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Electrochemical and structural properties of a protein system designed to generate tyrosine Pourbaix diagrams

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Abstract

This report describes a model protein specifically tailored to electrochemically study the reduction potential of protein tyrosine radicals as a function of pH. The model system is based on the 67residue $\alpha_3 Y$ three-helix bundle. $\alpha_3 Y$ contains a single buried tyrosine at position 32 and displays structural properties inherent to a protein. The present report presents differential pulse voltammograms obtained from α_3 Y at both acidic (pH 5.4) and alkaline (pH 8.3) conditions. The observed Faradaic response is uniquely associated with Y32, as shown by site-directed mutagenesis. This is the first time voltammetry is successfully applied to detect a redox-active tyrosine residing in a structured protein environment. Tyrosine is a proton coupled electrontransfer cofactor making voltammetry-based pH titrations a central experimental approach. A second set of experiments was performed to demonstrate that pH-dependent studies can be conducted on the redox-active tyrosine without introducing large-scale structural changes in the protein scaffold. $\alpha_3 Y$ was re-engineered with the specific aim to place the imidazole group of a histidine close to the Y32 phenol ring. α_3 Y-K29H and α_3 Y-K36H each contain a histidine residue which protonation perturbs the fluorescence of Y32. We show that these variants are stable and well-folded proteins whose helical content, tertiary structure, solution aggregation state and solvent-sequestered position of Y32 remain pH insensitive across a range of at least 3-4 pH units. These results confirm that the local environment of Y32 can be altered and the resulting radical site studied by voltammetry over a broad pH range without interference from long-range structural effects.

SUPPORTING INFORMATION AVAILABLE

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Figures showing cyclic and differential pulse voltammograms of $\alpha_3 Y$ and control samples; the $\alpha_3 W$ solution NMR structure; absorption and fluorescence spectra of NAYA, $\alpha_3 Y$ and single-site histidine variants; absorption spectra of $\alpha_3 Y$ at neutral and high pH; 1D NMR spectra of $\alpha_3 Y$ -K36H at acidic, neutral and alkaline pH; denaturation plots of $\alpha_3 Y$ and single-site histidine variants; size-exclusion chromatograms of $\alpha_3 Y$ and $\alpha_3 Y$ -K29H; ¹⁵N-HSQC spectra of $\alpha_3 Y$ -K36H. This material is available free of charge via the internet at http://pubs.acs.org.

INTRODUCTION

Amino-acid radical enzymes use selected glycine, cysteine, tyrosine and/or tryptophan residues as "in-house" one-electron redox cofactors.^{1,2} Functional glycine and cysteine radicals tend to be generated near or at the active site and directly involved in the catalytic chemistry. The two aromatic residues are used in catalytic redox reactions as well as serving as intermediate components in radical-transfer chains spanning tens of ångströms. Well-known examples of the former type are the Y_Z tyrosine in photosystem II (PSII)^{3,4} and the cross-linked Tyr-His species in the active site of cytochrome *c* oxidase,⁵ while the triple tryptophan electron-transfer wire in *E. coli* DNA photolyase,⁶ the ~ 35 Å radical-transfer chain in *E. coli* ribonucleotide reductase (RNR),^{7,8} and the putative multi heme/tryptophan electron/radical transfer chain in the MauG/pre-methylamine dehydrogenase complex⁹ represent examples of the latter type.

Virtually nothing is known as to how the protein matrix influences the thermodynamic properties of biocatalytic amino-acid radicals or the free-energy profiles of radical-transfer chains. It is very challenging to measure the reduction potentials of these highly oxidizing redox cofactors and, consequently, only a few estimates are available in the literature. An apparent potential of 1.0 V (pH 7.6) was reported for the tyrosine Y-O[•]/Y-OH(122) redox pair in *E. coli* RNR,¹⁰ although poor equilibria between the redox mediators in the bulk solvent and the radical site make this value unreliable. The formal potential of YZ in the large, membrane-bound PSII enzyme cannot be obtained directly by standard techniques such as redox titration or voltammetry-based methods but a time-dependent "operating" potential has been estimated from kinetically derived equilibrium constants.^{3,11} Assuming an operating potential of 1.25 V for $P_{680}^+/P_{680}^{,11,12}$ the Y_Z -O $^{\bullet}/Y_Z$ -OH potential falls between 1.13 and 1.22 V depending on the time elapsed after light excitation and the redox state of the adjacent Mn₄Ca cluster (pH 6.5 range; 13 and references therein). Likewise, an operating potential of 0.83 V (pH 6.0) has been estimated for the PSII Y_D-O[•]/Y_D-OH redox couple.¹¹ These indirectly derived, single-pH potentials are, to our knowledge, the only available estimates for unmodified tyrosine radical cofactors. Even less is known about the potentials of protein glycine, cysteine and tryptophan radical cofactors with experimental data available only for the two-electron heme/W191 redox reactions in cytochrome cperoxidase¹⁴ and the photolyase tryptophan-radical system.¹⁵ Thus, on the basis of existing experimental data it is not possible to make any firm predictions regarding how the protein environment influences the reduction potentials of amino-acid radical cofactors. The hydrogen-bonding status of the reduced and radical states, the chemical characteristics of hydrogen-bonding partners, electrostatic interactions, solvent accessibility and the effective dielectric of the protein environment are likely to play important roles^{3,16–19} but, at present, correlations between these parameters and amino-acid radical reduction potentials are unknown and unexplored.

The thermodynamic properties of aqueous tyrosine give rise to a few key predictions regarding tyrosine redox chemistry in proteins.^{2,3} The following half reactions, redox couples and acid dissociation constants are associated with tyrosine in water:

$\text{Y-OH}^{\bullet+} + e^- \Longleftrightarrow \text{Y-OH}$	Y-OH• ⁺ /Y-OH
$\text{Y-O}^{\bullet} + e^- + \text{H}^+ \Longleftrightarrow \text{Y-OH}$	Y-O•/Y-OH
$Y\text{-}O^{\bullet} + e^- \Longleftrightarrow Y\text{-}O^-$	Y-O•/Y-O ⁻
$\text{Y-OH}\bullet^+ \Longleftrightarrow \text{Y-O}\bullet + \text{H}^+$	$K_{\mathrm{oY}} = [\mathrm{Y}\text{-}\mathrm{O}^{\bullet}][\mathrm{H}^{+}]/[\mathrm{Y}\text{-}\mathrm{OH}^{\bullet+}]$
$\text{Y-OH} \Longleftrightarrow \text{Y-O}^- + \text{H}^+$	$K_{\rm rY} = [Y-O^-][H^+]/[Y-OH]$

The cation Y-OH^{•+}/Y-OH redox couple dominates at pH below pK_{oY} while the tyrosinate Y-O[•]/Y-O⁻ redox couple is observed at pH above pK_{rY} . There is no proton release or uptake associated with the change in redox state and thus the reduction potentials of the Y-OH^{•+}/Y-OH and Y-O[•]/Y-O⁻ redox pairs are pH independent. The neutral tyrosine Y-O[•]/Y-OH pair is the dominant redox couple in the $pK_{oY} < pH < pK_{rY}$ region. Since tyrosine oxidation/ reduction is a $1e^{-}/1H^+$ event in this pH region, the potential follows the Nernst equation and decreases by 59 mV per pH unit at 25° C. The pK_{oY} and pK_{rY} values are about -2 and 10 for aqueous tyrosine, respectively.²⁰ These values suggest that Y-O[•]/Y-OH is the dominant redox couple in a protein environment. Consequently, tyrosine oxidation/reduction reactions occurring inside proteins are expected to a large extent to involve proton-coupled electron transfer (PCET).

There are no electrochemical data of any kind available for a protein tyrosine radical, much less a full Pourbaix diagram representing the formal potential of this species over a significant pH range. The PCET characteristics of tyrosine and phenol redox systems predict that local interactions are critical for tuning their redox properties.^{3,8,21} Thus, in order to obtain a meaningful Pourbaix diagram it is essential that the measured potential is not strongly influenced by global changes occurring in the protein scaffold as a function of pH but rather reflects local conditions at the radical site. We have developed a tyrosine radical protein system that fulfills these criteria. The complexity of this task required a fairly extensive experimental approach including the use of voltammetry methods, protein design and engineering, absorption, fluorescence and CD spectroscopy as well as several NMR-based techniques. The obtained data are described and analyzed in two connected papers. In the current paper we describe the electrochemical and structural properties of a protein system designed to provide tyrosine Pourbaix diagrams according to the criteria listed above. In a follow-up study, tyrosine Pourbaix diagrams obtained from this protein system are described and discussed.²²

The tyrosine radical protein system is based on the *de novo* designed, 67-residue $\alpha_3 Y$ threehelix bundle, which contains a single buried tyrosine at position 32 and originally no histidine residues.¹⁶ In the present report we show that high-quality voltammetry data can be obtained from $\alpha_3 Y$ at both acidic and alkaline conditions. We show that the observed Faradaic response is uniquely associated with Y32. This is the first time voltammetry is successfully applied to detect a redox-active tyrosine residing in a structured protein environment. We also demonstrate a high-level structural control over $\alpha_3 Y$ by changing the environment of its redox-active tyrosine and proving that the resulting variants exhibit the required structural characteristics to support voltammetry-based pH titrations in a stable

protein background. This is significant since pH-based studies are central to characterizing proton-coupled ET reactions in proteins. More specifically, $\alpha_3 Y$ was re-engineered to make a histidine variant in which the imidazole ring of the introduced histidine interacts structurally with the phenol group of Y32. Our aim was to introduce an interaction at the site of the redox-active tyrosine and then specifically determine the effect of this interaction on the potential of Y32. This would demonstrate that the constructed tyrosine radical model system is capable of providing relevant radical site-specific information. Two variants of interest, a₃Y-K29H and a₃Y-K36H, were identified using protein modeling combined with optical and NMR-based spectroscopic screening. These two stable and well-structured proteins do not display any major changes in their secondary structures, tertiary structures or solution aggregation state across a range of 3-4 pH units. Their single tyrosine residue Y32 is located in a hydrophobic and structured environment. Moreover, protonation of the introduced histidines perturbs the fluorescence of Y32 suggesting that α_3 Y-K29H and α_3 Y-K36H contain an electrostatically coupled Y32/His pair. We conclude that α_3 Y-K29H and a₃Y-K36H exhibit the appropriate characteristics for probing the potential of Y32 as a function of local interactions and the solution pH.

MATERIALS AND METHODS

Protein expression and purification

The α_3 Y proteins were made by QuikChange (Stratagene) using a modified α_3 W/pET32b (Novagen) vector as template.¹⁸ α_3 Y has the following amino-acid sequence: GSRVKALEEKVKALEGKVKALGGGGRIEELKKKYEELKKKIEELGGGGEVKKVEEE VKKLEEEIKKL in which the N-terminal GS residues form part of a thrombin cleavage site. These two residues are labeled as -2 and -1 to keep the amino-acid numbering consistent with the chemically synthesized 65-residue $\alpha_3 Y$ protein.¹⁶ $\alpha_3 Y$ and variants were expressed in LB or minimal media and purified following standard protocols for His-tagged proteins (Novagen). Transformed BL21(DE3)pLysS or BL21-CodonPlus(DE3)-RIL cells were harvested either after a 3-4 hour IPTG induction period at 37° C (LB cultures) or after an overnight induction at 30° C (minimal media cultures) and stored at -20° C. Cells were resuspended in 20 mM Tris-HCl, 500 mM NaCl, 5 mM imidazole, pH 7.9 and lysed by sonication. The lysate was clarified by centrifugation, passed over a His•bind (Novagen) nickel column, and the thioredoxin- $\alpha_3 Y$ fusion protein eluted by an imidazole gradient. Thrombin (T6634; Sigma) was added to the thioredoxin- α_3 Y protein fraction and the resulting mixture dialyzed against 50 mM Tris-HCl, 500 mM NaCl, 2.5 mM CaCl₂, pH 8.0 at room temperature overnight. The digestion mixture was passed over a second nickel column to remove the His-tagged thioredoxin and any remaining undigested fusion protein. Target proteins were isolated by reverse-phase HPLC (semi-preparative TP2181010 column; Grace/Vydac) using an acetonitrile/water gradient containing 0.1% (w/v) trifluoroacetic acid and stored as lyophilized powder.

Absorption and fluorescence spectroscopy

Absorption spectra were collected on a Varian Cary 50 Bio or a Hitachi U-3000 UV/Vis spectrometer at room temperature. pH-titration samples were prepared by dissolving lyophilized protein in a 10 mM potassium phosphate, 10 mM HEPES, 10 mM borate, 10

mM CAPS, pH 7.0 buffer to an Abs₂₇₆ of 0.2 (10 mm path). The solution was split in two equal fractions and the pH carefully adjusted with 12 M HCl or 10 M NaOH. pH titrations were performed by constant volume titration. The apparent tyrosinate/tyrosine pK_a of Y32 was estimated by measuring at 293 nm (Abs_{max} for deprotonated Y32) and 400 nm (baseline) as a function of pH and fitting the resulting pH-titration curve to a single pK_a using the nonlinear curve fitting routines in KaleidaGraph (www.synergy.com).

Fluorescence spectra were collected on a Horiba Jobin Yvon Spex Fluorolog spectroflurometer at 23° C. pH-titration samples were prepared by dissolving lyophilized protein in a 10 mM sodium acetate, 10 mM potassium phosphate, 10 mm sodium borate (APB) pH 7.0 buffer to an Abs₂₇₆ of 0.2 (10 mm path), dividing the sample in two equal fractions, and adjusting the pH with either concentrated phosphoric acid or 10 M NaOH. The experiments were performed by constant volume titration and using an $\lambda_{ex} = 276$ nm and $\lambda_{em} = 285-445$ nm. The slit width for the excitation and emission light was 0.7 and 2.0 nm, respectively, and the averaging time for each 0.05 nm step was 0.2 seconds. Tyrosine emission center of mass was calculated as described in Ref. 23 and the resulting pH-titration curves fitted to a single p K_a .

Circular dichroism spectroscopy

CD studies were conducted on an Aviv 202 CD spectrometer at 25° C. For the α -helical measurements, lyophilized protein was dissolved in 10 mM APB pH 8.2 buffer to a concentration of ~ 50 μ M. To ensure accurate absorbance readings, the cuvette pathlength was 10 and 2 mm for the UV/Vis and CD measurements, respectively. The absolute degree of secondary structures was determined by using α_3 W as a reference (76 ± 1% α -helical between pH 4 and 10).^{16,18} pH titrations were conducted by constant volume titration and the samples prepared by dissolving protein powder in 10 mM APB pH 7.0 buffer, splitting the solution into two equal fractions, and adjusting the pH with either 12 M HCl or 10 M NaOH. Chemical denaturation was conducted by automated constant volume titration of a 10 M urea protein sample into a 0 M urea protein sample. Samples were prepared by diluting protein concentrations were 15–40 μ M for the pH and urea measurements. The degree of α -helical content and changes in this parameter were monitored by measuring the mean residue ellipticity at 222 nm ([Θ]₂₂₂). The chemical denaturation curves were fitted as described in Ref. 24.

Size-exclusion chromatography

Gel filtration was performed using an analytical SuperdexTM 75 column (GE Healthcare) equilibrated with 10 mM APB pH 7.0 buffer containing 100 mM KCl. The protein loading concentration was 250 μ M.

NMR spectroscopy

NMR data were collected using a Varian Inova 750 MHz spectrometer equipped with a conventional room temperature probe or a Bruker Advance III 750 MHz spectrometer fitted with a cryoprobe. pH-titration samples were prepared by dissolving lyophilized protein in a 10 mM deuterated sodium acetate, 10 mM sodium phosphate, 30 mM KCl, pH 4.0 buffer

and in a 10 mM sodium phosphate, 10 mM sodium borate, 30 mM KCl, pH 10.0 buffer. Buffers were prepared in D₂O, all samples were ~ $600 \,\mu\text{M}$ in protein and contained $500 \,\mu\text{M}$ 2,2-dimethyl-2-silapentane-5-sulfonate sodium salt (DSS) as a chemical-shift standard. pH titrations were conducted by equal volume titration and collecting one-dimensional (1D) spectra of the a₃Y-His proteins at 25° C. The chemical shift of the imidazole ring Hɛ1 resonance²⁵ was used to determine the apparent imidazole/imidazolium pK_a values of the single histidine in the α_3 Y-His proteins. Isotope effects were corrected using the following relationship: $pK(H_2O) = 0.929 \times pK(D_2O) + 0.42$.²⁶ Two-dimensional (2D) ¹⁵N-HSQC spectra²⁷ were obtained on ~ 250 μ M ¹⁵N-labeled α_3 Y-K29H and α_3 Y-K36H dissolved in either 20 mM deuterated sodium acetate, 20 mM sodium phosphate or 20 mM deuterated Tris. All samples contained 20 mM NaCl and 8% D₂O. ¹⁵N-HSQC spectra were acquired at 25° and 35° C using a spectral width of 14.0 ppm and 1024 complex points in the direct ¹H dimension and 18.0 ppm and 64 complex points in the indirect ¹⁵N dimension. 2D ¹H–¹H NOESY spectra²⁸ were obtained on ~ 200 μ M (α_3 Y-K29H and α_3 Y-K36H dissolved in D₂O containing 20 mM deuterated sodium acetate and 20 mM NaCl or 20 mM deuterated Tris and 20 mM NaCl. NOESY spectra were collected at 25° using a mixing time of 150 ms, a spectral width of 10.0 ppm and 2048 complex points in the direct ¹H dimension and 10.0 ppm and 800 complex points in the indirect ¹H dimension. NMR data processing was performed with the Felix95 software (Accelrys Inc., San Diego, CA).

Electrochemistry

Cyclic voltammetry²⁹ and differential pulse voltammetry^{29,30} were performed using an Autolab PGSTAT12 potentiostat. The electrochemical workstation was equipped with a temperature-controlled, Faraday-cage protected three-electrode micro-cell (Princeton Applied Research) containing a Ag/AgCl reference electrode, a platinum wire counter electrode, and 3 mm glassy carbon working electrode (all purchased from Advanced Measurements Inc.). The reference and counter electrodes were stored dry and routinely replaced after a limited set of experiments. They were prepared by filling the reference electrode with a 3 M KCl/saturated AgCl solution and the counter electrode with the buffer solution in which the sample was prepared. The surface of the working electrode was carefully polished between each measurement using a 0.05 µm alumina/water slurry on a glass-plate mounted microcloth pad (Bioanalytical systems Inc.). The electrode was manually polished for 60 sec., rinsed with water, sonicated first in ethanol for about 60 sec. and then in milli-Q water for another 60 sec., and finally rinsed with an excess of milli-Q water directed against the surface of the electrode. This protocol was repeated 2-3 times until a reproducible Faradaic response was observed from the sample. This cyclic procedure was typically performed only at the beginning of the experimental day and once the working electrode was conditioned, only a single polish/sonicate/rinse step was required to activate the surface between measurements. Voltammetry measurements were performed immediately following the electrode-activation treatment. The performance of the assembled three-electrode cell was routinely checked by collecting differential pulse voltammograms from a standard sample containing 300 µM N-acetyl-tryptophanamide in 10 mM APB, 125 mM KCl, pH 3.0. All samples were prepared using ultra-pure chemicals and the measurements made at 23° C under an argon atmosphere. Data analysis was performed using the Autolab GPES software and PeakFit (Systat Software Inc.).

RESULTS

Voltammetry studies of a₃Y

Panels A and B in Fig. 1 show cyclic voltammograms of α_3 Y and N-acetyl-L-tyrosinamide (NAYA), respectively. The two traces were obtained at very similar conditions, which are detailed in the figure legend. There is no observable Faradaic response from the protein sample while the NAYA sample gives rise to an easily detectable anodic waveform. As expected,^{31,32} the NAYA cyclic voltammogram is essentially completely irreversible. The poor response observed from α_3 Y when using cyclic voltammetry could not be improved by changing the protein concentration, buffer conditions or acquisition settings (e.g. see Fig. S1A in the supporting information). Thus unlike aqueous tyrosine³¹ and aqueous phenol,³³ electrochemical analysis using cyclic voltammetry is at the present not a feasible approach for the α_3 Y system.

However, a significant Faradaic response could be obtained from $\alpha_3 Y$ when using the more sensitive method of differential pulse voltammetry (DPV).^{29,30} Panels D and E in Fig. 1 show DP voltammograms obtained from $\alpha_3 Y$ (blue) and $\alpha_3 C$ (orange) at nearly identical conditions (see figure legend). $\alpha_3 C$ is a redox-inert Y32C variant of $\alpha_3 Y$.¹⁹ At both acidic (panel D) and alkaline (panel E) conditions $\alpha_3 Y$ gives rise to a distinct oxidation wave while $\alpha_3 C$ does not. Differential pulse voltammograms recorded on samples containing $\alpha_3 C$ look essentially identical to baseline voltammograms collected on buffer samples (e.g. Fig. S1B). We conclude that there is no Faradaic response from the three-helix bundle scaffold when Y32 is removed. Thus, the $\alpha_3 Y$ voltammograms displayed in panels D and E represent the unique tyrosine residing in the hydrophobic core of $\alpha_3 Y$.

Voltammograms obtained from aqueous samples at highly oxidizing potentials contain a prominent background current arising from water oxidation occurring at the surface of the working electrode. The magnitude of this current is influenced by the sample pH and by the type of working electrode used. Solvent oxidation effects are evident when comparing the raw and baseline-corrected α_3 Y pH 8.3 voltammogram displayed in panels E and F, respectively. α_3 Y DP voltammograms obtained at both acidic and alkaline pH display near symmetric waveforms with a well-defined peak potential and a width at half height of ~ 125 mV.

Panels C and F in Fig. 1 display baseline-corrected DPV traces of NAYA and α_3 Y, respectively. These traces were obtained at comparable sample conditions and using identical acquisition parameters (see figure legend). The only significant difference was the KCl concentration, which was 40 mM in the protein sample and 200 mM in the NAYA sample. At lower salt concentration the NAYA voltammogram displayed characteristics consistent with electrode absorption. α_3 Y gives rise to a current in the 0.1 µAmp range when using DPV while the NAYA sample provide a current on the 1.0 µAmp range. Thus, the peak current of the Y32 voltammogram is about one order of magnitude smaller relative to the peak current obtained from freely solvated NAYA.

We conclude that high-quality tyrosine DP voltammograms can be obtained from $\alpha_3 Y$ at both acidic and alkaline pH. The voltammogram collected at pH 5.6 is particularly

impressive considering the high potential involved. Overall, these results represent the first successful use of voltammetry to detect a tyrosine radical located in a stable and well-folded protein environment. They also suggest that the α_3 Y system can be used to obtain tyrosine Pourbaix diagrams spanning a significant pH range.

Protein modeling

 $\alpha_3 Y$ was re-engineered to make a histidine variant displaying a detectable interaction between the imidazole ring of the introduced histidine and the phenol head group of Y32. $\alpha_3 Y$ is a W32Y variant of the structurally characterized $\alpha_3 W$ three-helix bundle (see Fig. S2 in the supporting information).¹⁸ α_3 Y and α_3 W exhibit very similar structural properties including their a-helical content (~ 75%), pH stability (5-6 pH units), global stability (~ 4-5 kcal mol⁻¹) and solution aggregation state (monomeric).^{16,18} Moreover, 2D ¹³C-HSQC spectra reflecting the environment of protein core residues in these two proteins display comparable spectral linewidths and chemical-shift dispersion. These observations show that large-scale structural changes do not occur in the three-helix bundle scaffold upon changing tryptophan 32 to a tyrosine. On this basis, $\alpha_3 Y$ models with various Y32 χ^1 dihedral angles were made from the a_3 W NMR structure¹⁸ to identify sites where the incorporation of a histidine could place the imidazole ring within 5 Å of the Y32 phenol oxygen (Fig. 2). Histidine was modeled into each identified site and a broad range of Y32/His χ^1 rotamer combinations were visually inspected. This analysis promoted the generation of the following eight proteins: a₃Y-V9H, a₃Y-L12H, a₃Y-E13H, a₃Y-K29H, a₃Y-E33H, a₃Y-K36H, α_3 Y-L58H and α_3 Y-I62H.

Screening for Y32/His interactions in the a₃Y-His variants

It is long known that the optical properties of phenols are sensitive to the dielectric and hydrogen-bonding properties of solvating molecules.^{e.g.34} To provide an example, Fig. S3A in the supporting information displays the absorption spectra of $\alpha_3 Y$ (λ_{max} 277.8 nm) and NAYA (λ_{max} 275.3 nm) both obtained at neutral pH. The relative redshift of the $\alpha_3 Y$ spectrum indicates that the Y32 side chain is shielded from the bulk solvent. This is consistent with other characteristics of Y32¹⁶ and the sequestered position of the W32 residue (solvent accessible surface area of $2.6 \pm 1.4\%$ across the $\alpha_3 W$ NMR structural ensemble).^{18,35} Absorption and fluorescence spectra of the eight α_3 Y-His variants were compared to $\alpha_3 Y$ spectra obtained at corresponding conditions to search for changes in the microenvironment of Y32. Only minor shifts were detected in the Y32 absorption (Fig. S3B) while the fluorescence data provided more distinguishing information. We found α_3 Y-E13H, a₃Y-E33H and a₃Y-E58H excitation (data not shown) and emission (Fig. S4) spectra essentially identical to those of $\alpha_3 Y$ and these three proteins were excluded for further characterization. In contrast, excitation and/or emission spectra of α_3 Y-V9H, α_3 Y-L12H, α_3 Y-K29H, α_3 Y-K36H and α_3 Y-I62H are shifted relative to corresponding α_3 Y spectra (Fig. S5), which prompted further studies of these five proteins.

Alterations in the optical properties of Y32 may arise from a local change in the vicinity of the aromatic residue but may also reflect a global perturbation of the protein scaffold. A second screening step was performed to probe the structural integrity of the five selected α_3 Y-His proteins. p K_{app} values of Y32 and introduced histidines were determined optically

via the tyrosinate absorbance at 293 nm (Fig. S6) and by following spectroscopically the NMR chemical shift of the imidazole $\varepsilon 1$ proton (Fig. S7) as a function of pH. CD spectroscopy was used to determine absolute α -helical content, the pH-sensitivity in this parameter, and global protein stability. Only a minor decrease in the degree of helicity was observed for α_3 Y-V9H, α_3 Y-K29H and α_3 Y-K36H relative to α_3 Y (~ 5–10%; Fig. 3 and Table 1). These proteins also give rise to well-defined unfolding/folding transitions (Fig. S8A) from which the global protein stabilities could be determined (Table 1). In contrast, α_3 Y-L12H and α_3 Y-I62H show a loss of ~ 20% of their α -helical contents relative to α_3 Y (Table 1) and give rise to poorly defined denaturation curves (Fig. S8B). In addition, NMR resonances representing the H12 and H62 ring protons display relatively broad linewidths. This effect was particularly pronounced for α_3 Y-L12H with the imidazole H ϵ_1 resonance broadened beyond detection between pH 6.3 and 7.4 (Fig. 3C). These observations are all consistent with structurally perturbed protein folds and no additional data were collected on α_3 Y-L12H and α_3 Y-I62H.

A key requirement for the tyrosine radical model system is that pH-induced structural changes in the protein scaffold do not obscure detection and characterization of local interactions at the radical site. Thus for the histidine variants we wish to measure the potential of Y32 across the titratable pH range of the introduced histidine while avoiding large-scale structural changes. α_3 Y-K29H and α_3 Y-K36H meet this requirement well as they remain 66.8 ± 2.5 and 64.7 ± 0.8% α -helical, respectively, ± 2 pH units around the p K_{app} values of H29 and H36 (Table 1). Although the CD and NMR data collected on α_3 Y-V9H suggest a stable and well-folded protein, the p K_{app} of H9 is below the protein pH-stability range (Table 1). For this reason, α_3 Y-V9H was excluded from further characterization.

Spectroscopic evidence of Y32/His interactions in a_3 Y-K29H and a_3 Y-K36H

Fluorescence pH titrations were conducted to probe for more direct evidence of Y32/His interactions in α_3 Y-K29H and α_3 Y-K36H. Figure 4A shows the α_3 Y tyrosine emission center of mass as a function of pH. A 1.0 nm shift was observed between pH 5.1 and 9.1 and yielded a fitted single pK_{app} of 8.1 ± 0.1. This effect may arise from a pH-induced change in the electrostatic environment of the tyrosine, the hydrogen-bonding properties of the phenol OH group, or a combination thereof.³⁶ A more substantial shift of 4.3 nm was observed for α_3 Y-K36H (Fig. 4C). The pH-induced spectral changes titrate with a p K_{app} of 7.1 ± 0.1, which is equivalent to the pK_{app} value determined for H36 via NMR (Table 1). The correlation between the two pK_{app} values suggests that the Y32 fluorescence is sensitive to the protonation state of H36 and that the two aromatic side chains are in close proximity. Likewise, the α_3 Y-K29H emission center of mass titrates with a p K_{app} of 7.4 ± 0.1 (Fig. 4B). This value is similar to the NMR-derived H29 p K_{app} of 7.1 ± 0.1 suggesting a structural connection between Y32 and H29. The observed changes in the Y32 fluorescence most likely reflect electrostatic interactions between Y32 and the introduced histidine residues. Alternatively, changing the protonation state of the histidine leads to a change in the hydrogen-bonding environment of the Y32 OH group. Either of these explanations suggests that the imidazole group of the histidine residue introduced in α_3 Y-K29H and α_3 Y-K36H reside near or at the Y32 site.

Confirming the absence of pH-induced changes in aggregation state and tertiary structures

In preparation for voltammetry-based pH studies, it was essential to establish that α_3 Y-K29H and α_3 Y-K36H are monomeric and well structured across a significant pH range. Earlier work involving sedimentation equilibrium ultracentrifugation and analytical sizeexclusion chromatography have demonstrated that the aggregation state of $\alpha_3 Y$ in solution is that of a monomeric protein over a concentration range of at least 4-850 µM.¹⁶ Moreover, 1D and 2D NMR spectra collected on α_3 Y in the mM concentration range display characteristics consistent with a non-aggregated, uniquely folded protein. On the basis of these earlier observations, $\alpha_3 Y$ was used here as a standard to determine the aggregation states of a₃Y-K29H and a₃Y-K36H. Fig. S9 in the supporting information shows sizeexclusion chromatograms obtained from $\alpha_3 Y$ and $\alpha_3 Y$ -K29H at pH 7.0 and a protein loading concentration of 250 μ M. The α_3 Y and α_3 Y-K29H chromatograms display profiles consistent with a single major species (95% and 91% of the total 220 nm absorbance, respectively) eluting with a retention volume of 13.1 and 12.9 ml, respectively. An equivalent chromatogram obtained on a₃Y-K36H shows a single major species eluting at 13.1 ml (data not shown). We conclude that there are no significant differences in the hydrodynamic volume of the three proteins and, consequently, that they are all monomeric at neutral pH.

2D NMR spectroscopy was used to probe for changes in aggregation state and/or tertiary structure as a function of pH. ¹⁵N-HSQC spectra were collected from a₃Y-K29H and a₃Y-K36H at three different pH values (5.5, 7.0 and 8.5) and two different temperatures (25° and 35° C). Fig. 5B shows the α_3 Y-K29H ¹⁵N-HSQC spectrum obtained at pH 7.0 and 25° C. The displayed region of the 2D spectrum contains a set of 54 resolved ${}^{15}N$ - ${}^{1}H$ cross peaks. In this spectral region we expect to detect 56 cross peaks arising from backbone N-H groups in the helical regions of the protein. Amide resonances associated with the eight glycines in loop regions (¹⁵N 107–109 ppm) and L65 at the C-terminus (¹⁵N 126.6 ppm) occur outside the displayed region. The observed narrow spectral linewidths, overall chemical-shift dispersion and absence of minor peaks confirm that a₃Y-K29H is a monomeric, uniquely folded protein at these conditions. Moreover, when comparing panels A, B and C in Fig. 5 it is clear that there are no significant changes in the spectral linewidths as a function of pH, which, in turn, means that there is no significant change in the hydrodynamic volume of a₃Y-K29H. pH-induced electrostatic and hydrogen-exchange effects influence peak positions and intensities as expected but the overall chemical-shift dispersion remains essentially the same. Thus, α_3 Y-K29H remains well folded across the pH 5.5 to 8.5 range. The same conclusions can be made from the ¹⁵N-HSQC pH 5.5, 7.0 and 8.5 spectra obtained from a₃Y-K29H at 35° C (data not shown) and from a₃3Y-K36H at 25° C (data not shown) and 35° C (Fig. S10). There are no significant changes in spectral linewidths and chemical-shift dispersion as a function of protein, pH and temperature. We conclude that a₃Y-K29H and a₃Y-K36H are monomeric and well-structured proteins across a pH range of at least 5.5 to 8.5.

Confirming that Y32 is buried in a_3 Y-K29H and a_3 Y-K36H

Figure 6 shows ¹H-¹H NOESY spectra obtained from α_3 Y-K29H at pH 8.5 and 5.6 (panels A and B) and α_3 Y-K36H at pH 8.4 and 5.6 (panels C and D). The selected region of the NOESY spectrum display NOEs between the Y32 ring-protons and aliphatic protons located within a distance of ~ 5 Å. The $\delta 1 \& \delta 2$ (the *meta* positions of the aromatic ring) and $\epsilon 1 \& \epsilon 2$ (*ortho* positions) ring-protons are unambiguously assigned on the basis of the observed intraresidue NOE patterns. For example, consider the α_3 Y-K36H pH 8.4 spectrum (Fig. 6C). The unresolved resonances of the $\delta 1 \& \delta 2$ ring-protons at 6.89 ppm show strong NOE correlations to the Y32 β -protons at 2.99 and 3.15 ppm and Y32 α -proton at 3.98 ppm. The unresolved resonances of the $\epsilon 1 \& \epsilon 2$ ring-protons at 6.76 also show intra-residue NOEs to the Y32 β - and α -protons but with weaker intensities corresponding to the longer ¹H-¹H distances. The same intra-residue Y32 NOE correlation patterns are observed in panels A, B and D.

Figure 6 shows that the Y32 ring-protons are close to a dozen or more protons with resonances in the 0.59 to 0.97 ppm spectral region at both pH 8.5 and 5.6. The observed chemical shifts are consistent with resonances from methyl groups associated with valine, leucine and isoleucine residues. These types of residues form the hydrophobic packing layers above, below and at the level of W32 in the α_3 W structure.^{18,37} Depending on the pH, the α_3 Y-K29H aromatic ring-protons exhibit at least 12–26 additional inter-residue NOEs to aliphatic protons with resonances in the 1.40 to 4.17 ppm region (Figs. 6A and B). For α_3 Y-K36H at least 19–28 additional inter-residue NOEs are observed between the aromatic ring-protons and nearby aliphatic protons (Figs. 6C and D). These observations firmly suggest that Y32 is buried in the hydrophobic core of the two α_3 Y histidine variants, as expected on the basis of the protein design and the structural knowledge of α_3 W.

DISCUSSION

Key structural properties of the tyrosine radical protein system

Despite the fact that tyrosine radicals have been known to be involved in biological redox processes for more that three decades,³⁸ a basic Pourbaix diagram describing the reduction potential of a protein tyrosine radical as a function of the solution pH is not available. In fact, information regarding reduction potentials associated with protein tyrosine radicals is limited to only two indirectly derived, single-pH estimates (*vide supra*). This situation suggests that studies limited to the natural systems will not provide basic thermodynamic information of these essential PCET cofactors in a timely manner, or even at all. For this reason, we adopted a model protein approach.

It is vital to stress that although the 7.5 kDa three-helix bundle scaffold used here is small relative to most naturally occurring tyrosine radical proteins, it nonetheless exhibits all of the key characteristics of a protein system. Its folding is driven by the hydrophobic effect.^{18,35,37} Its unfolding/folding transition is reversible and cooperative (e.g. Fig. S8A). Importantly, the ensemble of conformers that represent the solution state of the three-helix bundle proteins occupy narrow wells as shown by the excellent structural statistics of $\alpha_3 W$ (see Fig. S2 legend) and by the characteristics of their HSQC spectra (e.g. Figs. 5 and Fig.

S10). Structural statistics of similar high quality were recently obtained for the NMR solution structures of two phenol-containing derivatives of α_3 Y (unpublished data). Finally, the tyrosine of interest Y32 is desolvated and maintained in a highly structured environment across a broad pH range. These structural properties uniquely separate the model system described here from all other tyrosine/phenol radical model systems described thus far in the literature. In these small-molecule^{e.g.39} and peptide^{e.g.32,40} model systems, the phenol side chain resides in a highly solvent-exposed and dynamic environment.

Electrochemical approach

Protein electrochemistry include classic redox titrations using chemical titrants and a redox cuvette,⁴¹ redox titrations using a thin-layer spectroelectrochemical cell controlled by a potentiostat,⁴² and protein voltammetry.⁴³ Redox titrations require a mediator system that covers the potential range of interest and a distinct spectroscopic feature that reflects the redox state of the cofactor. The size and the complexity of the redox protein generally do not provide an experimental barrier as long as the redox center is in equilibrium with the electrode via the redox mediators and give rise to a detectable signal that occurs outside the spectral envelope of the added mediators. In contrast, protein voltammetry does not require redox mediators or a spectroscopic probe of the redox system. This technique is however highly sensitive to the size of the protein and its orientation on the electrode surface since there must be electronic contact between the redox cofactor and the working electrode. The experimental time scale differs between the two approaches as redox titrations are conducted on the hours time scale while voltammetry data are typically collected on the minutes to ms time scale.

The main characteristics of $\alpha_3 Y$ all suggest that voltammetry is the feasible approach. The small size of the protein is an advantage and predicts that a functional Y32/electrode electron-tunneling distance can occur in multiple $\alpha_3 Y$ /electrode spatial orientations. $\alpha_3 Y$ is a high-potential system and an experimental time scale in the minutes rather than hours range reduces the risk of oxidative damage. Moreover, the weak optical features of $\alpha_3 Y$ (ϵ_{276} 1490 M^{-1} cm⁻¹ for reduced $\alpha_3 Y$, ϵ_{408} 2750 M^{-1} cm⁻¹ for oxidized tyrosine⁴⁴) combined with spectral overlap of redox mediators typically used in the high-potential range,^{10,45} will make spectroscopic monitoring of the $\alpha_3 Y$ redox state challenging. Finally, and most importantly, voltammetry can provide kinetic and mechanistic information in addition to thermodynamic parameters.^{33,43} Thus, the key demonstration of a reproducible Faradaic response from Y32 sets the stage for more detailed studies as we continue to develop the $\alpha_3 Y$ system.

The main challenge in protein voltammetry is to identify a working electrode system that provides the potential range required, allows direct cofactor/electrode electron transfer and promotes the folded protein to interact in a favorably manner with the electrode surface without adsorptive denaturation. A variety of electrode preparation strategies have been employed in order to modulate the strength of the protein/electrode interactions to, ideally, generate either a diffusion-controlled system or a stable protein film.⁴³

In this study, our first aim was to demonstrate that Y32 is redox active and that voltammetry can be used to probe the potential of the tyrosine in the folded protein at both acidic and alkaline conditions. A glassy carbon electrode was chosen as the working electrode system since it has an anodic potential window that extends to about +1.3 V vs. NHE in aqueous media.²⁹ The glassy carbon electrode was activated by polishing the surface in an $Al_2O_3/$ water slurry, which is predicted to generate a hydrophilic surface with C-O functionalities such as hydroxyls, carbonyls, ethers and carboxylates.⁴⁶ In a preliminary study on $\alpha_3 Y$, a Faradaic response could not be obtained from the folded protein but only at sample conditions where the protein is unfolded.¹⁶ By refining the pretreatment of the working electrode (see Materials and Methods section), we have now obtained differential pulse voltammograms from $\alpha_3 Y$ at sample conditions where the protein is known to be monomeric, folded and well structured. Once the electrode pretreatment protocol was obtained, the electrochemical response from $\alpha_3 Y$ was compared at high and low pH allowing sample conditions and acquisition parameters to be refined in an iterative manner yielding typical voltammograms of the quality shown in Figs. 1D to F. We showed that the Faradaic response is uniquely associated with Y32 and that it is abolished when the tyrosine is replaced by site-directed mutagenesis. Importantly, this is the first time voltammetry is successfully applied to detect a redox-active tyrosine residing in a structured protein environment.

Structural properties of a₃Y-K29H and a₃Y-K36H

Our second major goal was to demonstrate that Pourbaix diagrams derived from the tyrosine radical protein system reflect the *local* environment of the redox-active tyrosine. Overall, this goal represents a quite challenging task and was divided into three connected projects including the following: 1) To engineer and demonstrate the presence of a specific structural interaction at the site of the redox-active tyrosine. 2) To demonstrate that the structural properties of the re-engineered $\alpha_3 Y$ variants are appropriate in order to conduct voltammetry-based pH titrations. Thus, effects on the tyrosine radical potential from global events occurring in the protein scaffold, such as pH-induced changes in secondary and tertiary structures, must be minimized. In addition, the aggregation state of the protein should be pH independent to avoid a situation in which the distance between the radical site and the surface of the working electrode varies as a function of pH. This can lead to changes in the shape and position of the voltammogram representing the redox-active tyrosine and to an overall loss of the Faradaic signal.⁴⁷ 3) To derive potential *vs*. pH diagrams from $\alpha_3 Y$ and variants and determine the effects of the specifically engineered interaction on the tyrosine redox system. This study completes the first two objectives.

We re-engineered $\alpha_3 Y$ with the specific aim to place the imidazole group of a histidine residue close to the phenol ring of Y32 (Figs. S2 and 2). Eight $\alpha_3 Y$ -His variants were generated of which two displayed promising spectroscopic and structural characteristics. $\alpha_3 Y$ -K29H and $\alpha_3 Y$ -K36H each contain a histidine residue which protonation perturbs the fluorescence of Y32 (Table 1; Fig. 4). This observation suggests that the engineered histidine and the redox-active Y32 residue are in close proximity. Moreover, we could show that $\alpha_3 Y$ -K29H and $\alpha_3 Y$ -K36H are stable (Fig. S8: Table 1) and well-folded proteins whose

α-helical content (Fig. 3), tertiary structure (Figs. 5 and S10), solution aggregation state (Figs. 5, S9 and S10), and solvent-sequestered position of Y32 (Fig. 6) are pH independent or highly pH insensitive across a range of at least 3–4 pH units. These results demonstrate that we have achieved a tight structural control over the model protein hosting the redox-active tyrosine and that voltammetry measurements can be conducted across a broad pH range without large-scale structural changes occurring in the protein scaffold. Thus, the described model system is uniquely adapted to use voltammetry to study PCET reactions associated with tyrosine radical chemistry occurring in a solvent sequestered and well-structured protein environment. Such studies are described in a separate report.²²

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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REFERENCES

- 1. (a) Stubbe J, van der Donk WA. Chem. Rev. 1998; 98:705–762. [PubMed: 11848913] (b) Frey PA, Hegeman AD, Ruzicka FJ. Crit. Rev. Biochem. Mol. Biol. 2008; 43:63–88. [PubMed: 18307109]
- 2. Hoganson CW, Tommos C. Biochim. Biophys. Acta. 2004; 1655:116–122. [PubMed: 15100023]
- 3. Tommos C, Babcock GT. Biochim. Biophys. Acta. 2000; 1458:199–219. [PubMed: 10812034]
- 4. (a) Umena Y, Kawakami K, Shen J-R, Kamiya N. Nature. 2011; 473:55–60. [PubMed: 21499260]
 (b) Kawakami K, Umena Y, Kamiya N, Shen J-R. J. Photochem. Photobiol. B. 2011; 104:9–18. [PubMed: 21543235]
- (a) Proshlyakov DA, Pressler MA, DeMaso C, Leykam JF, DeWitt DL, Babcock GT. Science. 2000; 290:1588–1591. [PubMed: 11090359] (b) Hemp J, Robinson DE, Ganesan KB, Martinez TJ, Kelleher NL, Gennis RB. Biochemistry. 2006; 45:15405–15410. [PubMed: 17176062] (c) Gorbikova EA, Belevich I, Wikström M, Verkhovsky MI. Proc. Nat. Acad. Sci. U.S.A. 2008; 105:10733–10737.
- Aubert C, Vos MH, Mathis P, Eker APM, Brettel K. Nature. 2000; 405:586–590. [PubMed: 10850720]
- 7. (a) Sjöberg BM. Struct. Bond. 1997; 88:139–173.(b) Stubbe J, Nocera DG, Yee CS, Chang MCY. Chem. Rev. 2003; 103:2167–2201. [PubMed: 12797828]
- Reece SY, Hodgkiss JM, Stubbe J, Nocera DG. Phil. Trans. R. Soc. B. 2006; 1472:1351–1364. [PubMed: 16873123]
- Jensen LMR, Sanishvili R, Davidson VL, Wilmot CM. Science. 2010; 327:1392–1394. [PubMed: 20223990]
- Silva KE, Elgren TE, Que L, Stankovich MT. Biochemistry. 1995; 34:14093–14103. [PubMed: 7578006]
- 11. Rappaport F, Diner BA. Coord. Chem. Rev. 2008; 252:259–272.
- 12. Grabolle M, Dau H. Biochim. Biophys. Acta. 2005; 1708:209–218. [PubMed: 15878422]
- (a) Jeans C, Schilstra MJ, Klug DR. Biochemistry. 2002; 41:5015–5023. [PubMed: 11939798] (b) Buchta J, Grabolle M, Dau H. Biochim. Biophys. Acta. 2007; 1767:565–574. [PubMed: 17543884]
- 14. Mondal MS, Fuller HA, Armstrong FA. J. Am. Chem. Soc. 1996; 118:263-264.
- Byrdin M, Lukacs A, Thiagarajan V, Eker APM, Brettel K, Vos MH. J. Phys. Chem. A. 2010; 114:3207–3214. [PubMed: 19954157]

- Tommos C, Skalicky JJ, Pilloud DL, Wand AJ, Dutton PL. Biochemistry. 1999; 38:9495–9507. [PubMed: 10413527]
- 17. Tommos C. Phil. Trans. R. Soc. B. 2002; 357:1383–1394. [PubMed: 12437877]
- Dai Q-H, Tommos C, Fuentes EJ, Blomberg MRA, Dutton PL, Wand AJ. J. Am. Chem. Soc. 2002; 124:10952–10953. [PubMed: 12224922]
- 19. Hay S, Westerlund K, Tommos C. Biochemistry. 2005; 44:11891–11902. [PubMed: 16128591]
- 20. Dixon WT, Murphy DJ. Chem. Soc. Faraday. Trans. II. 1976; 72:1221-1230.
- (a) Huynh MHV, Meyer TJ. Chem. Rev. 2007; 107:5004–5064. [PubMed: 17999556] (b) Reece SY, Nocera DG. Annu. Rev. Biochem. 2009; 78:673–699. [PubMed: 19344235] (c) Warren JJ, Tronic TA, Mayer JM. Chem. Rev. 2010; 110:6961–7001. [PubMed: 20925411] (d) Dempsey JL, Winkler JR, Gray HB. Chem. Rev. 2010; 110:7024–7039. [PubMed: 21082865]
- 22. Berry BW, Martínez-Rivera MC, Tommos C. Submitted to the *Proceedings of the National Academy of Sciences USA*.
- 23. Ehrhardt MR, Erijman L, Weber G, Wand AJ. Biochemistry. 1996; 35:1599–1605. [PubMed: 8634291]
- 24. Santoro MM, Bolen DW. Biochemistry. 1988; 27:8063-8068. [PubMed: 3233195]
- 25. Markley JL, Bax A, Arata Y, Hilbers CW, Kaptein R, Sykes BD, Wright PE, Wüthrich K. J. Mol. Biol. 1998; 280:933–952. [PubMed: 9671561]
- 26. Krezel A, Bal W. J. Inorg. Biochem. 2004; 98:161–166. [PubMed: 14659645]
- 27. Cavanagh, J.; Fairbrother, WJ.; Palmer, AG.; Rance, M.; Skelton, NJ. Protein NMR spectroscopy: Principles and practice. 2nd. USA: Elsevier; 2006.
- Jeener J, Meier BH, Bachmann P, Ernst RR. (1979) Investigation of exchange processes by 2dimensional NMR-spectroscopy. J. Chem. Phys. 1979; 71:4546–4553.
- 29. Bard, AJ.; Faulkner, LR. Electrochemical methods: Fundamentals and applications. 2nd. USA: John Wiley & Sons, Inc; 2001.
- 30. Parry EP, Osteryoung RA. Anal. Chem. 1965; 37:1634–1637.
- 31. Harriman A. J. Phys. Chem. 1987; 91:6102-6104.
- 32. DeFelippis MR, Murthy CP, Broitman F, Weinraub D, Faraggi M, Klapper MH. J. Phys. Chem. 1991; 95:3416–3419.
- Costentin C, Louault C, Robert M, Savéant J-M. Proc. Nat. Acad. Sci U.S.A. 2009; 106:18143– 18148.
- 34. (a) Strickland EH, Wilchek M, Horwitz J, Billups CJ. J. Biol. Chem. 1972; 247:572–580.
 [PubMed: 5009702] (b) Lee JK, Ross RT. J. Phys. Chem. B. 1998; 102:4612–4618.(c) Noronha M, Lima JC, Lamosa P, Santos H, Maycock C, Ventura R, Macanita AL. J. Phys. Chem. A. 2004; 108:2155–2166.
- 35. Berry BW, Elvekrog MM, Tommos C. J. Am. Chem. Soc. 2007; 129:5308–5309. [PubMed: 17417844]
- 36. (a) Lakowicz, JR. Principles of Fluorescence Spectroscopy. New York, USA: Plenum Press; 1983.
 (b) Willis KJ, Szabo AG. J. Phys. Chem. 1991; 95:1585–1589.(b) Lee JK, Ross RT, Thampi S, Leurgans S. J. Phys. Chem. 1992; 96:9158–9162.
- Westerlund K, Berry BW, Privett HK, Tommos C. Biochim. Biophys. Acta. 2005; 1707:103–116. [PubMed: 15721609]
- 38. Sjöberg B-M, Reichard P, Gräslund A, Ehrenberg A. J. Biol. Chem. 1977; 252:536–541. [PubMed: 188819]
- 39. (a) Sjödin M, Styring S, Åkemark B, Sun L, Hammarström L. J. Am. Chem. Soc. 2000; 122:3932–3936.(b) Rhile IJ, Mayer JM. J. Am. Chem. Soc. 2004; 126:12718–12719. [PubMed: 15469234]
 (c) Benisvy L, Bittl R, Bothe E, Garner CD, McMaster J, Ross S, Teutloff C, Nesse F. Angew. Chem. Int. Ed. 2005; 44:5314–5317.(d) Costentin C, Robert M, Savéant J-M. J. Am. Chem. Soc. 2006; 128:4552–4553. [PubMed: 16594674] (e) Markle TF, Rhile IJ, DiPasquale AG, Mayer JM. Proc. Nat. Acad. Sci. U.S.A. 2008; 105:8185–8190.(f) Moore GF, Hambourger H, Kodis G, Michl W, Gust D, Moore TA, Moore AL. J. Phys. Chem. B. 2010; 114:14450–14457. [PubMed: 20476732] (g) Bonin J, Costentin C, Robert M, Routier M, Savéant J-M. Proc. Nat. Acad. Sci. U.S.A. 2010; 107:3367–3372.

- (a) Sibert R, Jesowicz M, Porcelli F, Veglia G, Range K, Barry BA. J. Am. Chem. Soc. 2007; 129:4393–4400. [PubMed: 17362010] (b) Sibert R, Jesowicz M, Barry BA. ACS Chem. Biol. 2010; 58:1157–1168. [PubMed: 20919724]
- 41. Dutton PL. Methods Enzymol. 1978; 54:411-435. [PubMed: 732578]
- 42. (a) Moss D, Nabedryk E, Breton J, Mäntele W. Eur. J. Biochem. 1990; 187:565–572. [PubMed: 2154376] (b) Rich PR, Iwaki M. Mol. Biosyst. 1997; 3:398–407. [PubMed: 17533453]
- 43. (a) Rusling JF. Acc. Chem. Res. 1998; 31:363–369.(b) Armstrong FA, Wilson GS. Electrochim. Acta. 2000; 45:2623–2645.(c) Armstrong FA. Curr. Opin. Chem. Biol. 2005; 9:110–117. [PubMed: 15811794]
- 44. Feitelson J, Hayton E. J. Phys. Chem. 1973; 77:10-15.
- 45. (a) Kálmál L, LoBrutto R, Allen JP, Williams JC. Nature. 1999; 402:696–699.(b) Bellér G, Lente G, Fábián I. Inorg. Chem. 2010; 49:3968–3970. [PubMed: 20415494]
- 46. (a) Chen P, McGreery RL. Anal. Chem. 1996; 68:3958–3965.(b) McGreery, RL. Electrochemical properties of carbon surfaces, Interfacial chemistry. Wiechowski, A., editor. USA: Dekker; 1999.
- 47. (a) Nicholson RS, Shain I. Anal. Chem. 1964; 36:706–723.(b) Laviron E. J. Electrochem. Chem. 1979; 101:19–28.(c) Tender L, Carter MT, Murray RW. Anal. Chem. 1994; 66:3173–3181.

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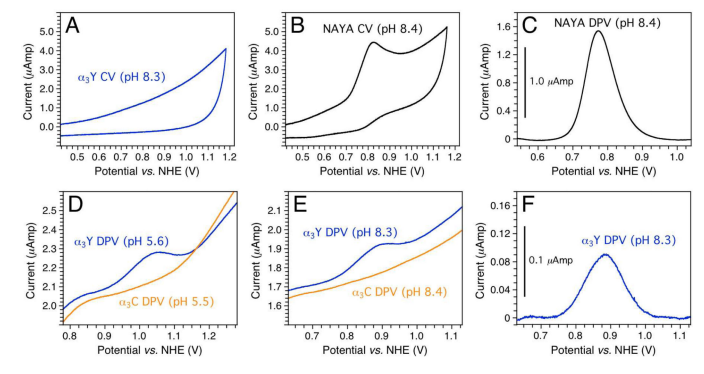
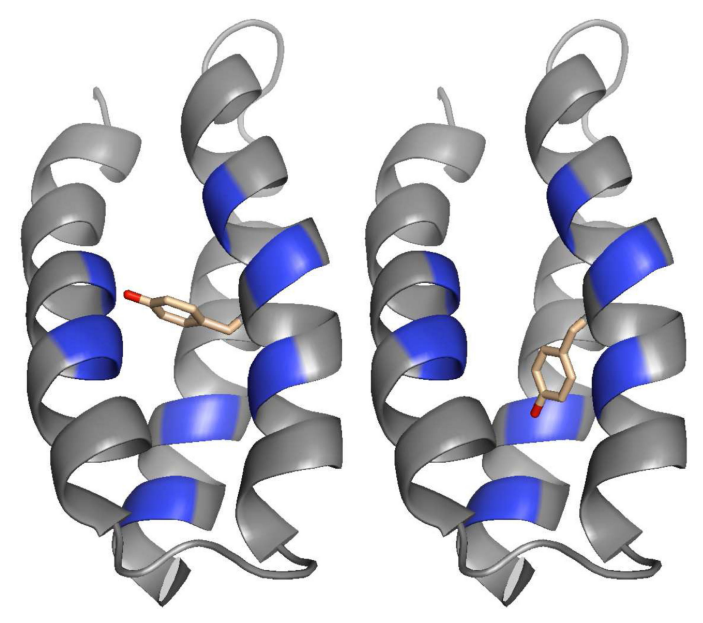


Fig. 1.

Electrochemical properties of $\alpha_3 Y$ and control samples. (A) Cyclic voltammogram of 210 μ M α_3 Y in 20 mM potassium phosphate, 20 mM sodium borate, 40 mM KCl, pH 8.33; scan rate 200 mV/s, iR-compensation 103 ohm. (B) Cyclic voltammogram of 200 µM NAYA in 20 mM potassium phosphate, 20 mM sodium borate, 200 mM KCl, pH 8.37; scan rate 200 mV s⁻¹, iR-compensation 103 ohm. (C) Differential pulse voltammogram of 200 µM NAYA in 20 mM potassium phosphate, 20 mM sodium borate, 200 mM KCl, pH 8.37; interval time 0.1 s, step potential 1.05 mV, scan rate 10.5 mV s^{-1} , modulation time 8 ms, modulation amplitude 50 mV. The trace has been baseline corrected. (D) Differential pulse voltammograms of (blue) 210 µM a₃Y in 20 mM sodium acetate, 20 mM potassium phosphate, 40 mM KCl, pH 5.56 and (orange) 200 µM a₃C in 20 mM sodium acetate, 20 mM potassium phosphate, 40 mM KCl, pH 5.46; interval time 0.1 s, step potential 1.05 mV, scan rate 10.5 mV s⁻¹, modulation time 5 ms, modulation amplitude 50 mV. (E) Differential pulse voltammograms of (blue) 210 µM a₃Y in 20 mM potassium phosphate, 20 mM sodium borate, 40 mM KCl, pH 8.33 and (orange) 200 µM a₃C in 20 mM potassium phosphate, 20 mM sodium borate, 40 mM KCl, pH 8.45; interval time 0.1 s, step potential 1.05 mV, scan rate 10.5 mV s⁻¹, modulation time 8 ms, modulation amplitude 50 mV. (F) Baseline-corrected trace of the $\alpha_3 Y$ differential pulse voltammogram shown in Panel E.





 α_3 Y models illustrating two possible orientations of the Y32 side chain. The α -carbons of residues changed in the α_3 Y-His variants are shown in blue.

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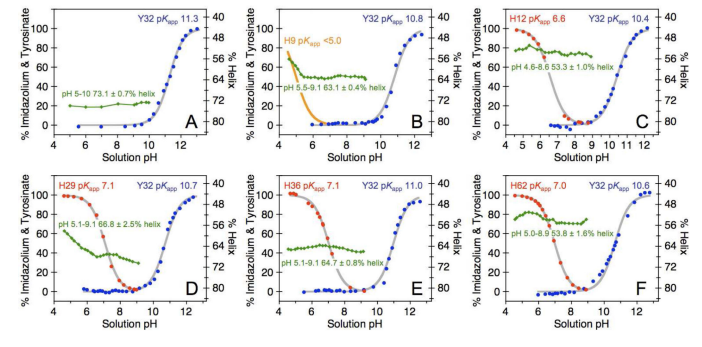


Fig. 3.

Physicochemical properties of $\alpha_3 Y$ and single-site histidine variants. pH titrations of Y32 (blue circles) and histidine residues (red circles) are shown for (A) $\alpha_3 Y$, (B) $\alpha_3 Y$ -V9H, (C) $\alpha_3 Y$ -L12H, (D) $\alpha_3 Y$ -K29H, (E) $\alpha_3 Y$ -K36H, and (F) $\alpha_3 Y$ -I62H. p K_{app} values were derived by nonlinear curve fitting (grey lines; Table 1). α -helical contents as a function of pH are shown in green (diamonds).

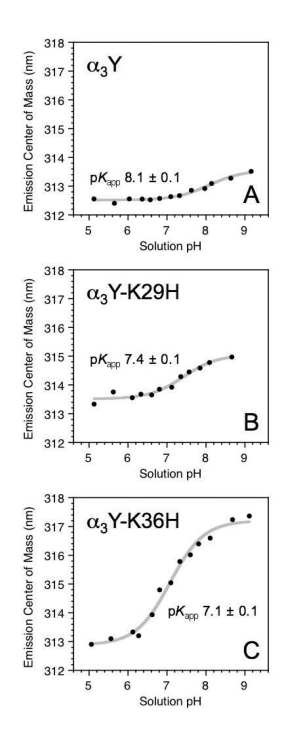


Fig. 4.

Fluorescence emission center of mass of Y32 in (A) α_3 Y, (B) α_3 Y-K29H and (C) α_3 Y-K36H as a function of pH. p K_{app} values were derived by nonlinear curve fitting (grey lines; Table 1).

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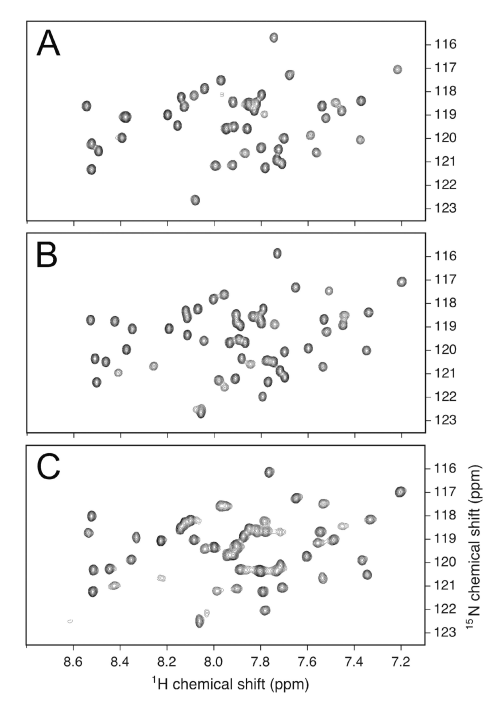
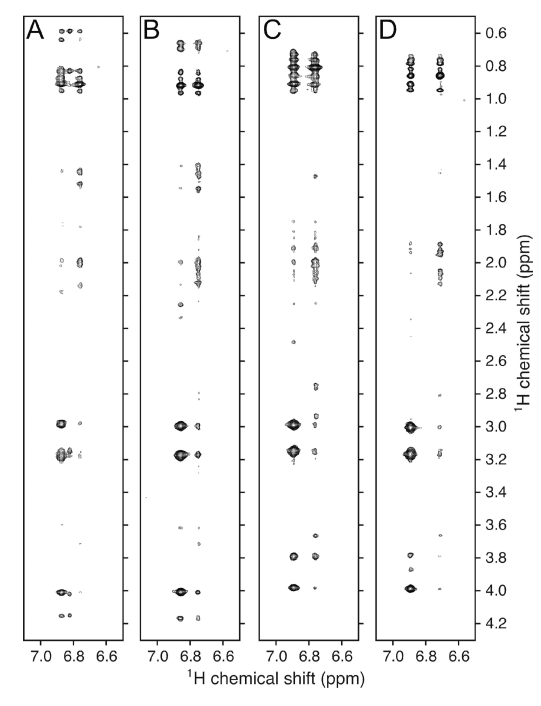
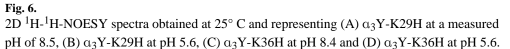


Fig. 5.

2D ¹⁵N-HSQC spectra of α_3 Y-K29H obtained at 25° C and with the pH at (A) 8.5, (B) 7.0 and (C) 5.5. These spectra, as well as those shown in Fig. S10, display a single set of peaks with no evidence of shadow peaks that would be indicative of minor conformers.





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Protein	$pK_{app} Y32^b$	pK_{app} His ^b	p K_{app} Y32 b p K_{app} His b Y32 Em(pH) c [Θ] ₂₂₂ d % Helix e	$[\Theta]_{222}^{d}$	% Helix ^e	5
$\alpha_3 Y$	11.3	I	8.1	-21.9	-21.9 73.1 ± 0.7 (pH 5.0-10.0) -3.7	-3.7
$a_3 Y$ -V9H	10.8	<5.0	n.d.	-18.7	$63.1 \pm 0.4 \text{ (pH } 5.5-9.1\text{)}$	-3.0
α_3 Y-L12H	10.4	6.6	n.d.	-16.0	$53.3 \pm 1.0 \text{ (pH 4.6-8.6)}$	n.d.
$\alpha_3 Y\text{-}K29H$	10.7	7.1	7.4	-20.4	$66.8 \pm 2.5 \text{ (pH 5.1–9.1)}$	-2.8
α_3 Y-K36H	11.0	7.0	7.1	-19.5	$64.7 \pm 0.8 \; (pH \; 5.1-9.1)$	-2.4
α ₃ Υ-I62H 10.6	10.6	7.0	n.d.	-16.5	$53.8 \pm 1.6 \text{ (pH } 5.0-8.9\text{)}$	n.d.

b Apparent tyrosinate/tyrosine and imidazole/imidazolium pKa values of Y32 and histidine residues obtained by fitting the pH-titration curves in Fig. 3 to a single pKa. Statistical errors 0.1.

 c Apparent pKa obtained by fitting the pH-titration curves in Fig. 4 to a single pKa. Statistical errors 0.1.

 d Mean residue ellipticity measured at pH 8.2 and 25° C. The α_3 W reference displays a [Θ]222 value of -22.6×10^3 deg cm² dmol⁻¹ at the same conditions.

 e Scaled relative to $\alpha 3$ W (76 \pm 1% $\alpha\text{-helical pH}$ 4–10),16,18

 $f_{
m Global}$ protein stabilities obtained by fitting the urea-denaturation curves in Fig. S8A. Data recorded at pH 8.2 and 25° C. Fitting standard errors < 0.03 kcal mol⁻¹.