Effect on Intrinsic Peroxidase Activity of Substituting Coevolved Residues from Ω -loop C of Human Cytochrome c into Yeast Iso-1-Cytochrome c

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ABSTRACT

Naturally-occurring variants of human cytochrome c (Cytc) that induce thrombocytopenia IV occur within Ω -loop C (residues 40 – 57). These variants enhance the peroxidase activity of human Cytc apparently by facilitating access to the heme by destabilizing Ω -loops C and D (residues 70 - 85). Given the importance of peroxidase activity in the early stages of apoptosis, we identified three sites with the EV mutation algorithm in or near Ω -loop C that coevolve and differ between yeast iso-1-Cytc and human Cytc. We prepared iso-1-Cytc variants with all possible combinations of the S40T, V57I and N63T substitutions to determine if these residues decrease the peroxidase activity of iso-1-Cytc to that of human Cytc producing an effective off state for a peroxidase signaling switch. At pH 6 and above, all variants significantly decreased peroxidase activity. However, the correlation of peroxidase activity with local and global stability, expected if cooperative unfolding of Ω -loops C and D is required for peroxidase activity, was generally poor. The *m*-values derived from the guanidine hydrochloride dependence of the kinetics of imidazole binding to horse Cytc, which is well-characterized by native-state hydrogen exchange methods, and K72A/K73A/K79A iso-1-Cytc show that local structural fluctuations and not subglobal cooperative unfolding of Ω -loops C and D are sufficient to permit binding of a small molecule like peroxide to the heme. A 2.46 Å structure of N63T iso-1-Cytc identifies a change to a hydrogen bond network linking Ω -loops C and D that could modulate the local fluctuations needed for the intrinsic peroxidase activity of Cytc.

Key Words

Cytochrome *c*, Protein dynamics, Peroxidase activity, Alkaline conformational transition, Apoptosis, Imidazole heme binding

1. Introduction

Cytochrome c (Cytc) is usually found near the mitochondrial membrane and is part of the Electron Transport Chain [1]. The heme of Cytc allows it to do redox chemistry, transporting electrons between Complex III and Complex IV. It also leaves the mitochondria as part of the intrinsic pathway of apoptosis [2,3]. Via its alternate function as a peroxidase, Cytc catalyzes the oxidation of the inner mitochondrial membrane lipid cardiolipin (CL), which causes CL to migrate to the outer mitochondrial membrane facilitating the release of Cytc from mitochondria [4]. Upon entering the cytosol, Cytc induces formation of a protein complex called the apoptosome initiating a cascade of caspases that ultimately leads to cell death [3,5].

Various mutations in Cytc are linked to thrombocytopenia type IV (THC4), which is a disease that is mainly characterized by low levels of blood platelets. Three specific mutations linked to THC4 have been discovered: G41S [6], Y48H [7], and A51V [8-10]. These mutations have been shown to promote peroxidase activity and to increase dynamics in omega loops C and D (residues 40-57 and 70-85, respectively) in the human protein [10-13]. However, recent molecular dynamics (MD) simulations on the THC4 variants indicate no correlation between the flexibility of Ω -loop C and peroxidase activity and in fact a negative correlation with the flexibility of Ω -loop D [14]. Peroxidase activity is also higher in yeast iso-1-Cytc when compared to human Cytc and other higher eukaryotes like horse [15-17]. The higher peroxidase activity of yeast iso-1-Cytc may function to convert H₂O₂ to H₂O, leading to protection from H₂O₂-induced apoptosis in yeast [18].

The hypothesis driving the current work is that the increased dynamics in the omega loops of Cytc leads to a more accessible heme and therefore higher peroxidase activity seen in both

variant forms of human Cytc and wild type yeast iso-1-Cytc. For a substrate like peroxide to access the heme iron, Met80 must be removed from its coordination site proximal to the heme iron to create a pentacoordinate species [19]. Previous work has shown that the oxidation of Met80 to a sulfoxide both enhances the peroxidase activity of Cytc [19-22] and allows hydrogen peroxide to access the heme more readily [20,21]. Cytc variants like those linked to THC4 in Ω -loop C more readily react with H_2O_2 to produce the Met80 sulfoxide and a direct correlation between the presence of Met80 sulfoxide and peroxidase activity is observed [22]. The role of Ω -loops C and D in mediating the dynamics needed to induce peroxidase activity is not surprising because they have been shown to be the two least stable substructures of horse Cytc [23,24].

Certain residues in the human Cytc omega loops appear to have coevolved (Fig. 1) [25]. For Ω -loop C, pairwise coevolution is most significant for residues Thr40 and Ile57 of human Cytc. Being at opposite ends of Ω -loop C, the side chains of these residues are in direct contact (Fig. 1). In yeast iso-1-Cytc, these positions are occupied by residues with one less methyl group (Ser40 and Val57). Equally significant pairwise coevolution occurs between Ile57 and Lys55. Other significant pairwise coevolution is observed between Ile57 and Thr63, Thr63 and Tyr74, and Thr40 and Lys55. As a group, Thr40, Lys55, Ile57, Thr63, and Tyr74 are proximal in the structure of human Cytc at the interface between Ω -loops C and D (Fig. 1). Lys55 and Tyr74 do not change between yeast iso-1-Cytc and human Cytc, although Asn63 in yeast iso-1-Cytc evolves to Thr63 in human Cytc.

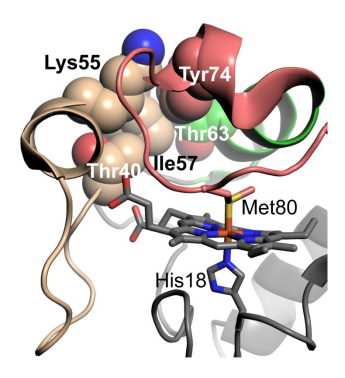


Fig. 1. Visualization of human Cytc (PDB ID: 3ZCF) [26] residues (space-filling models) in or near Ω -loop C (shown in wheat) that have coevolved. Ω -loop D is shown in salmon and the 60s helix in green. Shown as stick models are the heme and its axial ligands Met 80 and His 18.

As mentioned previously these residues are in a part of the protein important for regulating peroxidase activity [9-13,27]. We hypothesized that as eukaryotes evolved, tighter control of the intrinsic peroxidase activity Cytc was required, so that it could act as a signaling switch to initiate apoptosis. Therefore, the residues that have changed between the yeast and human proteins may be responsible for the large differences in intrinsic peroxidase activity seen between Cytc from these two species [27]. More specifically, coevolution at the interface of Ω -loops C and D has led to residues that are bulkier and more hydrophobic in human Cytc than in yeast iso-1-Cytc, which could limit the dynamics of these omega loops and therefore hinder heme access.

In the current work, we have mutated the coevolved Ω -loop C/D interface residues of yeast iso-1-Cytc to the amino acids observed in the human protein. These substitutions are S40T, V57I, and N63T, as well as all double substitution combinations and the triple substitution variant. Our goal was to better understand how these mutations affect the peroxidase activity of Cytc and the stability of the local substructures involved, and therefore how the intrinsic peroxidase activity of Cytc decreased as eukaryotes evolved.

2. Experimental section

2.1. Site-directed mutagenesis

Mutagenesis was done using the QuikChange Lightning mutagenesis kit (Agilent Technologies). All variants of yeast iso-1-Cytc contain a background mutation of C102S (WT) [28], which prevents intermolecular disulfide dimerization. To better mimic the human protein, we have chosen not to include a K72A mutation (variant usually referred to as WT*), which is often included because Lys72 is trimethylated in its native host, *Saccharomyces cerevisiae*, and is not when iso-1-Cytc is expressed from *Escherichia coli* [29]. For imidazole binding experiments, a K72A/K73A/K79A variant was prepared using DNA containing the WT* variant [30]. The template DNA for mutagenesis to produce all iso-1-Cytc variants was the pRbs_BTR1 vector [31], a derivative of pBTR1 [29,32] carrying an optimized ribosomal binding site [33]. Sequencing to confirm mutations was done either at the University of Montana Genomics Core or by Eurofins Genomics (Louisville, KY).

2.2. Protein expression and purification

Each pRbs_BTR1 vector carrying the desired mutated iso-1-Cytc gene was used to transform competent phage T1 resistant *E. coli* BL21(DE3) cells (New England Biolabs). Transformed

cells were inoculated into 2xYT media and grown for 30-36 hours at 37 °C with shaking at 150 rpm. Cells were pelleted and then suspended in lysis buffer consisting of 10 mM Tris, pH 7.5, 500 mM NaCl, 2 mM PMSF, and a small amount of DNAse. Cells were then lysed via sonication using a Qsonica Q700 sonicator. The lysate was centrifuged and then incubated overnight at 4 °C in 50% ammonium sulfate to precipitate out most protein impurities, which were removed via centrifugation. The supernatant was subjected to two rounds of dialysis for 8 to 24 hours each in buffer consisting of 12.5 mM sodium phosphate, pH 7.2, 1 mM disodium EDTA, and 2 mM β-mercaptoethanol (βME). The dialyzed solution was then loaded onto a CM Sepharose column equilibrated to 50 mM sodium phosphate, pH 7.2, 1 mM EDTA, and 2 mM BME. The protein was eluted using a linear gradient of the same buffer with 0.8 M NaCl added. The eluate was then exchanged into a low salt solution consisting of 50 mM sodium phosphate, pH 7.0, before flash freezing and storing in -80 °C. At most 24 hours before experiments, the protein was thawed and further purified using a HiTrap HP SP 5 mL cation exchange column attached to an AKTAprime plus chromatography system (GE Healthcare Life Sciences) and was eluted using a 50 mL 0-50 %B gradient with 50 mM sodium phosphate buffer, pH 7.0, as buffer A and 50 mM sodium phosphate buffer, pH 7.0, with 1 M NaCl added as buffer B. The eluate was concentrated by centrifuge ultrafiltration (Amicon Ultra-15, 10k MWCO, Millipore) and the high salt removed by several rounds of exchange into MilliQ water.

2.3. Alkaline conformational transition

To evaluate the alkaline transition, each iso-1-Cytc variant was oxidized with potassium ferricyanide at room temperature for 20-30 minutes. The protein was then separated from the potassium ferricyanide using a G25 size exclusion column (GE Healthcare) with 200 mM NaCl as the running buffer. The solution at the start of the experiment had a final protein concentration

of 100 μ M in 100 mM NaCl. As HCl or NaOH was added to change the pH of the solution, an equal amount of a 2-fold concentrated stock solution (200 μ M protein in 200 mM NaCl) was added to maintain the concentrations of protein and NaCl in the titration solution. Absorbance measurements were collected using a Beckman Coulter DU 800 UV-Vis instrument. Wavelength scans were from 750 nm to 500 nm in a quartz suprasil cuvette. A Denver Instrument DU-10 pH meter with a Fisher Scientific Accumet Double Junction Semi-Micro pH probe was used for pH measurements. All measurements were taken at 21 ± 2 °C. Absorbance at 695 nm, A₆₉₅, was used to monitor the conformational transition. Subtraction of 750 nm absorbance values, A₇₅₀, was used as a baseline correction to account for instrument drift during the period of the titration. Corrected 695 nm absorbance values, A_{695corr} (A₆₉₅ – A₇₅₀), were then plotted as a function of pH, and a modified form of the Henderson-Hasselbalch equation was fit to the data that allows the number of protons associated with the alkaline transition (*n*) to be determined as well as p K_a values for the alkaline transition [27].

2.4. Guanidine hydrochloride denaturation

Iso-1-Cytc variants were oxidized as described for the alkaline transition, except that the buffer for G25 chromatography was 20 mM Tris, 40 mM NaCl and 1 mM EDTA at pH 7.5.

Denatured (D) and Native (N) stocks were prepared with final concentrations of 4 µM protein in 20 mM Tris, 40 mM NaCl, and 1 mM EDTA at pH 7.5. The D stock also contained 6 M guanidine hydrochloride (GdnHCl). For circular dichroism (CD) measurements, 2 mL N stock was placed in a 1 cm square quartz cuvette with a stir bar and tubing for withdrawing N stock and adding D stock, and with masking to ensure that light does not scatter off the stir bar or tubing. The titration was carried out by increasing GdnHCl concentration in 0.1 M increments with a Hamilton titrator interfaced to the CD spectrometer. Ellipticity measurements were taken

using an Applied Photophysics Chirascan spectrometer at 25 °C. Data were collected at wavelengths of 250 nm and 222 nm, with 250 nm ellipticity values subtracted from the 222 nm values as a baseline correction, $\theta_{222\text{corr}}$. A two-state model assuming a linear relationship between the free energy of unfolding, ΔG_u , and GdnHCl concentration (eq 1) was fit to plots of $\theta_{222\text{corr}}$ vs GdnHCl concentration using nonlinear least squares methods.

(1)
$$\theta_{222\text{corr}} = \frac{\theta_N + (\theta_D + m_D[\text{GdnHCl}]) * exp^{\frac{m[\text{GdnHCl}] - \Delta G_u^{0'}(\text{H}_2\text{O})}{RT}}}{1 + exp^{\frac{m[\text{GdnHCl}] - \Delta G_u^{0'}(\text{H}_2\text{O})}{RT}}}$$

In eq 1, $\theta_{\rm N}$ is the native state baseline, $\theta_{\rm D}$ and $m_{\rm D}$ are the intercept and slope of the denatured state baseline, m is the rate of change of $\Delta G_{\rm u}$ with respect to GdnHCl concentration and $\Delta G_{\rm u}^{\rm o'}({\rm H_2O})$ is the free energy of unfolding in the absence of GdnHCl.

2.5. Peroxidase activity measurements

Iso-1-Cytc variants were oxidized as described for the alkaline transition. Peroxidase activity was measured by the oxidation of guaiacol to tetraguaiacol [34], which absorbs strongly at 470 nm. Different 50 mM buffers were used for different pH values: sodium acetate for pH 5, 2-(N-morpholino)ethanesulfonic acid (MES) for pH 6, sodium phosphate for pH 7, and tris(hydroxymethyl)aminomethane (Tris) for pH 8. Each buffer was adjusted to the desired pH using HCl or NaOH. The solutions were degassed under argon for 30-45 minutes to limit non-enzymatic oxidation of guaiacol by O_2 . The degassed solutions were then used to make three separate solutions of 4 μ M protein, 400 μ M guaiacol, and 100 ± 5 mM hydrogen peroxide such that H_2O_2 is above the Michaelis constant, K_M , with respect to H_2O_2 for peroxidase activity (~37 mM for human Cytc) [35,36]. The concentrations of guaiacol and hydrogen peroxide were

measured by absorbance using $\varepsilon_{274} = 2150 \text{ M}^{-1} \text{ cm}^{-1} [37]$, and $\varepsilon_{240} = 41.5 \text{ M}^{-1} \text{ cm}^{-1} [38,39]$, respectively. The protein solution, the degassed buffer, and differing amounts of the guaiacol solution were mixed into a single solution immediately prior to measurements. This solution and the hydrogen peroxide solution were mixed 1:1 with an Applied Photophysics SX20 stoppedflow spectrometer and time-dependent changes in absorbance at 470 nm (A₄₇₀) were recorded. Final protein concentration was 1 µM, final hydrogen peroxide concentration was 50 mM in 50 mM buffer. Data were collected at 25 ± 0.1 °C. For each trial, A_{470} was plotted as a function of time and the linear region of the resulting graph with the highest slope was used to determine the initial enzyme velocity, v. Five runs were done for each guaiacol concentration, and the slopes were averaged then multiplied by 4 (oxidation of guaiacol requires removal of 4 electrons [34,40], divided by the protein concentration and the reported extinction coefficient [34,40,41] of 26.6 mM⁻¹ cm⁻¹ for tetraguaiacol (3,3'-dimethoxy-4,4'-biphenoquinone) [42] to obtain v/[Cytc] values at each guaiacol concentration. These v/[Cytc] values were plotted as a function of guaiacol concentration and the Michaelis-Menten equation was fit to the data to obtain the Michaelis constant, $K_{\rm M}$, and the catalytic rate constant, $k_{\rm cat}$.

2.6. Equilibrium and kinetics of imidazole binding

Imidazole binding experiments were carried out either by equilibrium titration monitored by UV-Vis spectroscopy (Beckman DU 800 UV-Vis Spectrometer) or by stopped-flow mixing methods (Applied Photophysics SX20 Stopped-flow Spectrometer). Horse heart Cytc (Sigma used without further purification) and K72A (WT*) and K72A/K73A/K79A variants of yeast iso-1-Cytc were used for these experiments. All experiments were done in 20 mM 3-(N-morpholino)propanesulfonic acid (MOPS), pH 7 at constant ionic strength. NaCl was used to maintain constant ionic strength as the concentration of imidazole was varied.

For equilibrium experiments, individual samples were prepared for each data point by mixing together two solutions at the same ionic strength, both containing 10 μ M Cytc, one containing imidazole and one not containing imidazole. The solutions were mixed in proportions to produce a set of solutions covering the desired range of imidazole concentrations. Samples were equilibrated at room temperature (21 \pm 2 °C) for at least two hours before measuring UV-Vis spectra.

Stopped-flow experiments were carried out by 1:1 mixing of Cytc with imidazole. Both solutions contained 20 mM MOPS, pH 7. The concentration of the imidazole solution was varied. The ionic strengths of both the Cytc and imidazole solutions were adjusted to the same value with NaCl. The final concentration of Cytc after mixing was 10 μM. Stopped-flow experiments in the presence of GdnHCl were carried out at an ionic strength of 1.25 M with GdnHCl replacing NaCl to keep ionic strength constant in the protein and imidazole containing solutions. Binding of imidazole was followed at 405 nm, A₄₀₅, as a function of time. Exponential functions were fit to plots of A₄₀₅ versus time.

2.7. Crystallization and structure determination of N63T iso-1-Cytc

The N63T variant of iso-1-Cytc was purified as described above, oxidized with 5 mg of K₃[Fe(CN)₆] per mg of protein, separated from K₃[Fe(CN)₆] using a sephadex G25 column equilibrated to and run with 50 mM sodium phosphate pH 6.0, and then concentrated to 37.2 mg/mL using an Amicon Ultra-15 (10k MWCO) centrifuge ultrafiltration device. Qiagen's PEGs I Suite, PEGs II Suite, JCSG Core III Suite, and JCSG Core IV Suite were used to screen for crystallization conditions. Needle-like crystals were obtained from 1:1 sitting drops incubated at 20 °C for several screening conditions including: PEGs II-16 (0.1 M Tris pH 8.5, 30% w/v

polyethylene glycol (PEG) 1000), JCSG III-33 (0.1 M 4-(2-hydroxyethyl)-1piperazineethanesulfonic acid (HEPES) pH 7.5, 30% v/v PEG 400, 5% v/v PEG 3000, 10% glycerol), JCSG III-61 (0.1 M MES pH 6.0, 40% v/v PEG 400, 5% PEG 3000) and JCSG IV-40 (0.1 M HEPES pH 7.5, 50% v/v PEG 200). Crystals were harvested directly from the crystal trial plates and cryoprotected with 20% glycerol. Crystals that diffracted to 2.46 Å resolution were obtained from the PEGs II-16 screening condition. X-ray diffraction data were collected at the Stanford Synchrotron Radiation Lightsource beamline 12-2 with a DECTRIS PILATUS 6M detector. The data were indexed, integrated and scaled in the P4₁ space group using iMosflm [43] and Aimless [44]. The structure was solved by molecular replacement using Molrep [45], integrated into the CCP4i2 software suite [46]. Model building was accomplished in REFMAC [47] and PHENIX [48] and the structure was refined through iterative cycles of manual adjustment in Coot [49] and refinement of atomic positions, real space, occupancy, and thermal parameters in PHENIX, yielding $R_{\text{work}} = 0.165$ and $R_{\text{free}} = 0.217$. Data collection and refinement statistics are provided in Table S1. Coordinates have been deposited at the Protein Data Bank (www.rcsb.org) under PDB code: 7MRI. All figures were produced with PyMol [50].

3. Results

3.1 Stability of iso-1-Cytc variants

Fig. 2 shows typical GdnHCl denaturation data for several iso-1-Cytc variants with humanlike mutations. It is evident from the data in Fig. 2 that the effects of humanlike substitutions at positions 40, 57 and 63 are modest. In these plots, the open data points were not used in data fitting because of the initial downward curvature of ellipicity at 222 nm at low GdnHCl concentrations. This effect is a result of the high charge density of iso-1-Cytc, which causes initial additions of GdnHCl to stabilize helical structure [51,52]. To further mitigate the

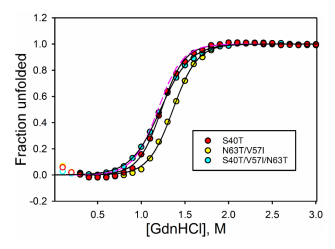


Fig. 2. Plots of fraction of protein unfolded for selected iso-1-Cytc variants as a function of GdnHCl concentration. Fraction unfolded was calculated from fits of eq 1 to $\theta_{222\text{corr}}$ vs GdnHCl concentration data. Open circles are data points that were not used in the fit. Solid curves are fits of a two-state model (eq 1) to the data. Data are shown for the S40T (red circles), N63T/V57I (yellow circles) and S40T/V57I/N63T (cyan circles) variants. The dashed magenta line is fraction unfolded for WT iso-1-Cytc using the parameters in Table 1.

Table 1Thermodynamic parameters for GdnHCl unfolding of iso-1-Cytc variants at 25 °C.

Variant	ΔG _u °′(H ₂ O) (kcal/mol)	m (kcal/mol×M)	C _m (M)
WT ^a	5.05 ± 0.30	4.24 ± 0.13	1.19 ± 0.04
V57I	4.22 ± 0.13	3.54 ± 0.12	1.193 ± 0.007
S40T	4.99 ± 0.09	3.94 ± 0.16	1.27 ± 0.03
N63T	5.38 ± 0.12	4.01 ± 0.10	1.343 ± 0.005
N63T/V57I	5.40 ± 0.16	3.87 ± 0.12	1.40 ± 0.03
S40T/V57I	4.23 ± 0.07	3.66 ± 0.22	1.16 ± 0.06
S40T/N63T	4.50 ± 0.12	3.85 ± 0.07	1.17 ± 0.01
S40T/V57I/N63T	4.197 ± 0.004	3.49 ± 0.05	1.20 ± 0.02

^a Parameters from Duncan et al. 2009 [31].

effects of this behavior, a native state baseline that is invariant with GdnHCl concentration was used.

As can be seen from the parameters obtained from fitting eq 1 to the data in this manner (Table 1), the differences in global stability are small. Only the V57I, S40T/V57I, andS40T/V57I/N63T have significant changes in $\Delta G_u^{o\prime}(H_2O)$ relative to WT iso-1-Cytc. The m values for the variants are significantly smaller than that of WT iso-1-Cytc, decreasing to values similar to human Cytc (3.5 – 3.7 kcal/mol×M) [15]. The GdnHCl concentration at the midpoint of unfolding, C_m , increases for some of the single and double mutant variants, but is the same for WT iso-1-Cytc and the final S40T/V57I/N63T variant.

The GdnHCl denaturation results overall indicate that the global stability of the variants is not affected strongly with the exception of the V57I, S40T/V57I, and S40T/V57I/N63T variants. All of these variants contain the V57I mutation, which appears to be important for decreasing the *m*-value (compare S40T data to S40T/V57I/N63T data in Fig. 2). In general, all variants unfold less cooperatively than WT iso-1-Cytc, indicating that the differences between the human and yeast proteins may have more to do with changes in substructure dynamics and stability than with global stability.

3.2 Alkaline conformational transition of iso-1-Cytc variants

The alkaline transition of Cytc happens under mildly basic conditions when one of the lysine residues in Ω -loop D replaces the methionine that interacts with the iron of the heme. The p K_a of the alkaline transition can provide information about the local stability of Ω -loop D (residues 70 – 85) [53,54]. Fig. 3 shows typical data for the alkaline conformational transition of the iso-1-Cytc variants with humanlike substitutions at residues 40, 57 and 63, derived from spectra

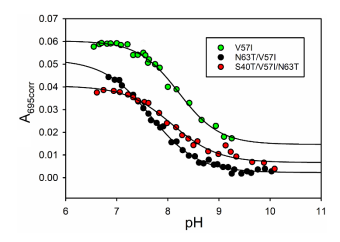


Fig. 3. Plot of corrected absorbance at 695nm, A_{695corr}, versus pH for iso-1-Cytc variants. Black lines are fits of a modified form of the Henderson-Hasselbalch equation to the data. Parameters from these fits are provided in Table 2. Data for the V57I (green circles), N63T/V57I (black circles), and S40T/V57I/N63T (red circles) are shown.

Table 2 Alkaline transition parameters of iso-1-Cytc variants at 25 °C.

23 C.		
Variant	pK_a	n
WT ^a	8.00 ± 0.05	0.98 ± 0.01
V57I	8.17 ± 0.06	1.13 ± 0.12
S40T	7.99 ± 0.06	0.98 ± 0.17
N63T	8.01 ± 0.05	0.95 ± 0.06
N63T/V57I	7.64 ± 0.04	0.96 ± 0.05
S40T/V57I	7.94 ± 0.16	1.05 ± 0.13
S40T/N63T	7.84 ± 0.11	0.98 ± 0.12
S40T/V57I/N63T	7.96 ± 0.10	0.89 ± 0.08
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^a Parameters from Duncan et al. 2009 [31].

measured from 500 to 750 nm (Fig. S1). As with the global unfolding of these variants, the effects on the local stability of Ω -loop D as measured by the alkaline transition appear to be small. Fits of the Henderson-Hasselbalch equation to alkaline transition data yield p K_a values

similar to the WT protein (Table 2). The N63T/V57I variant shows a significant decrease in the pK_a of the alkaline transition relative to WT iso-1-Cytc, whereas the V57I and S40T/N63T variants show a small increase and a small decrease, respectively, in the pK_a of the alkaline transition. The n values (Table 2) in all cases are approximately 1, as is expected based on the net deprotonation of a single lysine. Overall, the alkaline transition data indicate that the stability of the Ω -loop that contains Met80 and the nearby lysine residues is similar for WT iso-1-Cytc and all variants.

3.3 Peroxidase activity of iso-1-Cytc variants

The peroxidase activities of the variants of iso-1-Cytc were measured from pH 5 to 8 using guaiacol as substrate [15,16,27,34,55,56]. Michaelis-Menten plots for peroxidase activity at pH 8 are shown in Fig. 4 for theV57I, N63T/V57I, and S40T/V57I/N63T variants. More significant changes in k_{cat} are evident in Fig. 4 than might be expected based on the changes in the global and local stability described above. The K_{M} and k_{cat} values from fits of the Michaelis-Menten

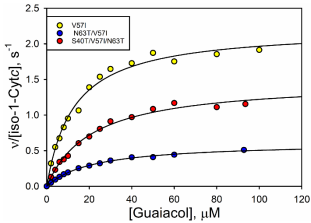


Fig. 4. Michaelis-Menten plots at pH 8 for variants V57I (yellow), N63T/V57I (blue) and S40T/V57I/N63T (red). Black lines are fits of the Michaelis-Menten equation to the data.

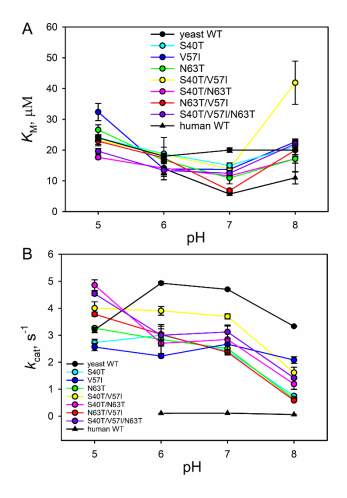


Fig. 5. Plots of (A) K_M and (B) k_{cat} versus pH for all variants. For comparison, K_M and k_{cat} values for WT yeast iso-1-Cytc (black circles) and WT human Cytc (black triangles) are provided.

equation to peroxidase activity data are provided in Table S2 and plotted in Fig. 5 as a function of pH for WT and variant forms of iso-1-Cytc and for human Cytc. The $K_{\rm M}$ values do not differ much among the WT and variant forms of yeast iso-1-Cytc. The $K_{\rm M}$ values for all variants and WT are higher at more acidic pH values, decrease as the pH becomes more alkaline until pH 8, where most variants show an increase relative to pH 7. With regard to $k_{\rm cat}$, the WT iso-1-Cytc value is in the middle of the range observed for the variants at pH 5, but at higher pH, the variants show significantly lower $k_{\rm cat}$ values relative to WT iso-1-Cytc. However, the decrease in $k_{\rm cat}$ for the variants does not approach that observed for human Cytc. Interestingly at pH 8, peroxidase activity as measured by $k_{\rm cat}$ becomes much lower for all the variants. Peroxidase

measurements at pH 9 were attempted, but the peroxidase activity for most variants was so low that it became difficult to distinguish signal from noise in the data. The pH 9 results are likely due to the increasing presence of the alkaline form of iso-1-Cytc, which cannot perform peroxidase reactions due to binding of the strong field ligand lysine to the heme [34].

3.4. Imidazole binding as a function of GdnHCl concentration

In hydrogen-deuterium exchange (HX) experiments, it is known that exchange can occur via subglobal unfolding or by lower energy local structural fluctuations [57]. Recent work on equine Cytc has shown that some protein dynamics may not be detectable by the HX methods used to study the dynamics of Ω -loops C and D [58]. The observation that the effects of the Ω -loop C mutations on peroxidase activity were more pronounced than their effects on local and global stability led us to test the extent of structural rearrangement necessary for a small ligand like peroxide to bind to the heme. In analogy to HX experiments on horse heart Cytc [23,24], the extent of unfolding of Ω -loop D can be assessed by measuring the GdnHCl-dependence of the free energy of binding (m-value) for the small exogenous ligand imidazole (Im). For these studies, we used horse heart Cytc to allow direct comparison with the m-value for opening of the Ω -loop D substructure as measured by HX methods [23,24]. We also used two variants of the yeast protein, K72A (WT*) and K72A/K73A/K79A. The latter variant was used to eliminate interference from alkaline conformers on the dynamics of Ω -loop D that mediate Im binding. The equilibria and kinetics of binding of Im as a function of ionic strength, I, were measured first to find optimal conditions for measurements as a function of GdnHCl concentration.

Equilibrium binding of Im to the WT* and the K72A/K73A/K79A iso-1-Cytc variants was measured by absorbance spectroscopy (Fig. S2). Data were analyzed as a function of total

imidazole concentration to allow direct comparison with values reported, previously [59,60]. Because it is deprotonated Im that binds to the heme of Cytc, and Im is only about 50% ionized at pH 7, all reported equilibrium constants are apparent equilibrium constants for Im binding. Measurements at ionic strengths of 0.05 to 1.25 M show that the apparent equilibrium constant for Im binding, $K_{\text{Im,equil}}$, increases slowly up to I = 1 M for WT* and then levels off (Fig. S3). The magnitude of $K_{\text{Im,equil}}$ for the K72A/K73A/K79A variant initially decreased as ionic strength increased, but then increased again leveling off at I = 1 M (Fig. S3). At I = 1.0 M, $K_{\text{Im,equil}}$ is 313 \pm 35 M⁻¹ for WT* iso-1-Cytc and 503 \pm 56 M⁻¹ for K72A/K73A/K79A iso-1-Cytc. These values are 20-fold and 30-fold larger, respectively, than the previously reported value of 15.3 \pm 0.5 M⁻¹ for the binding constant of Im for horse heart Cytc at I = 1.0 M [59].

A kinetic scheme involving an initial rearrangement of Cytc to a conformer competent for Im binding, Cytc[†], has been found to be consistent with the dependence of the observed rate constant, k_{obs} , on Im concentration [59,60]. In this mechanism (Scheme 1), k_{obs} levels off at high Im concentration because the observed rate constant, k_{obs} , for Im binding (eq 2) becomes limited

Cytc
$$\frac{k_1}{k_{-1}}$$
 Cytc[†] + Im $\frac{k_2}{k_{-2}}$ Cytc-Im

Scheme 1. Kinetics mechanism of Im binding to Cytc.

(2)
$$k_{\text{obs}} = \frac{k_1 \left(\frac{k_2}{k_{-1}}\right) \times [Im] + k_{-2}}{1 + \left(\frac{k_2}{k_{-1}}\right) \times [Im]}$$

by the rate constant, k_1 , for formation of Cyt c^{\dagger} . We investigated the kinetics of Im binding to WT* and K72A/K73A/K79A iso-1-Cytc and horse heart Cytc at 1.25 M ionic strength. Fits of the kinetic data are consistent with a major fast phase due to Im binding and a slow phase that is on the time scale expected for proline isomerization (Fig. S4 and Table S3) [61]. Proline

isomerization phases are also observed near neutral pH for the alkaline transition of iso-1-Cytc mediated by His73 in Ω-loop D for a K73H variant of iso-1-Cytc [62]. Therefore, we believe the slow phase is due to proline isomerization within the protein following imidazole binding. Plots of total amplitude at 405 nm versus total Im concentration show that the affinity of Im for yeast iso-1-Cytc is considerably higher than for horse heart Cytc (Fig. S5). The data for horse heart Cytc yield apparent binding constants similar to those reported by Sutin and Yandell at 1.0 M ionic strength (Table S4) [59]. Apparent binding constants from amplitude data for the yeast iso-1-Cytc variants are consistent with values from our equilibrium titrations (Table S4).

Plots of $k_{\rm obs}$ versus Im concentration for K72A/K73A/K79A iso-1-Cytc and horse heart Cytc at 1.25 M ionic strength fit well to eq 2 (Fig. S6, Fig. S7 and Table S5). However, the data for WT* iso-1-Cytc do not deviate detectably from linearity. Therefore, we used horse heart Cytc and K72A/K73A/K79A iso-1-Cytc to measure the GdnHCl dependence of the binding of Im to Cytc. Interestingly, in line with the ionic strength dependence of $K_{\rm Im,equil}$ observed for the yeast iso-1-Cytc variants, comparison of our kinetic data with previous data for horse heart Cytc suggest that the kinetics of Im binding to Cytc depend on ionic strength (Fig. S6). Therefore, measurements of the kinetics of Im binding as a function of GdnHCl concentration were carried out at a constant ionic strength of 1.25 M.

Fig. 6 shows plots of $k_{\rm obs}$ versus total Im concentration at 0, 0.25 and 0.50 M GdnHCl for K72A/K73A/K79A iso-1-Cytc and horse heart Cytc. It is evident that $k_{\rm obs}$ at each Im concentration increases as GdnHCl concentration increases and that overall the $k_{\rm obs}$ values are larger for the yeast protein compared to the horse protein. From plots of amplitude data as a function of total Im concentration, the affinity of Im for horse Cytc can be seen to increase as GdnHCl concentration increases (Fig. S8). Fits of the data to a single site binding model confirm

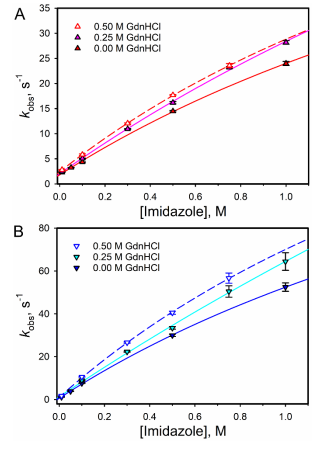


Fig. 6. Plots of k_{obs} versus Im concentration at 0, 0.25 and 0.50 M GdnHCl for (A) horse heart Cytc and (B) K72A/K73A/K79A iso-1-Cytc. All data were measured at a constant ionic strength of 1.25 M. Solid curves are fits of eq 2 to the data.

this observation for both horse heart Cytc and K72A/K73A/K79A iso-1-Cytc, although the error bars are large for the yeast variant (Table 3). Plots of the apparent free energy of Im binding, $\Delta G^{\circ\prime}_{Im}$ (Table 3), as a function of GdnHCl concentration are shown in Fig. 7A. The plots are linear yielding *m*-values in the range of 0.4-0.7 kcal mol⁻¹ M⁻¹ for both proteins (Table 4) indicating that Im binding causes only a small structural rearrangement. By comparison, unfolding of the Ω -loop D substructure as detected by hydrogen exchange [24,63] or the alkaline transition [53,64] yield GdnHCl *m*-values of 1.5-1.7 kcal mol⁻¹ M⁻¹.

Table 3 Thermodynamic parameters for imidazole binding to Cytc as a function of GdnHCl at I = 1.25 M, pH 7 and 25 °C.

Parameter	[GdnHCl] (M)	Protein	
		Horse Heart Cytc	K72A/K72A/K79A iso-1-Cyt <i>c</i>
$K_{ m Im,Amp}$ $^{ m a}$	0	17 ± 2	570 ± 430
(M)	0.25	20 ± 2	720 ± 490
	0.50	25 ± 3	1000 ± 760
$\Delta G^{\circ}{'_{\operatorname{Im}}}^{b}$	0	-1.68 ± 0.06	-3.7 ± 0.5
(kcal/mol)	0.25	$\textbf{-}1.78 \pm 0.06$	-3.9 ± 0.4
	0.50	$\textbf{-}1.90 \pm 0.07$	-4.1 ± 0.5

^a $K_{\text{Im,Amp}}$ is the apparent binding constant for Im derived from kinetic amplitude data. Error is the standard error reported by SigmaPlot for fits to eq S2 in Fig. S5.

The GdnHCl concentration dependence of the activation free energy can be obtained from plots of RTln(k) versus GdnHCl for the rate constants k_1 and k_{-2} obtained from fits of eq 2 to the data in Fig. 6 (Table S6). The slopes of these plots are small (Fig. 7B). The m-value for k_1 , m_{k_1} , which is a measure of the extent of the conformational change needed for Im binding is within error zero. The m-value for k_{-2} , $m_{k_{-2}}$, which is a measure of the extent of the conformational change needed to break the Im bond to the iron of the heme is similar in magnitude to the equilibrium m-value for Im binding, $m_{\rm Im}$ (Table 4).

^b Error is propagated from the error in $K_{\text{Im,Amp}}$.

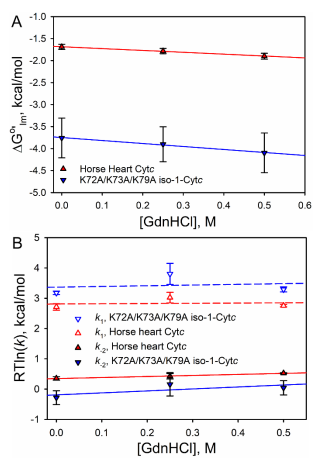


Fig. 7. Plots of (A) $\Delta G^{\circ}'_{Im}$ and (B) RTln(k) for k_1 and k_2 versus GdnHCl for horse heart Cytc and K72A/K73A/K79A iso-1-Cytc. Solid and dashed curves are fits of a linear free energy relationship to the data. The m-values from these fits are provided in Table 4.

Table 4 GdnHCl *m*-values for $\Delta G^{\circ\prime}_{Im}$, k_1 and k_{-2} for imidazole binding to Cytc at I = 1.25 M, pH 7 and 25 °C.

Protein	$m_{\rm Im}$ (kcal mol ⁻¹ M ⁻¹)	m_{k_1} (kcal mol ⁻¹ M ⁻¹)	$m_{k_{.2}}$ (kcal mol ⁻¹ M ⁻¹)
Horse heart Cytc	0.42 ± 0.02	0.08 ± 0.67	0.34 ± 0.03
K72A/K73A/K79A iso-1-Cyt <i>c</i>	0.68 ± 0.06	0.2 ± 1.3	0.65 ± 0.64

3.5 Structure of the N63T variant

Fig. 8 shows alignments of the X-ray structure of N63T iso-1-Cytc with those of WT iso-1-Cytc and human Cytc. The backbone RMSD values obtained from PyMol show that N63T when aligned with WT yeast iso-1-Cytc has a backbone RMSD of 0.405 Å, while it is 0.323 Å when aligned with WT human Cytc, indicating that the backbone of the N63T variant aligns slightly better with that of human Cytc than with yeast iso-1-Cytc. The all atom RMSD values show the same trend: the RMSD for the N63T variant aligned to yeast iso-1-Cytc is 1.626 Å, while it is 0.871 Å for N63T iso-1-Cytc aligned to WT human Cytc.

There are also changes in hydrogen bonding that result from the N63T mutation to iso-1-Cytc. For WT yeast iso-1-Cytc, there is a hydrogen bond from the carbonyl of the Asn63 side chain to the -OH group of Tyr74 and from the -NH₂ of the Asn63 side chain to the carbonyl of Leu58 (Fig. 8A). Thus, for WT iso-1-Cytc, the N-terminal end of Ω -loop D is anchored to the

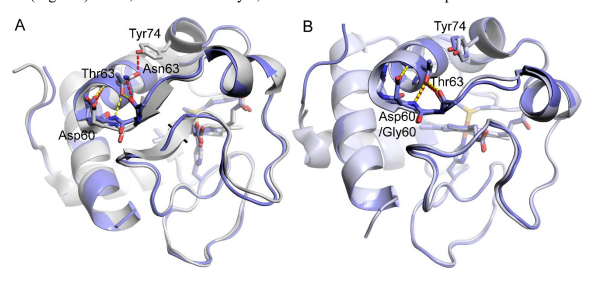


Fig. 8. Structure of N63T iso-1-Cytc (PDB ID: 7MRI) overlaid with (A) WT iso-1-Cytc (PDB ID: 2YCC) [65] and with (B) WT human Cytc (PDB ID: 3ZCF) [26]. In both panels hydrogen bonds for N63T iso-1-Cytc are shown as yellow dashed lines and for WT yeast and human proteins as red dashed lines. In both panels, the backbone atoms from Leu58 to Thr63 (Asn63) are shown as sticks.

short neck of β -sheet at either end of Ω -loop C. For N63T iso-1-Cytc, as for human Cytc, the side chain of Thr63 maintains a hydrogen bond to the carbonyl of Leu58. However, the hydrogen bond to the side chain -OH of Tyr 74 is lost (Fig. 8B), disrupting the connection between Ω -loop D and the β -sheet neck at the ends of Ω -loop C. The side chain of Thr63 in the N63T variant also forms a hydrogen bond to the backbone amide NH of Asp60, which is the N-cap residue of the 60s helix. The side chain of Asp60 caps the backbone amide NH of Thr63 (Fig. 8B). This reciprocal interaction constitutes a classic "capping box" motif, where the sidechains of the N3 residue of the helix (Thr63) and of the N-cap residue of the helix (Asp60) are each hydrogen bonded to the backbone amide NH of the other [66,67]. For WT human Cytc, the residue at position 60 is Gly, which is incapable of forming this "capping box" motif (Fig. 8B). WT and N63T yeast iso-1-Cytc form a hydrogen bond from the side chain of Asp60 to the amide NH of Asn62 also capping this amide NH. The hydrogen bond is stronger (2.95 Å) for the N63T variant than for WT iso-1-Cytc (3.29 Å). The N63T variant may have unique properties as a result of this "capping box" motif. However, all three proteins have the hydrophobic interaction between the 4^{th} residue of the helix (Met64 in yeast, Leu64 in human) and the N' residue of the helix cap (Trp59) typical of the "capping box" motif [66].

4. Discussion

4.1. Effects of coevolved mutations on local and global stability and the peroxidase activity of iso-1-Cytc.

There is considerable evidence that the effects of mutations on protein function depend upon sequence context [68]. Sites that coevolve in a protein, while distant in sequence are observed to

be proximal in the three-dimensional structure [68-70], as observed for the set of three residues used for mutational analysis in this work. The coevolved sites in or near Ω -loop C of Cytc were predicted using the EVmutation algorithm [25]. EVmutation is particularly effective at accounting for the effects of mutation on enzymatic activity, yielding Spearman correlation coefficients of 0.80 and 0.87 between mutation and function for the M.Hae III DNA methyltransferase and β -lactamase, respectively [25]. EVmutation also successfully predicted the effects of mutations on function in other proteins, such as poly(A)-binding protein [25]. EVmutation analyzes the sequence of a protein in organisms that have evolved earlier and compares it to the protein sequence in organisms that evolved more recently. Given the prominent role that the dynamics of Ω -loop C plays in the peroxidase activity of Cytc [10-13], it seemed likely that the three coevolved residues studied here at positions 40, 57 and 63 might explain the large difference in peroxidase activity between yeast and human Cytc.

As is evident from our data, the effects of single, double and triple mutant variants at the coevolved positions, 40/57 and 57/63, on global and local stability, in particular, are relatively small. With regard to local stability, while individual single or double mutation variants perturb the p K_a of the alkaline transition, there is no net effect of all three mutations on the local stability of Ω -loop D as measured by the p K_a of the alkaline transition (Table 2). Thus, evolutionary pressure appears to act to maintain the instability of the least stable Ω -loop C and Ω -loop D substructures of Cytc. In support of the notion that evolution has preserved the properties of Ω -loop C, the stability of this least stable substructure is similar in yeast iso-1-Cytc versus equine Cytc (Fig. 9) [31,71]. By contrast, the stability of the most stable substructure increases by more than 7 kcal/mol in equine Cytc compared to yeast iso-1-Cytc. The net result is a much smaller free energy range for the partial unfolding of the substructures of yeast iso-1-Cytc compared to

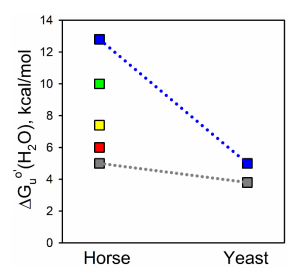


Fig. 9. Substructure energy levels for horse Cytc and yeast iso-1-Cytc. The foldons as determined for horse Cytc, using Englander's nomenclature, are from lowest to highest energy, the infrared (Ω -loop C, shown in gray), red (Ω -loop D), yellow (residues 37 – 39 and 58 – 60), green (residue 19 – 36 Ω -loop and 60s helix), and blue (N- and C-terminal helices) foldons [72]. Limited proteolysis was used to determine the energy of the least stable substructure of yeast iso-1-Cytc [31], which precludes measuring the energies of the other substructures. $\Delta G_u^{o'}(H_2O)$ for unfolding of iso-1-Cytc by GdnHCl was used for the stability of the most stable substructure because the *m*-value for global unfolding of the yeast protein is similar to that for the most stable structure of horse Cytc obtained from HX experiments.

equine Cytc [23,24,31,72]. The broad free energy range for the substructures of horse Cytc is presumably maintained in human Cytc, which is somewhat more stable than equine Cytc [73].

Denaturant *m*-values have been shown to correlate with the changes in the solvent-accessible surface area that occurs upon unfolding [74,75]. As a result, mutation-induced changes in stability have often been interpreted in terms of changes in the compactness of the denatured state [76,77], which in some cases result from a reverse hydrophobic effect [78,79]. However, mutation-induced changes in denaturant *m*-values for apparent two-state unfolding can also be caused by changes in the population of intermediates [80,81]. For horse Cytc, it has been shown that the large difference in substructure stabilities leads to low populations of intermediates during the GdnHCl unfolding transition. The result is a decreases in the cooperativity of the

apparent two-state transition as measured by the *m*-value of unfolding of horse Cyt*c* when monitored by CD or fluorescence relative to the true two-state *m*-value of 4.6 kcal/mol×M measured by HX methods [82]. Denaturant *m*-values monitored by standard optical methods yield lower *m*-values for GdnHCl unfolding of horse (3.0 kcal/mol×M) [83] and human Cyt*c* (3.5 – 3.7 kcal/mol×M) [15] relative to yeast iso-1-Cyt*c* (4.2 – 5.1 kcal/mol×M) [31,84,85]. The *m*-value of WT iso-1-Cyt*c* is closer to that of the *m*-value for true two-state unfolding of Cyt*c* measured by HX, consistent with the tight spacing of the substructure stabilities leading to GdnHCl unfolding that is more closely two-state (more cooperative, larger *m*-value). The decrease in the *m*-values for the substitution of the human residues at positions 40, 57 and 63 of yeast iso-1-Cyt*c* suggests that unfolding of these variants is less cooperative. The loss of the Asn63 to Tyr74 hydrogen bond in the X-ray structure of the N63T variant (see section 3.5) is consistent with the observed loss of cooperativity.

At pH 6 and above, the peroxidase activity of iso-1-Cytc is affected significantly by the substitutions at positions 40, 57 and 63. However, k_{cat} is not decreased to the level expected if the coevolved residues were solely responsible for the differences in intrinsic peroxidase activity between the yeast and human proteins. It is possible that pairwise coevolution revealed by the EVmutation algorithm [25] is insufficient to detect a broader set of coevolving residues necessary to decrease the intrinsic peroxidase reactivity to the level observed for human Cytc. In the highly conserved Ω -loop D, the I81A mutation in human Cytc [86], which substitutes the yeast residue alanine for the human residue isoleucine, had much larger effects on peroxidase activity than the A81I mutation in yeast iso-1-Cytc [27]. This observation indicates that a broader context of sequence differences between the yeast and human proteins are needed to change the dynamics of the two least stable substructures in a manner that limits the intrinsic

peroxidase activity of Cytc. Mutations in Ω -loop C, may have smaller effects on stability and function in the context of the yeast iso-1-Cytc sequence than they do in equine or human Cytc because of the smaller energy differences in the substructures of the yeast protein [31,71].

4.2. Correlation between Peroxidase Activity and Local and Global Stability

Plots of the change in the activation free energy for $k_{\rm cat}$, $\Delta\Delta G^{\ddagger}(k_{\rm cat})$, versus $\Delta G_{\rm u}^{\circ\prime}({\rm H_2O})$ from GdnHCl denaturation or the change in the free energy of the alkaline transition, $\Delta\Delta G^{\circ\prime}_{\rm alk}$, show very little correlation, except at pH 8 (Fig. 10) and for $\Delta G_{\rm u}^{\circ\prime}({\rm H_2O})$ at pH 7 (Fig. S9). A plot of $\Delta\Delta G^{\ddagger}(k_{\rm cat})$ at pH 8 versus $\Delta G_{\rm u}^{\circ\prime}({\rm H_2O})$ (Fig. 10A) that excludes WT iso-1-Cytc gives a correlation coefficient, R, of 0.96. This correlation suggests that this set of substitutions affects the properties of iso-1-Cytc so that they are distinct from WT iso-1-Cytc. The slope of the correlation is only 0.5 at pH 8 and even lower at pH 7 (~0.1) indicating that the change in $\Delta G_{\rm u}^{\circ\prime}({\rm H_2O})$ is not the only modulator of peroxidase activity.

A plot of $\Delta\Delta G^{\ddagger}(k_{cat})$ at pH 8 versus the change in the free energy of the alkaline transition, $\Delta\Delta G^{\circ}{}'_{alk}$, (Fig. 10B) that excludes WT iso-1-Cytc gives a correlation coefficient, R, of 0.57 (dotted line). A significantly better correlation coefficient (0.96) can be obtained if the S40T and N63T variants are not included in the correlation. This observation suggests that N63T and S40T variants may differ from the WT protein and from the other variants in some manner that affects peroxidase activity. The inverse correlation observed between $\Delta\Delta G^{\ddagger}(k_{cat})$ and $\Delta\Delta G_{alk}$ at pH 8 is also consistent with Lys-heme binding near the midpoint of the alkaline transition for these variants slowing formation of the open coordination site required for peroxide binding [34]. Recent studies showing that H₂O₂-induced carbonylation of lysines involved in the alkaline transition may be important for activating the peroxidase activity of Cytc [19] is also consistent

with this correlation. The observation that the correlation is not absolute indicates that other aspects of the dynamics of Ω -loops C and D are important in regulating the activation of peroxidase activity.

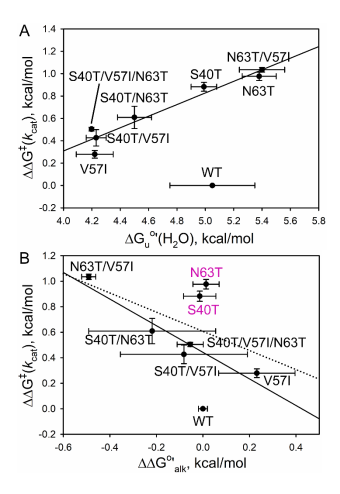


Fig. 10. (A) Plot of $\Delta\Delta G^{\ddagger}(k_{cat})$ at pH 8 versus $\Delta G_u^{o\prime}(H_2O)$ for all variants. The solid line is the correlation without WT iso-1-Cytc (R = 0.96). (B) Plot of $\Delta\Delta G^{\ddagger}(k_{cat})$ at pH 8 versus $\Delta\Delta G^{o\prime}_{alk}$. The dotted line is the correlation without WT iso-1-Cytc (R = 0.57). The solid correlation line (R = 0.96) also does not include the S40T and N63T variants. $\Delta\Delta G^{\ddagger}(k_{cat}) = -RTln(k_{cat,mut}/k_{cat,WT})$. The error bars are propagated from the errors in k_{cat} . $\Delta\Delta G^{o\prime}_{alk} = 2.3RT(pK_{a,mut} - pK_{a,WT})$. The error bars are propagated from the errors in the p K_a for the alkaline transition.

4.3. Relationship between structure and dynamics

Hydrogen-exchange (HX) data from previous work shows that human wild type Cytc [11] has larger protection factors at most residues than yeast iso-1-Cytc [87]. The protection factors, P, for Ω -loop D of yeast iso-1-Cytc measured at pD = 4.6 are generally low or unassigned, except for trimethyllysine 72, Lys73 and Tyr74 with LogP values of 3 – 4 and Thr78 and Phe82 which have LogP around 2 [87]. The protection factors for human Cytc in this same region measured at pD = 6.5 are also low, however residues Tyr74 and Ile75 have LogP values of almost 3, while residues Lys73, Lys79, Met80, and Ile85 have LogP values around 1 [11]. Although the measurements were carried out at different pD values, unlike Ω -loop C, the protection factors for Ω -loop D are not strongly dependent on pD [63]. Thus, the data suggest that the dynamics of Ω -loop D near Lys 73 and Lys 79 are similar for human Cytc, whereas for yeast iso-1-Cytc Ω -loop D is more dynamic near Lys79 than Lys73. Equilibrium data for the alkaline transition of horse Cytc indicate that Lys79 predominates over Lys73 in the alkaline state and that Lys72 does not contribute [54]. Studies comparing WT and K72A human Cytc also indicate that Lys72 is not a significant contributor to the alkaline state but are consistent with Lys73 rather than Lys79 being the predominant ligand for the kinetics of the alkaline transition. Equilibrium data for the alkaline transition of yeast iso-1-Cytc suggest that Lys73 is a somewhat stronger ligand for the alkaline state than Lys79, in the presence of trimetyllysine72 [88]. Lysines 72, 73 and 79 all contribute to the alkaline transition when Lys72 of iso-1-Cytc is not trimethylated [29] with mutational studies indicating that Lys72 has the lowest alkaline transition p K_a when only a single Lys is available in Ω -loop D [89]. However, kinetic data point to Lys79 as the predominate ligand for the kinetics of the alkaline transition of iso-1-Cytc [27]. The dynamics of Ω -loop D as measured by the alkaline transition and HX methods are often equated

[53,54]. The comparison is not straightforward if the GdnHCl-dependence of HX is not measured because HX can occur by local structural fluctuations rather than cooperative unfolding of a foldon like Ω -loop D. However, the hydrogen exchange data are qualitatively consistent with kinetic data for alkaline transition which indicate that for yeast iso-1-Cytc, Lys79 dominates the kinetics [27], while for human Cytc Lys73 is dominant in the kinetics of the alkaline transition [15].

For human Cytc (PDB: 3ZCF), the Thr63 side chain hydrogen bonds to main chain atoms of Gly60 and Ile58, whereas in yeast iso-1-Cytc (PDB: 2YCC), the Asn63 side chain hydrogen bonds to the side chain hydroxyl of Tyr74 and the main chain of Leu58. Because Tyr74 is in Ω -loop D, this residue could be a link between the two loops that modulates the dynamics that control opening of the heme crevice. The structure of N63T iso-1-Cytc shows that the hydrogen bond to the side -OH of Tyr74 is lost (Fig. 8) such that the side chain at position 63 no longer bridges Ω -loops C and D. If this hydrogen bond causes the larger protection factor at Lys73 compared to near Lys79 for iso-1-Cytc, one might expect that the N63T variant would have similar dynamics near both Lys73 and Lys 79, as indicated by the similar protection factors for these residues in human Cytc. Mutation-induced switching in the preferred ligand for the alkaline conformer may explain the poor correlation between the p K_a of the alkaline transition and peroxidase activity (Fig. 10 and Fig. S9). The formation of a "capping box" motif in the N63T iso-1-Cytc variant (Fig. 8) not present in WT iso-1-Cytc or human Cytc (Fig. 8) could further confound this correlation.

Our previous results for an I81A variant in Ω -loop D of human Cytc showed that the effect on the kinetics of the alkaline transition was selective for one of two detectable alkaline conformers [86]. Selective effects on the kinetics of one alkaline conformer were also observed

for the naturally-occurring A51V variant of human Cytc [9]. Based on the effects of A81I and G83V variants in Ω -loop D of iso-1-Cytc on the kinetics of the alkaline transition [27], we tentatively assigned the Lys79 alkaline conformer as the one that was preferentially affected by Ω -loop D substitutions. For iso-1-Cytc, a good correlation between the rate constant for forming the Lys79 alkaline conformer and the k_{cat} for peroxidase activity of the Ω -loop D variants was observed [27] suggesting that backbone dynamics near Lys79 might correlate with the dynamics needed for peroxidase activity. The decreased peroxidase activity for the Ω -loop C variants also may result from a redistribution of the relative backbone dynamics near Lys79 versus Lys 73. Recent MD simulations show a somewhat significant (R = 0.67) correlation between the flexibility of Ω -loop D and the p K_a of the alkaline transition of THC4 variants of human Cytc [14]. The fact that this correlation is imperfect may reflect the tendency of mutations in Ω -loops C and D to affect the dynamics of different parts of Ω -loop D.

4.4. Imidazole binding as a function of GdnHCl concentration

The effect of mutations on the thermodynamics of the alkaline transition correlate well with the thermodynamics of the cooperative unfolding of the Ω -loop D substructure of Cytc as measured by HX methods [53,54]. However, all substructures of Cytc have amides that exchange via lower energy local fluctuations [57]. Thus, it is possible that the dynamics required for peroxide binding to the heme go through lower energy pathways that do not require cooperative unfolding of the Ω -loop C and D substructures. In Scheme 1, formation of Cytc[†] is the limiting step in the reaction for Cytc with small molecules. We used imidazole in this work, but similar behavior is expected for the binding of hydrogen peroxide as it relates to the peroxidase activity of human Cytc. Plots of both $\Delta G^{o'}_{Im}$ and the activation free energy for

forming $\operatorname{Cyt}c^\dagger$ as a function of GdnHCl concentration (Fig. 7), yield m-values much less than observed for cooperative unfolding of the Ω -loop D substructure (Table 4). Thus, it can be concluded that Ω -loop D does not need to fully unfold for small molecules to bind to the heme. The GdnHCl dependence of the activation free energy for forming $\operatorname{Cyt}c^\dagger$ (Fig. 10B) is zero within error (Table 4) for both the horse and yeast proteins indicating that the conformational change to allow a small ligand to bind to the heme is minimal. These results are consistent with the small perturbation to the structure of horse $\operatorname{Cyt}c$ with imidazole bound [90], and the minimal perturbation to the structure of yeast iso-1- $\operatorname{Cyt}c$ when Met80 is displaced by hydroxide at moderately alkaline pH [16]. Thus, it is likely that mutation-induced effects on activation of the peroxidase activity of $\operatorname{Cyt}c$ will correlate better with local backbone fluctuations than with the pK_a of the alkaline transition or with global stability.

5. Conclusion

Decreases in the p K_a of the alkaline transition of Cytc often occur for mutations that increase the intrinsic peroxidase activity of Cytc. This observation has led to the assumption that cooperative unfolding of Ω -loop D linked to the dynamics of the adjacent Ω -loop C is needed to permit Cytc to act as a peroxidase. Substituting human residues into yeast iso-1-Cytc at three coevolved sites in or near Ω -loop C, we observed poor correlations between peroxidase activity and both the p K_a of the alkaline transition and global stability. The GdnHCl dependence of the kinetics of Im binding to horse and yeast cytochromes c confirmed that small local structural fluctuations of the backbone of Cytc are sufficient to allow small ligands like peroxide to bind to the heme. These results suggest that subtle mutation-induced changes in backbone dynamics may be sufficient to decrease the susceptibility of Cytc to pre-activation by reactive oxygen species so as to maintain the off state of the peroxidase signaling switch for apoptosis, particularly since the

stability of Ω -loop D appears to be robust to mutations at coevolved sites in and near Ω -loop C. In summary, previous work has focused on the unfolding of the entire Ω -loop D as being necessary for the binding of hydrogen peroxide and other small molecules, but our work indicates that the unfolding of the entire substructure is not necessary to activate the peroxidase activity of Cytc.

Abbreviations

βME β-mercaptoethanol

CD circular dichroism

CL cardiolipin

Cyt*c* cytochrome *c*

 $\Delta G^{\circ \prime}_{Im}$ Apparent free energy of binding of imidazole to Cytc

GdnHCl guanidine hydrochloride

HX hydrogen/deuterium exchange

Im imidazole

iso-1-Cytc yeast iso-1-cytochrome c

 $K_{\text{Im,equil}}$ apparent equilibrium constant for Im binding to Cytc

MD molecular dynamics

THC4 thrombocytopenia type IV

WT* variant of yeast iso-1-Cytc with a K72A substitution

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at

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