(F/Y)-βC19:α2**

The goal of observing and measuring radical transfer processes from Y356 of modified peptides such as [Re]-Y-βC19 requires that the peptide remain bound to α2 for a sufficiently long time so that radical injection can occur. In other terms, the dissociation rate constant for the [Re]-Y-βC19:α2 complex ($k_{off}$) must be slower than the PCET rate constant. To this end, we have undertaken experiments to measure the rates of association and dissociation of the peptide:α2 complex.

The data in Figure 3 shows that the emission of [Re]-Y-βC19 increases in the presence of α2 and thus loss of emission intensity could provide a reasonable method to measure the dissociation of the peptide from α2:[Re]-F-βC19 using stopped-flow spectroscopy. The baseline [Re] emission was determined by mixing Re-F-βC19 in one syringe with an equal
Discussion

Diffusion Limited Rate Constants

The upper limit for $k_{\text{off}}$ and $k_{\text{on}}$ may be ascertained from the determination of the diffusion coefficient for Re-Y-βC19 and α2 and an estimate of the binding site between the two using eq 6. Multiple-photon fluorescence recovery after photobleaching (MP-FRaP) was used to determine the diffusion coefficient of Re-Y-βC19 ($d_{\text{fit}}$). Figure S3 shows the recovery of the emission after an intense pulse of laser light was used to photo-bleach a defined area of the sample. The recovery is attributed to the diffusion of Re-Y-βC19 into the bleached area. A fit of recovery curve yields $d_{\text{fit}} = 6.1 \times 10^{-7}$ cm$^2$/s$^{-1}$ (35).

The diffusion coefficient of α2 ($d_{\alpha2}$) was determined using the Einstein-Stokes relationship in eq 4 using the radius found by DLS and that estimated from the crystal structure of α2. The average hydrodynamic radius obtained from DLS from a mass-weighted size distribution analysis was 55.8 Å. The radius of α2 was also estimated as 40.4 Å using eq 5 with $d_1$, $d_2$ and $d_3$ as 100.7 Å, 71.4 Å, and 73.2 Å, respectively. These two radii values yield $k_{\text{off}}$ for α2 as $3.9 \times 10^{-7}$ cm$^2$/s$^{-1}$ and $5.4 \times 10^{-7}$ cm$^2$/s$^{-1}$, respectively.

The active site of the α2:Re-Y/F-βC19 complex was estimated from the crystal structure of Ac-Y-βC19 bound to α2 which has only 16 amino acid residues visible by X-ray crystallography (18). Using eq. 5, the estimated radius of the active site was determined to be 8.97 Å. With eq. 6, the diffusion controlled rate constant for the [Re]-Y/F-βC19:α2 complex association ($k_{\text{on}}$) was found to be $6.8 - 7.8 \times 10^7$ M$^{-1}$/s$^{-1}$. Using $K_d = 9 \mu$M, a limiting rate constant for dissociation ($k_{\text{off}}$) was computed to be between 6100 s$^{-1}$ and 7000 s$^{-1}$. With these results and the SF kinetic data, the kinetics of association and dissociation for the [Re]-Y/F-βC19:α2 complex can limited to $7.8 \times 10^8$ M$^{-1}$/s$^{-1}$ > $k_{\text{on}}$ > $4.4 \times 10^7$ M$^{-1}$/s$^{-1}$ and 7000 s$^{-1}$ > $k_{\text{off}}$ > 400 s$^{-1}$.

Discussion

The [Re] complex within the [Re]-Y/F/3,5-F2Y-βC19 peptide serves as a probe of both peptide conformation and the pK$_a$ of the adjacent Y residue. Excitation of [Re] produces a metal-to-ligand charge transfer (MLCT) excited state, in which a metal $d$-orbital electron is promoted to an orbital of π* symmetry primarily localized on the bpy ligand resulting in an effective Re$^{1+}$[((bpy)$^*$)] electronic configuration. $^3[\text{Re}((\text{bpy})^*)]^*$ serves as a potent excited state oxidant ($E^0([\text{Re}^{1+}((\text{bpy})^*)]^*/[\text{Re}((\text{bpy})^*)]) = 1.59$ V vs. NHE (30)) that can oxidize deprotonated tyrosinate according to the mechanism:
The emission of the photoexcited [Re] center is therefore quenched by the transfer of an electron from the adjacent tyrosine in [Re]-Y-βC19 upon deprotonation of the tyrosine phenol (22). Thus monitoring of the emission band intensity as a function of pH probes the pK_a for the tyrosine phenol within the peptide and bound to α2. The solution pK_a value of tyrosine is maintained upon incorporation into the peptide and upon binding of the peptide to α2 (pK_a(3,5-F_2Y) = 7.2, pK_a([Re]-3,5-F_2Y-βC19) = 7.3, pK_a([Re]-3,5-F_2Y-βC19:α2) = 7.3). These results show that the photoRNR construct faithfully models RNR in that the pK_a of tyrosine at amino acid position 356 is unperturbed relative to the value in solution (42).

MLCT excitation of d^6 polypyridyl complexes is associated with a marked increase in the molecular dipole owing to the intramolecular charge separation. For this reason, the emission energy and intensity are strongly dependent upon polarity of the surrounding environment (43). High polarity solvents tend to lower the energy of the thermally equilibrated excited state, thus resulting in a red shift of the emission. Protic solvents, such as alcohols and water, also strongly quench the excited state lifetime through nonradiative decay mechanisms mediated by vibrational overlap of the excited state with the high frequency O–H stretch of the solvent (40). [Re] attached to the F_n Y-βC19 peptide therefore serves as an effective probe of peptide conformation and binding to class I RNRs. Removal of water from the solvation sphere of [Re] upon peptide conformational changes or binding events serves to both blue shift the energy of and increase the lifetime for the emission. The monotonic increase of the emission band to higher energy with a concomitant increase in intensity (Figure 3) is consistent with the exclusion of water from the solvation sphere of the [Re] probe upon binding of the peptide to the protein. The shift in the emission profile with addition of the protein therefore provides a direct measure of peptide binding to α2. The similarity of the K_D of [Re]-Y-βC19 relative to that measured for Ac-Y-βC19 by competitive inhibition suggests that modification of the peptide with the [Re] moiety does not greatly perturb the interaction of the peptide to the protein. The multiple emission lifetimes observed for [Re]-Y-βC19:α2 in the presence of RNR substrate and effector are consistent with multiple solvation environments of the [Re] chromophore, generally partitioning between protein-bound and solvent exposed. Several other peptides containing photooxidants of different sizes and shapes have been generated, all of which bind to α2 with K_D in the micromolar range (20, 21). In addition, the N-terminal peptide residues (356-360) are not located in the X-ray structure of Ac-Y-βC19:α2 (Figure 1; (18, 44)). Taken together these results are consistent with a flexible peptide N-terminus when bound to α2.

The observation of conformational flexibility within the photoRNR provides a connection to conceptualize the conformational gating of RNR activity at the molecular level. We have shown that the electron and proton are committed to a colinear pathway in α2 (21) for radical transport. The reaction rate for particle transfer is highly dependent on transfer distance because the electron and proton must tunnel. Whereas the electron can transfer over large distances, the proton cannot (9); hence PCET may provide a basis for conformational gating in RNR. The binding of α2 to β2 is known to trigger radical transport from •Y122 to Y356, but only in the presence of RNR substrate and effector (26, 27). Conformational flexibility of the 356-360 region of β2 within the α2:β2 complex would then allow the •Y356 radical to lock into the PCET pathway for radical transport across the subunit interface. Inasmuch as the [Re]-3,5-F_2Y-βC19: α2 complex has been shown to turnover upon irradiation (22), hole injection must be faster than the peptide off rate, which has now been determined to be in the range of 7000 s^{-1} > k_{off} > 400 s^{-1}. The conformational changes

\[
\text{[Re]}^I - \text{Y}^- \xrightarrow{\text{hv}} 3 \text{[Re]}^I - \text{Y}^- \xrightarrow{k_{ET}} \text{[Re]}^0 - \text{Y}^\bullet
\]
in the N-terminal peptide are likely to effect the efficient oxidation of Y731 in α2, making its detection challenging. Studies are now underway to examine how conformational changes of the N-terminal tail affect PCET in the photoRNR.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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**References**


Figure 1.
Molecular structure of [Re]-Y-βC19 bound to α2. Peptide and protein coordinates are adapted from the x-ray structure of α2 with Y-βC19 peptide bound (18). The Y731-Y730 and Y730-C439 distances are 3.3 and 3.4 Å, respectively. Y-βC19 residues 359-356 are not located in the structure, thus the Y356-Y731 distance is unknown.
Figure 2.
Emission spectra of [Re]-3,5-F2Y-βC19 as a function of pH, which was titrated from 4.5 to 11.5. Black curves correspond to emission at pH 4.5, 5.0 and 11.5. Inset: pH Titration curve of integrated emission intensity from pH 5.0 to 11.5.
Figure 3.
Emission spectra of [Re]-Y-βC19 (5 μM) at pH 7.5 as the α2 concentration is increased from 0 → 65 μM. Inset: Fraction of peptide bound as a function of α concentration (○) with fit (→) to binding model in eq 3.
Figure 4.
Emission spectra of [Re]-3,5-F₂Y-βC₁₉ (5 μM) and α₂ (20 μM) as the pH is titrated from 9.0 to 6.3. The solid black curves correspond to the endpoints of the titration at pH 9.0 and pH 6.3. The dark grey traces are spectra recorded at pH 6.3 (lower spectrum) and pH 9.0 (upper spectrum) in the absence of α₂. Inset: Integrated emission intensity as a function of pH (○) with fit (—) to monoprotic titration curve.
Figure 5.
Time resolved emission decay trace recorded at 600 nm for [Re]-Y-βC19 (5 μM) in the presence of α2 (20 μM) and ATP (3 mM).
Figure 6.
TA spectra recorded for 100 μM solutions of [Re]-F-βC19 (top) and [Re]-Y-βC19 (bottom) in the presence of 135 μM α2, 1 mM CDP, and 3 mM ATP in 20% glycerol at pH 7.5. Spectra are recorded at 65, 115, 215, and 415 ns (top) and at 115, 215, and 415 ns (bottom). Insets: Single wavelength absorbance kinetics data measured at 480 nm.