

NMR Investigations of the Rieske Protein from *Thermus thermophilus* Support a Coupled Proton and Electron Transfer Mechanism

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Abstract: The Rieske protein component of the cytochrome *bc* complex contains a [2Fe–2S] cluster ligated by two cysteines and two histidines. We report here the pK_a values of each of the imidazole rings of the two ligating histidines (His134 and His154) in the oxidized and reduced states of the Rieske protein from *Thermus thermophilus* (*TtRp*) as determined by NMR spectroscopy. Knowledge of these pK_a values is of critical interest because of their pertinence to the mechanism of electron and proton transfer in the bifurcated Q-cycle. Although we earlier had observed the pH dependence of a ^{15}N NMR signal from each of the two ligand histidines in oxidized *TtRp* (Lin, I. J.; Chen, Y.; Fee, J. A.; Song, J.; Westler, W. M.; Markley, J. L. *J. Am. Chem. Soc.* **2006**, *128*, 10672–10673), the strong paramagnetism of the [2Fe–2S] cluster prevented the assignment of these signals by conventional methods. Our approach here was to take advantage of the unique histidine–leucine (His134–Leu135) sequence and to use residue-selective labeling to establish a key sequence-specific assignment, which was then extended. Analysis of the pH dependence of assigned $^{13}\text{C}'$, $^{13}\text{C}^\alpha$, and $^{15}\text{N}^{\varepsilon 2}$ signals from the two histidine cluster ligands led to unambiguous assignment of the pK_a values of oxidized and reduced *TtRp*. The results showed that the pK_a of His134 changes from 9.1 in oxidized to ~12.3 in reduced *TtRp*, whereas the pK_a of His154 changes from 7.4 in oxidized to ~12.6 in reduced *TtRp*. This establishes His154, which is close to the quinone when the Rieske protein is in the cytochrome *b* site, as the residue experiencing the remarkable redox-dependent pK_a shift. Secondary structural analysis of oxidized and reduced *TtRp* based upon our extensive chemical shift assignments rules out a large conformational change between the oxidized and reduced states. Therefore, *TtRp* likely translocates between the cytochrome *b* and cytochrome *c* sites by passive diffusion. Our results are most consistent with a mechanism involving the coupled transfer of an electron and transfer of the proton across the hydrogen bond between the hydroquinone and His154 at the cytochrome *b* site.

Introduction

Thermus thermophilus Rieske protein (*TtRp*) is an indispensable subunit of the bacterial cytochrome *bc* complex, which has a counterpart in the mitochondrial cytochrome *bc*₁ complex (complex III) (EC 1.10.2.2).^{1–5} Complex III consists of three catalytic subunits (with cofactors): cytochrome *b* (two hemes), cytochrome *c*₍₁₎ (heme), and the Rieske iron–sulfur protein

([2Fe–2S] cluster). Complex III is a membrane-bound protein complex that serves as a hub to accept electrons from hydroquinone and to donate the electrons to cytochrome *c*₍₁₎. The electron transfer mechanism is known as the bifurcated Q-cycle.^{6–10} In the half-reaction that takes place in the bacterium, the Rieske protein first accepts an electron from a hydromenaquinone molecule, and then donates the electron to cytochrome *c*.^{9,10} The second electron is recycled to reduce the menaquinone in the cytosolic side. The rate-limiting step occurs in the first electron transfer to the Rieske protein.^{9–11} The net reaction is to oxidize the hydromenaquinone to menaquinone, reduce two cytochromes *c*, release two protons into the

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periplasmic space, and transfer two protons across the membrane, against the proton gradient.^{6,7}

Inhibitors of complex III, such as stigmatellin and myxothiazol, have been used to study the function of the complex.^{12–15} X-ray crystal structures show interactions among cytochrome *b*, stigmatellin, and the Rieske protein. The oxygen of stigmatellin (an inhibitor that mimics ubiquinone) is observed in the X-ray structure to be within hydrogen-bonding distance to the N^{ε2} atom of *His161*.^{12,13} (We use italics here to indicate the bovine numbering system and normal font to indicate the *T. thermophilus* numbering system; *His161* corresponds to His154 in *TtRp*.) Models have been proposed to explain the catalytic mechanism, under the assumption that the ternary complex represents an intermediate in the enzymatic turnover.^{1,9,10,16}

The redox potential of the Rieske protein is known to be coupled to the protonation states of the imidazole rings of the iron-ligated histidines.^{17–20} In the oxidized state, one histidine has a high p*K*_a value (~9.65) and the other has a value (~7.85) near physiological pH; in the reduced state, both have very high p*K*_a values (~12.5).¹⁷ Therefore, the elucidation of the detailed electron and proton transfer mechanism depends critically on assigning p*K*_a values to the individual iron-ligated histidines in the oxidized Rieske protein. If the p*K*_a of His154 is ~7.9, then it likely to be deprotonated before binding the hydroquinone, or before formation of the reaction complex, and then electron transfer can be coupled to proton transfer.^{9,16} If the p*K*_a His154 is ~9.7, then electron transfer would appear to be decoupled from proton transfer, and the hydroquinone deprotonation step likely occurs separately.^{1,2}

X-ray structures identify two primary binding sites on the *bc*₍₁₎ complex for the Rieske protein: one, named the *b*-site, is close to cytochrome *b*, and the other, named the *c*-site, is close to cytochrome *c*₍₁₎. The *b*-site is the hydroquinone oxidation site.

After the proton and electron transfer at the *b*-site, the protein has to move and donate the electron to cytochrome *c*₍₁₎ at the *c*-site.^{1,2} Disruption of the translocation eliminates the enzyme activity owing to large distance between cytochromes *b* and *c*₍₁₎.²¹ Two major models have been advanced to explain the transfer mechanism: (i) on the basis of multiple conformations of the Rieske protein observed in different X-ray structures, Brandt et al. suggested that the protein transfers between sites by large, redox state coupled, conformational changes;^{1,12,22,23} (ii) others have proposed that passive diffusion between sites is more likely.^{8,24}

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Materials and Methods

Chemicals, Bacterial Strains, and Vectors. [¹⁵N]NH₄Cl (99%), [13C]-D-glucose (99%), D₂O (99%), [¹⁵N]-leucine (98%), [13C,¹⁵N]-histidine (98%), and sodium 2,2-dimethyl-2-silapentane-5-sulfonate (DSS) were purchased from Cambridge Isotope Laboratories (Andover, MA). Unlabeled amino acids, BME vitamins, and corresponding antibiotics were purchased from Sigma Aldrich (St. Louis, MO). Isopropyl β-D-thiogalactoside (IPTG) was purchased from LabScientific (Livingston, NJ). Water used in all samples and buffers was purified with an Elga Maxima water purifier (Elga, Inc.; Topsfield, MA). *Escherichia coli* strain BL21(DE3)pLysS, Origami(DE3) (with leucine deficiency), Origami B (DE3)pLysS (with leucine deficiency), and λ-(DE3) lysogenization kit were purchased from Novagen (Madison, WI). Expression vector, pET17b carrying cDNA encoding *TtRp* (pET17b/*TtRp*), was provided by Dr. James A. Fee (Scripps Research Institute, CA). *E. coli* strain JM2071 (with leucine and histidine deficiencies) was purchased from the Coli Genetic Stock Center (Yale University, New Haven, CT).

Protein Expression and Labeling. [15N]-*TtRp* and [13C,¹⁵N]-*TtRp* was produced by BL21(DE3)pLysS cells containing pET17b/*TtRp* grown in M9 medium containing 1 g/L ¹⁵NH₄Cl as described previously³ but with modifications: 0.3 mM FeCl₃ and 1% (v/v) BME vitamins were added to the growth media; [13C]-glucose was added in two aliquots with 2 g/L added at the beginning and 2 g/L added when OD₆₀₀ reached 2.5 (immediately after IPTG induction). [¹⁵N]-Leu-*TtRp* was produced by Origami(DE3) cells containing pET17b/*TtRp* in a rich synthetic medium described previously,²⁵ with the modification that 50% of the amino acids, including the selectively labeled amino acids, was added initially, and the remaining 50% was added with the IPTG. The strain JM2071(DE3) was derived from JM2071 by using the standard protocol with a lysogenization kit (Novagen). [13C,¹⁵N]-His,[¹⁵N]-Leu-*TtRp* was produced by JM2071(DE3) cells containing pET17b/*TtRp* grown in a similar way to [15N],[NA]-Leu-*TtRp*, except that the histidine was enriched with ¹³C and ¹⁵N and the cells were grown initially in 2 L of LB medium to a high cell density (OD of ~6) before the transfer to the synthetic medium. The yield in this medium was ~100 mg/L culture. An auxotrophic strain of *E. coli* (Origami(DE3) or JM2071(DE3)) was required for leucine labeling. Details concerning other labeling patterns and the purification protocols used are presented in the Supporting Information.

NMR Sample Preparation. Amicon (Millipore, Billerica, MA) tubes were used for protein concentration and buffer exchange following protein purification. The NMR buffer was 20 mM phosphate, 20 mM Tris, and 20 mM borate, in 9% D₂O at pH 8.5; the solution contained ferricyanide as the oxidizing agent and DSS as the internal chemical shift standard. Sodium dithionite was added to samples used for pH titrations or conformational analysis. Shigemi (Shigemi, Inc., Allison Park, PA) susceptibility matched NMR tubes were used for NMR samples in the oxidized state. NMR samples in the reduced state were prepared in an anaerobic chamber and transferred to NMR tubes with a J. Young valve (Wilmad, Vineland, NJ). The pH meter was calibrated before taking readings. Readings were taken before and after the data collection, and the averaged pH value was reported. In addition to the common pH standards, the pH electrode in the anaerobic chamber was calibrated against freshly prepared, saturated calcium hydroxide, as a pH 12.50 standard, and 0.05 M, 0.1 M, 0.5 M, and 1.0 M solutions of sodium hydroxide, respectively, as pH 12.67, 12.94, 13.52, and 13.73 standards.

NMR Spectroscopy. The temperature of samples of oxidized *TtRp* used for ¹³C-detection experiments was controlled at 278 K, because this temperature provided the clearest separation of the

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A

T₄₆P**EKEPLKPGDILVYAQGGGEPKPIRLEELKPGDPFVLAYPMMDPKTKVVKSGEA**
KNTLLVARFDPEELAPEVAQH₁₂₀**AAEGVVAYS****AVCTH****134LGCIVSQF**₁₄₂**VADEEAA**
LCPCH₁₅₄**G****GVYDLRH**₁₆₂**GAQVIAGPPPRPVPQLPVRVEDGVLVAAGEFLGPVGV**
QA₂₀₁

B

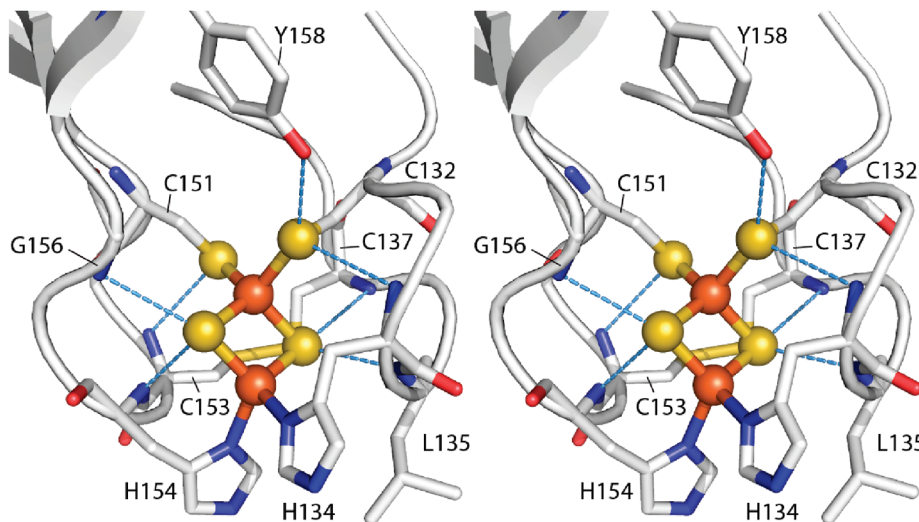


Figure 1. Sequence and structure of *TtRp*. (A) Primary sequence of the water-soluble fragment of the *T. thermophilus* Rieske protein (*TtRp*) investigated here. The construct was truncated to remove the N-terminal transmembrane region (residues 1–45) and nine C-terminal residues (202–210); in addition, it contains the point mutation W142F. The sequence contains four histidine residues: two of these (His134 and His154, shown in bold) each ligates an iron atom in the Fe–S cluster; the other two (His120 and His162) are not involved in Fe–S cluster interactions. Residues identified by X-ray crystallography as hydrogen bond donors to the iron sulfide cluster are underlined. Residues with NMR assignments in both oxidized and reduced forms of the protein are highlighted in green. (B) Cross-eyed stereoscopic view of the Fe–S cluster of *TtRp* and surrounding amino acid residues from the X-ray structure.³ Blue dashed lines indicate hydrogen bonds to the cluster.

histidine ¹³C' signals. The ¹⁵N-detected pH titration studies were carried out at both 278 K (to match conditions for ¹³C data collection) and 298 K (to match conditions used earlier).¹⁸ NMR data collection with all samples of reduced *TtRp* was carried out at 298 K. Technical details are presented in the Supporting Information.

Conventional triple resonance data, including ¹⁵N-HSQC, HNCACB, CBCACONH, and CCONH, were collected and used in determining spectral assignments.

NMR Data Processing and Analysis. Proton chemical shifts were referenced relative to internal DSS (taken as 0 ppm); carbon and nitrogen spectra were referenced indirectly by the canonical ratios.²⁶ All spectra were processed by XWIN-NMR software (Bruker) with exponential line broadening. COSY spectra were apodized by applying a Gaussian window function. Additional details are provided in Supporting Information. SPARKY software²⁷ was used for peak picking. Origin 7.0 software (OriginLab, Northampton, MA) was used to carry out nonlinear fitting of the titration data according to eq 1,²⁸

$$\delta_{\text{obs}} = \delta_B + [\Delta\delta \times 10^{n(pK_a - \text{pH})}] / [1 + 10^{n(pK_a - \text{pH})}] \quad (1)$$

in which δ_{obs} and δ_B represent the observed chemical shift of the signal at a certain pH and the signal at deprotonated state,

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respectively, $\Delta\delta$ represents the titration shift of the signal from the deprotonated state to the protonated state, and n is the Hill coefficient. Unless otherwise noted, all curve fitting was carried out with the Hill coefficient fixed at 1.0. The errors reported are those from fitting the experimental points.

Results and Analysis

Assignment Strategy. *TtRp* contains four histidine residues (Figure 1A): His120, His134, His154, and His162. Signals from the iron–sulfur cluster ligating histidines (His134 and His154, Figure 1B) were easily identified on the basis of their paramagnetic line broadening. It has long been known that backbone signals from large proteins can be assigned through the selective labeling of ¹³C'-¹⁵N_(i+1) linkages.²⁹ Upon examination of the dipeptide sequences involving the four histidines of *TtRp* (His120–Ala121, His134–Leu135, His154–Gly155, and His162–Gly163), it was clear that the ¹³C' of His134 could be identified by incorporating [¹⁵N]-Leu and [U-¹³C,U-¹⁵N]-His into *TtRp*. Once this carbon was identified, the plan was to extend this assignment through paramagnetic optimized single-bond correlations to as much of the His134 spin system as possible. Then by comparison of the pH dependence of the assigned signals with those from the ¹⁵N NMR titration curves for the ligated histidines, the specific assignment of the two pK_a values to His134 and His154 could be accomplished.

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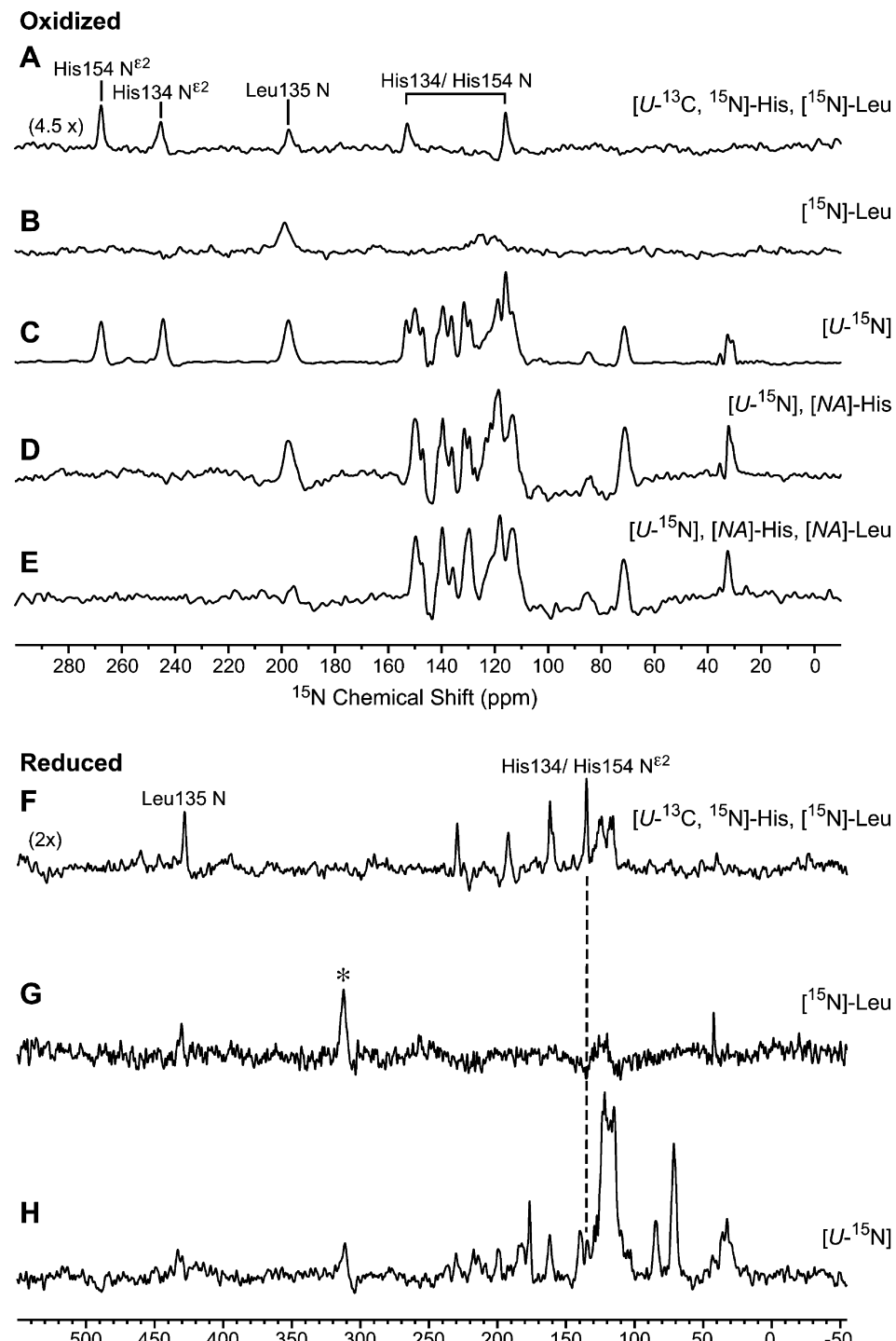


Figure 2. Nitrogen-15 NMR spectra of the variously labeled samples of *TtRp* used in assigning signals to iron–sulfur cluster ligands. One-dimensional ¹⁵N NMR spectra of ~10 mM oxidized *T. thermophilus* Rieske protein (*TtRp*) with different selective labeling and reverse selective labeling patterns at pH ~8.5 in the oxidized state (A–E) and pH 8.0 in the reduced state (F–H): (A) 17 mM [^{13}C , ^{15}N]-His, [^{15}N]-Leu *TtRp*; 86 $\times 10^3$ scans; (B) ~7 mM [^{15}N]-Leu *TtRp*; 1.8 $\times 10^6$ scans; (C) ~15 mM [^{15}N] ^{14}N *TtRp*; 1.9 $\times 10^6$ scans. (D) ~8 mM [^{15}N],[NA-Leu] *TtRp*; 2.1 $\times 10^6$ scans; (E) ~7 mM [^{15}N],[NA-Leu],[NA-His] *TtRp*; 4.2 $\times 10^6$ scans; (F) 14 mM [^{13}C , ^{15}N]-His, [^{15}N]-Leu *TtRp*; 1 $\times 10^6$ scans; (G) 9 mM [^{15}N]-Leu *TtRp*; 1.6 $\times 10^6$ scans; the asterisk denotes the $^{15}\text{N}^{14}\text{N}$ in the air; (H) 5.5 mM [^{15}N] ^{14}N *TtRp*; 2.1 $\times 10^6$ scans. Technical details were presented in Supporting Information.

Assignments. In order to carry out the strategy outlined above, it was necessary first to identify the ¹⁵N NMR signal from Leu135. Because of its participation in the hydrogen-bonding network at the iron–sulfur cluster³ (Figure 1B), we expected the ¹⁵N signal to be hyperfine-shifted. Our approach was to compare directly observed ¹⁵N NMR spectra, collected with short recycling delays as optimal for paramagnetic signals, of

TtRp samples containing the following labeling patterns: [^{13}C , ^{15}N]-His, [^{15}N]-Leu (Figure 2A,F); [^{15}N]-Leu (Figure 2B,G); [^{15}N] (Figure 2C,H); [^{15}N],[NA-His] (Figure 2D); and [^{15}N],[NA-His],[NA-Leu] (Figure 2E).

In spectra of selectively labeled oxidized *TtRp*, the ¹⁵N signal at 194.4 ppm, which was observed with all samples except for that with excess natural abundance leucine (Figure 2E) was

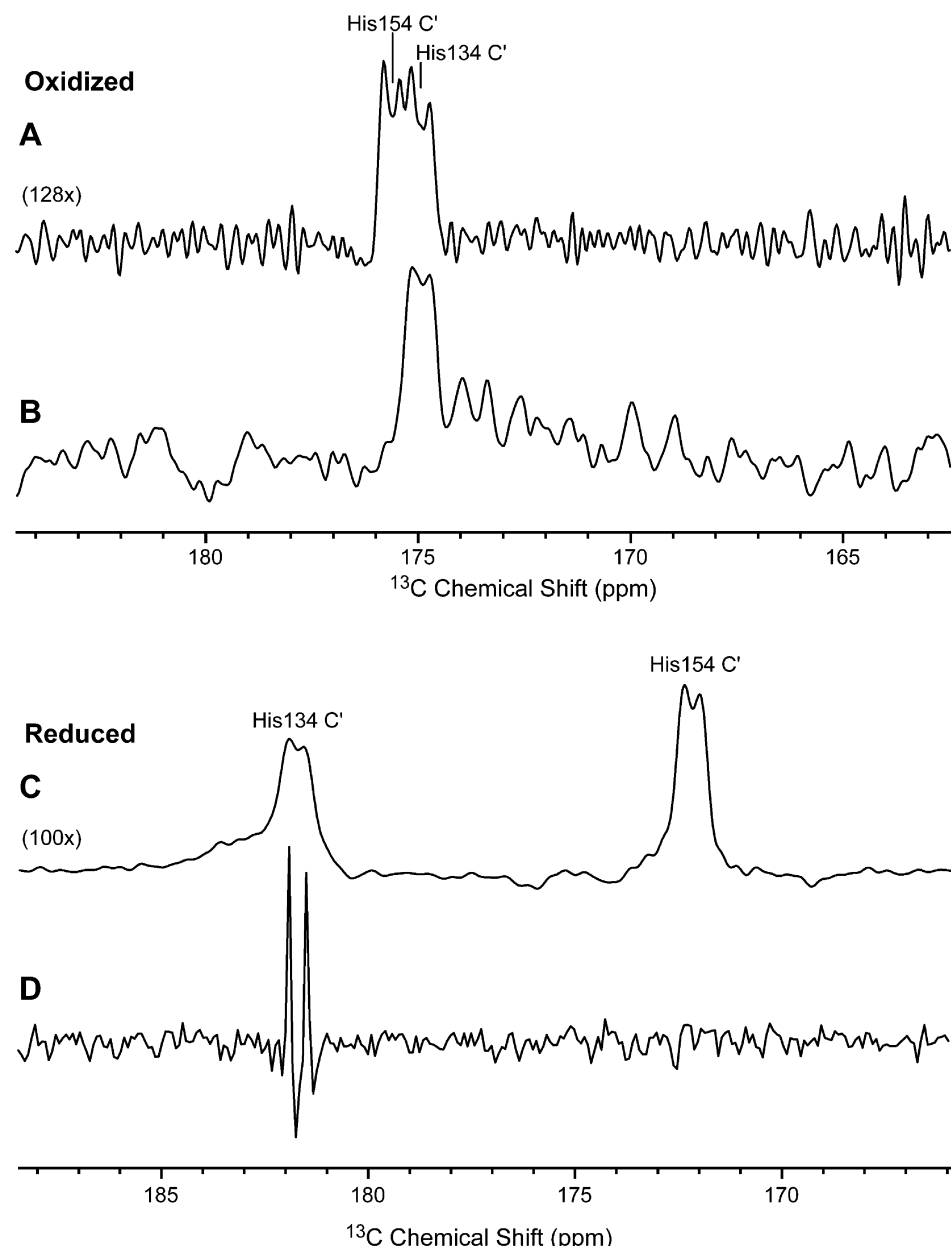


Figure 3. Carbon-13 [nitrogen-15] selective decoupling experiment leading to sequence-specific assignments: (A) 1D-¹³C-SW spectrum of 17 mM oxidized [U-¹³C, ¹⁵N]-His, [¹⁵N]-Leu TtRp acquired at 278 K and pH 8.5; locations of the two peaks were verified by a 2D ¹³C-[¹³C] CT-COSY-SW experiment; (B) double resonance ¹³C-[¹⁵N] difference decoupling experiment; the decoupler was set to be on-resonance at the ¹⁵N of Leu135 in the first spectrum and off-resonance in the second spectrum; the spectrum shown is the difference between the two spectra; (C) 1D-¹³C-SW spectrum of 14 mM reduced [U-¹³C, ¹⁵N]-His, [¹⁵N]-Leu TtRp acquired at 298 K and pH 7.9; (D) double resonance ¹³C-[¹⁵N] difference decoupling experiment of the reduced [U-¹³C, ¹⁵N]-His, [¹⁵N]-Leu TtRp. See the Supporting Information for technical details.

assigned tentatively to Leu135. The same set of spectra also served to confirm the identity of ¹⁵N signals from the backbone and side chain nitrogens of His134 and 154 (Figure 2A).

Spectra of selectively labeled reduced TtRp, by comparison with previous results,¹⁸ allowed the assignment of the signal at ~133 ppm (Figure 2F) to overlapped ¹⁵N^{ε2} peaks from His135 and His154. The signal at 423.6 ppm was tentatively assigned to Leu135 N (Figure 2G). The spectrum of uniformly ¹⁵N-labeled reduced TtRp (Figure 2H) was consistent with the assignment of these peaks.

The key step in assigning the iron-ligating histidines was a double resonance decoupling experiment with TtRp incorporating [U-¹³C, U-¹⁵N]-His, [¹⁵N]-Leu (Figure 3), which was used to identify the carbonyl carbon of His134. The directly observed

undecoupled ¹³C NMR spectrum of this sample in the oxidized state exhibited a pair of broad doublets at ~175.3 ppm (Figure 3A). When irradiated with ¹⁵N at 194.4 ppm (the suspected resonance frequency of Leu135), one of the doublets decreased in intensity as shown by the difference spectrum (off resonance – on resonance) (Figure 3B). This experiment identified the doublet at lower frequency as that from ¹³C' of His134. The other doublet was assigned by difference to ¹³C' of His154. The origin of each doublet was coupling to the adjacent ¹³C^a. Control experiments (Figure S1, Supporting Information) served to show that the effect observed was not an artifact.

We used the same approach to assign the ¹³C' peaks from His134 and His154 in reduced TtRp (Figure 3C). Selective irradiation of the ¹⁵N signal tentatively assigned to Leu135N,

Table 1. Nuclei Observed and Their Properties^a

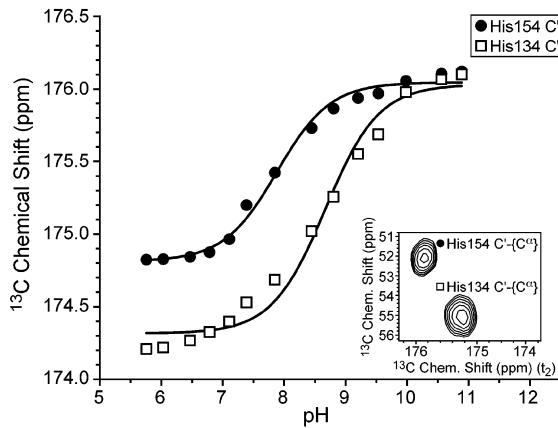
| atom | number of bonds (H-bonds) to the nearest iron atom | distance to the closest iron atom (Å) | distance to the His134 N ^{ε2} (Å) | distance to the His154 N ^{ε2} (Å) |
|-----------------------|--|---------------------------------------|--|--|
| His134 C' | 5 | 4.8 | 4.36 | 7.58 |
| His154 C' | 5 | 4.8 | 7.26 | 5.96 |
| His134 C ^α | 4 | 4.7 | 4.36 | 8.04 |
| His154 C ^α | 4 | 4.1 | 6.66 | 4.51 |
| Leu135 N | 6 (2) | 4.1 | 4.67 | 6.80 |

^a Distances are the average of those in the A and B models from the X-ray structure.³

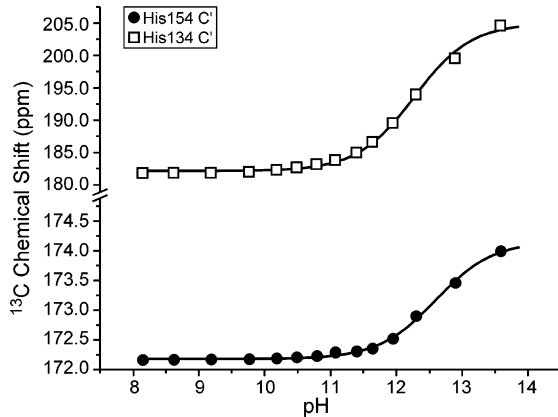
gave rise to a sharp doublet in the difference spectrum, which served to assign the higher frequency signal to His134C' (Figure 3D). Again, the other peak was assigned by difference to His154C' (Figure 3C).

The decoupling experiment confirmed the tentative Leu135¹⁵N assignments: 194.4 ppm in oxidized and 423.6 ppm in reduced state. In addition, by comparison of ¹³C NMR spectra taken under conditions optimized for diamagnetic signals with those optimized for paramagnetic signals, it was possible to clearly distinguish the ¹³C' signals of the diamagnetic histidines (His120 and His162) from those of His134 and 154 (Figure S2, Supporting Information).

A Oxidized



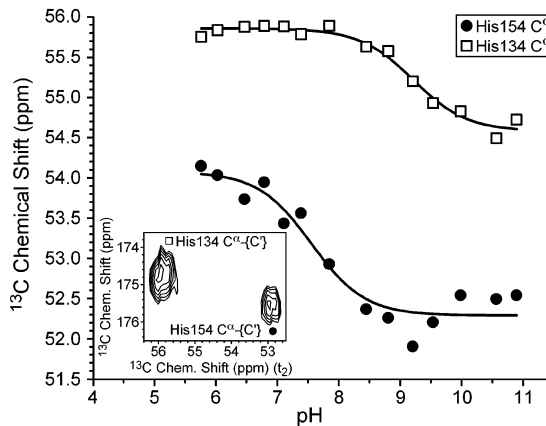
C Reduced



We extended the primary histidine ¹³C' assignments to the ¹³C^α atoms of His134 and His154 (Table 1) on the basis of connectivities observed in the constant-time 2D ¹³C-[¹³C] COSY-superWEFT³⁰ spectra of ([U-¹³C,U-¹⁵N]-His,[¹⁵N]-Leu)-TtRp (see insets in Figure 4A,B,D). However, attempts to extend these assignments to additional atoms in the histidine spin systems proved unsuccessful.

pH Titrations. We collected 2D ¹³C-[¹³C] COSY-superWEFT spectra of oxidized and reduced ([U-¹³C,U-¹⁵N]-His,[¹⁵N]-Leu)-TtRp at multiple pH values (Figure 4). The ¹³C chemical shifts used as input for titration analysis were measured from the directly detected dimension, with the exception of those for ¹³C^α in reduced TtRp, which were measured from the indirect dimension because of its higher signal-to-noise ratios. For comparison, directly observed 1D ¹⁵N NMR spectra of oxidized (Figure 5A) and reduced (Figure 5C) [¹⁵N-His] TtRp were collected as a function of pH. The histidine ¹⁵N^{ε2} signals from oxidized TtRp were used to generate titration curves (Figure 5B), but the corresponding signals from reduced TtRp disappeared in spectra of samples above pH 10.48, unlike the ¹³C' (Figure 4C) and ¹³C^α (Figure 4D) signals, which could be followed above pH 13.5. As a means of verifying that the ¹⁵N and ¹³C' signals were monitoring the same titration step, we

B Oxidized



D Reduced

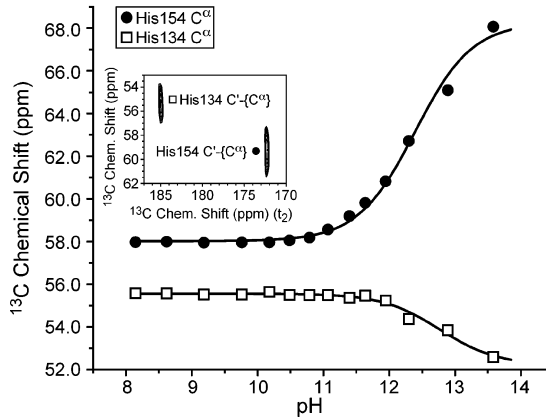


Figure 4. Carbon-13 titration curves and analysis: (A, B) data from 2D ¹³C-[¹³C] CT-COSY-SW spectra of oxidized 21 mM [U-¹³C,¹⁵N]-His,[¹⁵N]-Leu TtRp collected at 278 K at various pH values; (C, D) data from 2D ¹³C-[¹³C] CT-COSY-SW spectra of reduced ~5 mM [U-¹³C,¹⁵N]-His,[¹⁵N]-Leu TtRp collected at 298 K at various pH values. Open squares correspond to His134, and filled circles correspond to His154. (A) Titration data for the ¹³C' NMR signals. The inset shows the 2D spectrum at pH 8.81. Fitting of the data points (curves) yielded His134 pK_a = 8.68 ± 0.08 and His154 pK_a = 7.89 ± 0.07. (B) Titration data for the ¹³C^α NMR signals. The inset shows the 2D spectrum at pH 7.85. Fitting of the data points (curves) yielded His134 pK_a = 9.22 ± 0.09 and His154 pK_a = 7.56 ± 0.18. (C) Titration data for the ¹³C' NMR signals. Fitting of the data points yielded His134 pK_a = 12.27 ± 0.03 and His154 pK_a = 12.59 ± 0.03. (D) Titration data for the ¹³C^α NMR signals. Fitting of the data points (curves) yielded His134 pK_a = 12.39 ± 0.05 and His154 pK_a = 12.75 ± 0.09. The inset shows the 2D spectrum at pH 11.4. Fitting of the data points yielded His134 pK_a = 12.27 ± 0.03 and His154 pK_a = 12.59 ± 0.03.

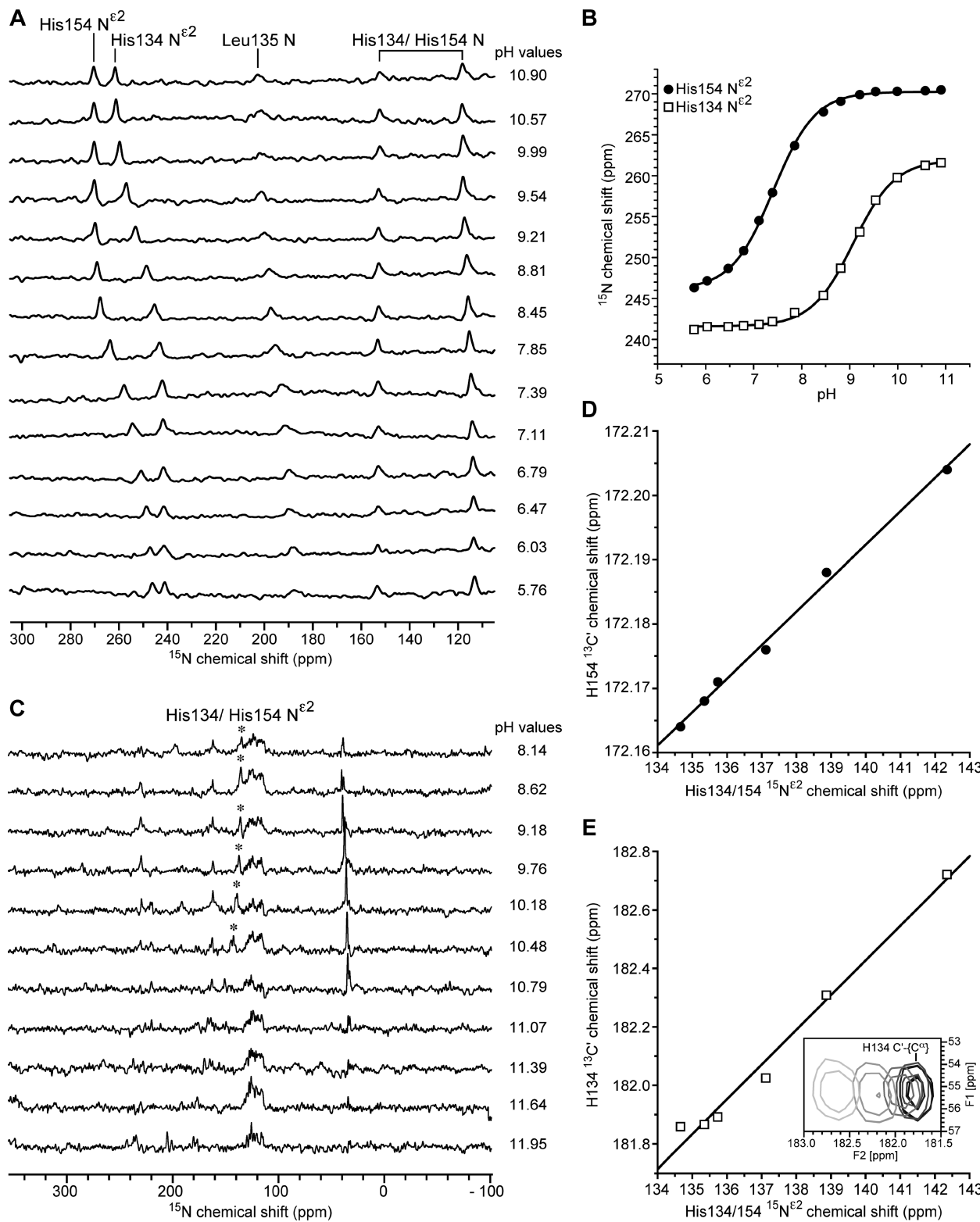


Figure 5. Nitrogen-15 titration curves and analysis. (A, B) Data from 1D ¹⁵N NMR spectra of 12 mM oxidized [¹³C, ¹⁵N]-His, ^{[15}N]-Leu *TtRp* collected at 278 K at various pH values. On the basis of data shown in Figures 4 and 5, the lower pK_a value was assigned to His154 and the higher pK_a value was assigned to His134 (Table 2). (C–E) Data from 1D ¹⁵N NMR spectra of ~5 mM reduced [¹³C, ¹⁵N]-His, ^{[15}N]-Leu *TtRp* collected at 298 K at various pH values. (A) Representative ¹⁵N NMR spectra of oxidized *TtRp*. Peak assignments are shown over the top trace. (B) The pH dependence of the chemical shifts of the peaks assigned to the ¹⁵N^{ε2} of His134 (open squares) and the ¹⁵N^{ε2} of His154 (filled circles) of oxidized *TtRp*. Fitting of the data points (curves) yielded His134 $pK_a = 9.16 \pm 0.02$ and His154 $pK_a = 7.58 \pm 0.01$. (C) Representative ¹⁵N NMR spectra of reduced *TtRp*. The ¹⁵N^{ε2} signals of His134 and His154 (denoted by *) overlapped between pH 8.14 and 10.48 and disappeared from spectra at higher pH. (D) Correlation between the His154 ¹³C' and ¹⁵N^{ε2} chemical shifts for the first six titration points. Linear regression yielded a correlation coefficient of 0.998. (E) Correlation between His134 C' and the ¹⁵N^{ε2} for the first six points. Linear regression line yielded a correlation coefficient of 0.992. The inset shows the 2D ¹³C'-¹³C^α cross peak assigned to His134 at different pH values (cross peaks move to the left with increasing pH) collected at the pH values corresponding to the top 6 spectra in panel C.

Table 2. Summary of Data Used to Assign the Cluster Ligated Histidine with the Higher pK_a Value to His134 and That with the Lower pK_a Value to His154

| nucleus observed | pK_a | figure number | experiment |
|-------------------------------|--------------|---------------|--|
| Oxidized <i>TtRp</i> at 278 K | | | |
| His134 C' | 8.68 ± 0.08 | 4A | ^{13}C -[^{13}C]-CT-COSY |
| His134 C α | 9.22 ± 0.09 | 4B | ^{13}C -[^{13}C]-CT-COSY |
| His154 C' | 7.89 ± 0.07 | 4A | ^{13}C -[^{13}C]-CT-COSY |
| His154 C α | 7.56 ± 0.18 | 4B | ^{13}C -[^{13}C]-CT-COSY |
| His134 N ϵ^2 | 9.16 ± 0.02 | 5B | 1D- ^{15}N rapid pulsing |
| His154 N ϵ^2 | 7.58 ± 0.01 | 5B | 1D- ^{15}N rapid pulsing |
| Oxidized <i>TtRp</i> at 298 K | | | |
| His134 N ϵ^2 | 9.07 ± 0.02 | S3 | 1D- ^{15}N rapid pulsing |
| His154 N ϵ^2 | 7.41 ± 0.01 | S3 | 1D- ^{15}N rapid pulsing |
| Reduced <i>TtRp</i> at 298 K | | | |
| His134 C' | 12.27 ± 0.03 | 4C | ^{13}C -[^{13}C]-CT-COSY |
| His134 C α | 12.39 ± 0.05 | 4D | ^{13}C -[^{13}C]-CT-COSY |
| His154 C' | 12.59 ± 0.03 | 4C | ^{13}C -[^{13}C]-CT-COSY |
| His154 C α | 12.75 ± 0.09 | 4D | ^{13}C -[^{13}C]-CT-COSY |

examined the linearity of their chemical shift relationship: the linear correlation coefficients were 0.998 and 0.992 for His154 and His134, respectively (Figure 5D,E). Because the carbon signals could be detected throughout the titration, were not overlapped, and were separately assigned, they provided the means for determining the pK_a values of the two ligand histidines in reduced *TtRp*. The results of the titration studies are summarized in Table 2.

Redox State Dependent Conformational Differences. Comparison of the ^1H - ^{15}N HSQC spectra of oxidized and reduced *TtRp* indicated chemical shift changes for numerous cross peaks. We established conditions at pH 5.2 under which *TtRp* was half-reduced and used a 2D NMR exchange experiment to cross assign signals corresponding to specific backbone ^1H - ^{15}N groups in the oxidized and reduced states.

Our next step was to carry out assignments for oxidized and reduced *TtRp* at pH 5.2. Conventional triple resonance NMR data sets were collected, including HNCACB, CBCACONH, ^{15}N -HSQC, and CCONH. We supplemented these experiments with ^1H - ^{15}N HSQC data from samples of *TtRp* labeled selectively with ^{15}N -Leu and ^{15}N -His. We used the PINE server³¹ to convert peak lists from these experiments, along with the protein sequence, into probabilistic assignments. The resulting assignments were refined by manual examination.

We succeeded in assigning signals to 122 of the 156 residues of *TtRp* in both oxidation states. Apart from the 18 residues belonging to the two loops (131–140 and 150–157) that surround the Fe–S cluster, which we did not expect to be able to assign, the assignments account for 87% of the residues of *TtRp*.

We made use of the $^{13}\text{C}^\alpha$ and $^{13}\text{C}^\beta$ assignments to compare the secondary structure of the oxidized and reduced forms of *TtRp* according to eq 2:^{32,33}

$$\Delta\Delta\delta_i = (\delta_{\text{C}\alpha,i} - \delta_{\text{C}\alpha,\text{rc},i}) - (\delta_{\text{C}\beta,i} - \delta_{\text{C}\beta,\text{rc},i}) \quad (2)$$

where $\delta_{\text{C}\alpha,i}$ is the $^{13}\text{C}^\alpha$ chemical shift of the i th residue of *TtRp*, $\delta_{\text{C}\beta,i}$ is the $^{13}\text{C}^\beta$ chemical shift of the i th residue of *TtRp*, $\delta_{\text{C}\alpha,\text{rc},i}$ is the random coil $^{13}\text{C}^\alpha$ chemical shift for residue i , and $\delta_{\text{C}\beta,\text{rc},i}$ is the random coil $^{13}\text{C}^\beta$ chemical shift for residue i downloaded

from BMRB Web site.³⁴ Positive values of $\Delta\Delta\delta_i$ indicate α -helices, and negative values indicate β -strands.^{32,33}

The results of this analysis (Figure 6A) showed no evidence for any large difference in the secondary structures of oxidized and reduced *TtRp*. We also examined the differences ($\Delta\Delta\delta_{i,\text{red}} - \Delta\Delta\delta_{i,\text{oxid}}$) (Figure 6B) and found, with few exceptions, that they were small. Other than the glycine loop (Gly63–Gly65) and the residues near or between the iron–sulfur binding loop, none of the assigned carbons exhibited redox state dependent ^{13}C NMR chemical shift differences greater than 0.5 ppm (data not shown). It appears that oxidized and reduced *TtRp* have very similar conformations, a result in agreement with the passive diffusion model for the translocation between *b*-site and *c*-site.²⁴

Discussion

NMR Assignments. In an earlier study,¹⁸ we incorporated selectively labeled histidine into the *TtRp* protein to distinguish between signals from the $\text{N}^{\delta 1}$ and $\text{N}^{\epsilon 2}$ atoms of the imidazole rings. Although signals from the $\text{N}^{\delta 1}$ atoms of the two imidazole rings that ligate iron were unobservable, because of the strong paramagnetism, signals from the $\text{N}^{\epsilon 2}$ atoms could be observed and their chemical shifts followed as a function of pH to yield distinct pK_a values for the two histidine residues in the oxidized state. We speculated on the basis of solvent-accessibility arguments that the lower pK_a value corresponded to His134 and outlined the mechanistic consequences of this assignment. Because of its importance, we undertook here a rigorous assignment of these signals. These assignments required the preparation of protein samples labeled selectively with ^{15}N and ^{13}C (Figures 2 and 3). Contrary to our earlier speculation,¹⁸ the results proved that His154 has the lower pK_a value. Our current NMR assignments in oxidized and reduced *TtRp* include the $^{15}\text{N}^{\epsilon 2}$, $^{13}\text{C}'$, and $^{13}\text{C}^\alpha$ atoms of each ligand histidine. These extended assignments allowed us to determine pK_a values for the ligand histidines in reduced *TtRp*. We used conventional methods with protein labeled uniformly with ^{13}C and ^{15}N to carry out extensive assignments of NMR signals in both oxidized and reduced *TtRp*. Several of these assignments were verified by selective ^{15}N labeling. The NMR assignments have been deposited at the BMRB under accession numbers 16787 (reduced) and 16804 (oxidized).

The similarity of the diamagnetic chemical shifts of oxidized and reduced *TtRp* rule out a large conformational change of the type observed by X-ray crystallography of cytochrome *bc*₁ complexes in the presence and absence of various inhibitors.¹²

pH Titration Studies. Our assignments required that the oxidized protein be studied at 278 K, in order to achieve separation of the $^{13}\text{C}'$ signals of the ligand histidines. Because the earlier pH titration study was carried out at 298 K, we repeated the ^{15}N titration study at this temperature (Figure S3, Supporting Information).

The shapes of the ^{15}N titration curves reported here differ slightly from those reported by Lin et al.¹⁸ The midpoint broadening of the peak yielding the lower pK_a observed earlier was not reproduced, and the splitting observed at high pH (attributed to interaction with titrating Tyr158)¹⁸ occurred at higher pH (11.0 instead of 10.5). These differences probably

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(34) <http://www.bmrb.wisc.edu/>.

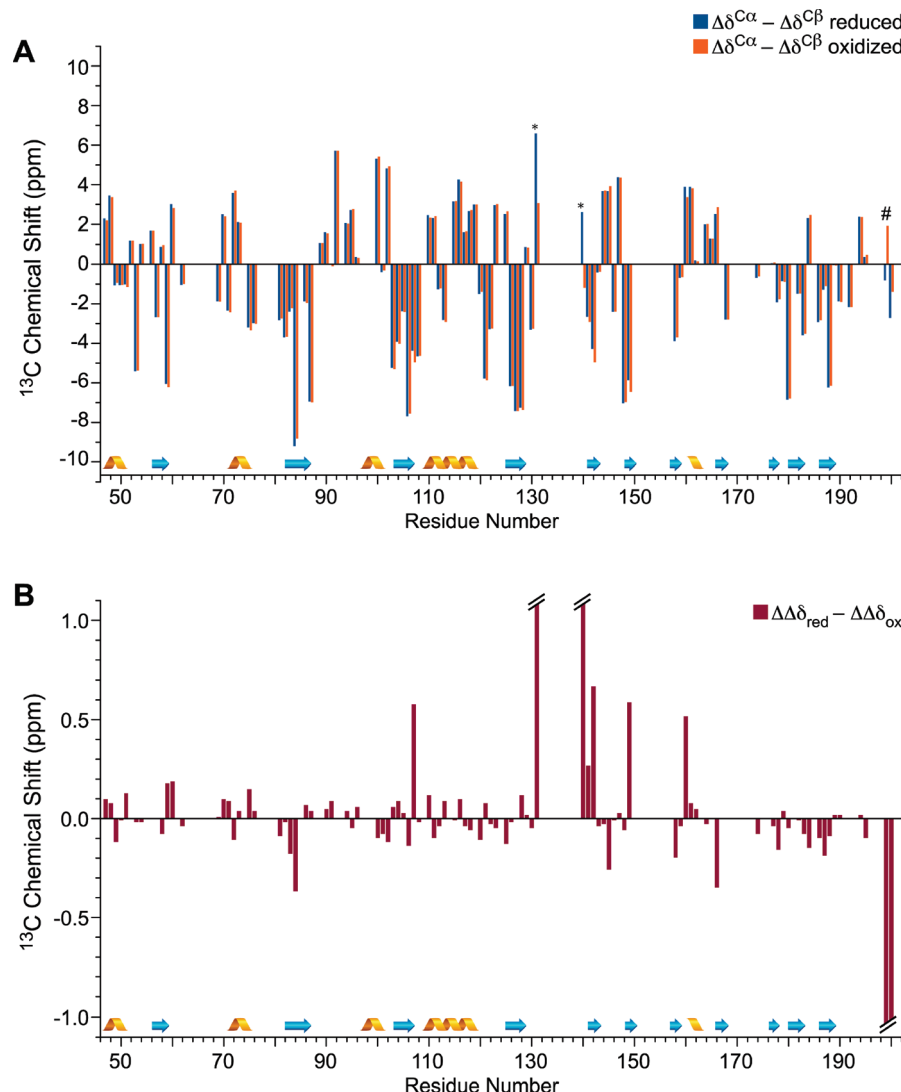


Figure 6. Secondary structure analysis of oxidized and reduced *TtRp*. Chemical shift data from 8.5 mM [$U^{-13}\text{C}, ^{15}\text{N}$] *TtRp* at pH 5.2, 298 K, in both oxidized (orange) and reduced (blue) states are plotted as a function of residue number. Secondary structural elements derived from the X-ray structure³ are (represented at the bottom of the figure) are in agreement with the chemical shift results. (A) For each residue of *TtRp*, the bar represents the difference between the secondary $^{13}\text{C}^\alpha$ chemical shift (experimental shift minus random coil shift, $\Delta\delta_{^{13}\text{C}^\alpha}$) and the secondary $^{13}\text{C}^\beta$ chemical shift (experimental shift minus random coil shift, $\Delta\delta_{^{13}\text{C}^\beta}$). The symbol * denotes residues that are close to the iron–sulfur binding site; # denotes the carboxyl terminus. (B) For each residue of *TtRp*, the bar represents the secondary shift difference ($\Delta\delta_{^{13}\text{C}^\alpha} - \Delta\delta_{^{13}\text{C}^\beta}$) in the reduced protein minus that in the oxidized protein. Large values corresponding to residues experiencing hyperfine shifts are truncated in the figure.

are a consequence of the presence of ferricyanide, used here but not by Lin et al., which appears to catalyze the protonation–deprotonation steps so that they become fast on the NMR time scale. The pK_a values determined here from the ^{15}N data at 298 K (7.41 ± 0.01 and 9.07 ± 0.02 , Figure S3, Supporting Information) are similar to those reported by Lin et al. (7.46 ± 0.02 and 9.24 ± 0.02) at the same temperature.¹⁸

The pK_a values derived from the pH dependence of NMR signals from different histidine atoms exhibited some variation (Table 2). Values derived from the chemical shifts of the imidazole atom (histidine $^{15}\text{N}^{\epsilon 2}$) are most likely to reflect the true pK_a for the residue. The pK_a values derived from the $^{13}\text{C}^\alpha$ chemical shifts in 2D spectra were in good agreement with these; however, the pK_a values derived from the chemical shifts of the $^{13}\text{C}'$ atoms, which are farther from the site of protonation–deprotonation, showed minor deviations (Table 2). In all cases, the higher pK_a corresponded to His134 and the lower pK_a to His154. The chemical shifts of the backbone nitrogens of His134, Leu135, and His154 in oxidized *TtRp* also were found

to be pH dependent (Figure S4, Supporting Information), but these shallow titration curves were not analyzed further.

The $\text{N}^{\epsilon 2}$ signals from His134 and His154 of reduced *TtRp* overlapped over the pH range where they were visible and disappeared from spectra of samples with pH > 10.5 (Figure 5C). However, the titration could be followed from the chemical shifts of the peaks assigned to the $^{13}\text{C}'$ and $^{13}\text{C}^\alpha$ atoms of His134 and His154 of reduced *TtRp* (Figure 4C,D). The fitted curves for the $^{13}\text{C}'$ signals, which exhibited larger (and hence more reliable) titration shifts than the $^{13}\text{C}^\alpha$ signals (which showed opposite pH dependence), yielded pK_a values of 12.3 ± 0.3 for His134 and 12.6 ± 0.3 for His154. These values are similar to those derived earlier from electrochemical studies: pK_a values ~ 12.5 .¹⁷

His154 exhibited a remarkably large redox state dependent shift in its pK_a , from 7.6 ± 0.2 in oxidized to 12.6 ± 0.3 in reduced *TtRp*; this represents a 10^5 increase in the proton affinity of the reduced protein over the oxidized. Although the shift in the pK_a of His134 is smaller, 9.2 ± 0.1 in oxidized vs $12.3 \pm$

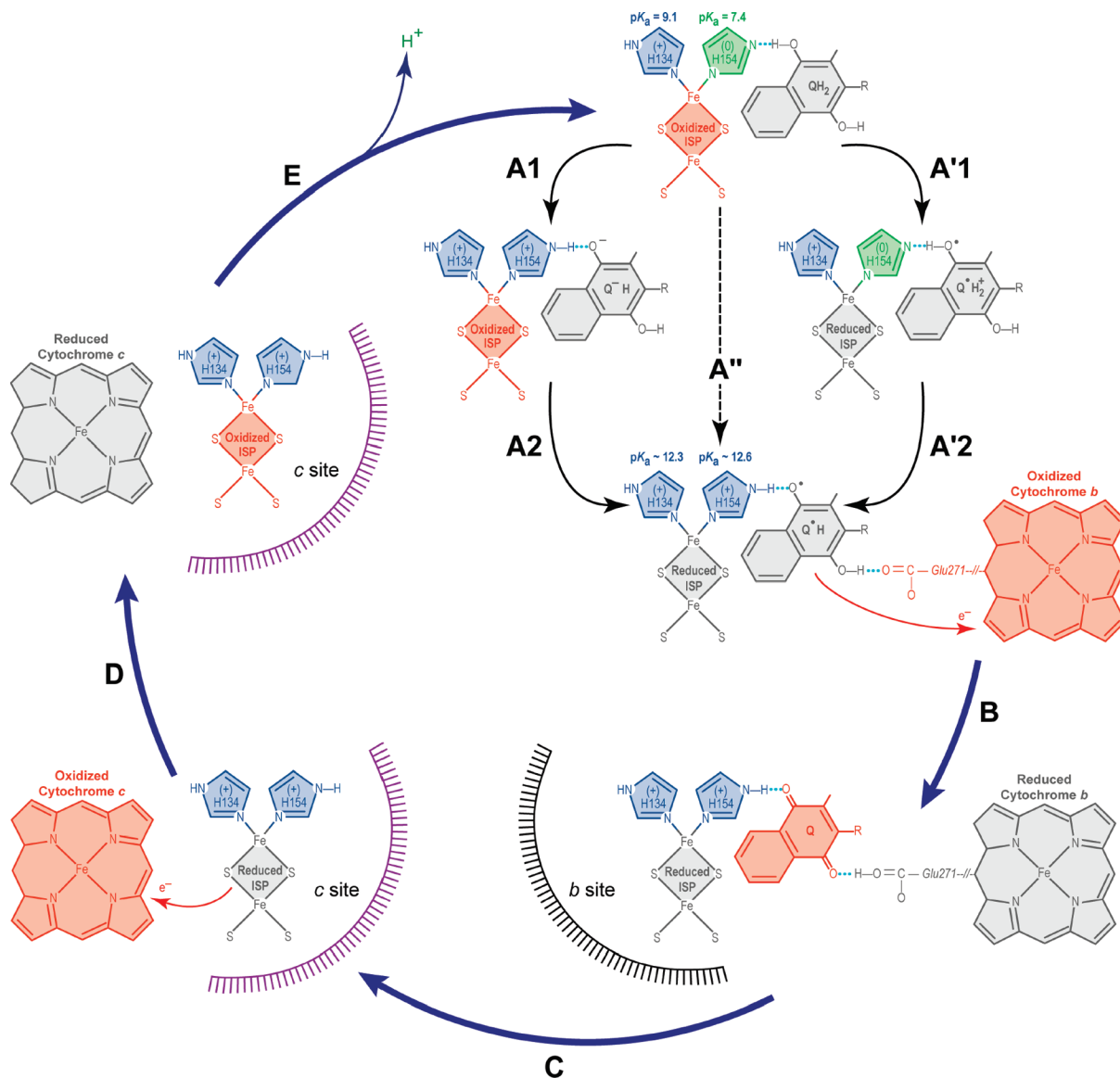


Figure 7. Model for proton and electron transfer at the Q_o site on the cytochrome bc complex. Pentagons represent the imidazole rings of the histidines (green deprotonated, blue protonated). Oxidized species are depicted in red and reduced species in gray. See Discussion for details. (A) The hydro(mena)quinone hydrogen bonds to the deprotonated His154 of the oxidized ISP ($TtRp$, iron–sulfur protein) (top); then the electron and the proton are transferred by one of three possible pathways: proton transfer first/electron transfer second (PT/ET), $A_1 \rightarrow A_2$; electron transfer first/proton transfer second (ET/PT), $A'_1 \rightarrow A'_2$; or coupled electron and proton transfer (ET+PT), A'' . In each case, the intermediate formed at the end is the semi(mena)quinone radical. (B) The semiquinone radical donates the electron to oxidized cytochrome b . (C) ISP translocates between the b -site and c -site to donate an electron to oxidized cytochrome c . (D) ISP donates an electron to cytochrome c . (E) ISP translocates from site- c to site- b , and only the deprotonated ISP is capable of forming the reaction complex.

0.3 in reduced $TtRp$, the increase represents a $>10^3$ increase in its proton affinity. Because of its high pK_a in both oxidized and reduced $TtRp$, His134 probably remains protonated throughout the Q-cycle. The NMR data, which yield separate titration curves with Hill coefficients near unity, rule out cooperativity between protonation–deprotonation at His134 and His154. Because no other proton binding sites are within ~ 9 Å of the iron–sulfur cluster, His154 is likely to be the terminal proton carrier.

Model for Proton and Electron Transfer. The model shown in Figure 7 is based on the pK_a values determined here, structural information about the b -site,³⁵ and three assumptions concerning the classical semi(mena)quinone model. The assumptions are that reactions are *in vitro* at 298 K, that the menaquinone and ubiquinone species have the same redox potential differences

as their Q/QH_2 state, such that menaquinone species are all 170 mV lower than the ubiquinone species, and that pK_a values of these two quinone species are the same.

The proton-first/electron-second (PT/ET) pathway involves steps A_1 (21.6 kJ mol⁻¹; $\Delta pK_a = 3.85$; Figure 7) and A_2 (-9.5 kJ mol⁻¹; $\Delta E = 0.1$ V). The electron-first/proton-second (ET/PT) pathway involves steps A'_1 (82.8 kJ mol⁻¹; $\Delta E = -0.86$ V) and A'_2 (-70.7 kJ mol⁻¹, $\Delta pK_a = -12.6$). The direct mechanism A'' involving simultaneous electron and proton transfer (ET+PT) has a free energy of 12.1 kJ mol⁻¹. Both initial intermediates in the stepwise pathway (A_1 and A'_1) are energetically unstable compared with the reactants, and both represent larger energy increases than the concerted reaction A'' (9.5 kJ mol⁻¹ for A_1 ; 70.7 kJ mol⁻¹ for A'_1). In addition, as we show here, adding an electron and a proton to $TtRp$ does not involve a large conformational change. Thus, the most likely

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mechanism is the energetically favored concerted reaction A'',^{17,36} in which electron transfer is coupled to proton transfer across the hydrogen bond between hydro(mena)quinone and His154 of the iron–sulfur protein. Theoretical arguments based on the Born–Oppenheimer approximation³⁷ and the Hohenberg–Kohn theorem³⁸ require concerted movement of the proton and electron. Our experimental results interpreted as shown above are inconsistent with models proposed by Link and Brandt^{39,40} that involve, as the first step, the deprotonation of hydro(ubi)quinone with a pK_a of >11.3 in solution. The resulting high-energy semiquinone is rapidly depleted by electron transfer reaction B, thus removing the reactive semiquinone, which could contribute to oxidative stress. This, in turn, stabilizes the reduced, protonated Rieske protein from returning to the oxidized state so that it can undergo the much slower translocation from site-*b* to site-*c* (reaction C, Figure 7).

In the biological environment, however, the binding of the substrates cannot be neglected (hydromenaquinone and the oxidized, deprotonated *TtRp* bind to the enzyme before the reaction). In *Rhodobacter sphaeroides*, the binding affects the rate of the turnover and modulates the pK_a of the Rieske protein and the redox potential of the hydro(ubi)quinone.¹⁶ The change is ~0.5–1 pH unit and ~50 mV.⁴¹ First, menaquinone binding appears to slightly lower the energy gap of the A'1 step (although it remains highly unfavored). Second, menaquinone binding makes the A1 step less favored as discussed elsewhere.¹⁶

The Proton Pathway. From the pK_a assignments, it is clear that the Rieske protein is the primary proton carrier. The complete proton transfer pathway against the proton gradient involves the following steps (Figure 7). First, menaquinone accepts two electrons and two protons in the cytosolic side (this is another half-reaction to the Q-cycle in the menaquinone-reducing-site). Second, by flip–flop to the periplasmic space, it binds and forms the reaction complex there (reactant in Figure 7A). Third, hydro(mena)quinone donates one electron and one proton to *TtRp* (Figure 7A). *TtRp* carries the electron and proton until it donates the electron to the cytochrome *c* at the *c*-site and releases the proton because of the large decrease in the pK_a of His154. Finally the *b*-site complex reforms.

Summary. Although the general features of the bifurcated Q-cycle mechanism and the proton motive gradient have been accepted, the detailed mechanism has remained controversial.^{42,43} Even after the confirmation of the important intermediate,

ubisemiquinone,⁴⁴ debates still continued over the detailed mechanisms. The primary evidence for the intermediate complex involving cytochrome *b*, ubihydroquinone, and the Rieske protein has come from the structure of the quinonoid inhibitor complex. Prior to this study, there was no direct evidence for the protonation state of the key histidine, His154 (corresponding to the bovine *His161*), of the Rieske protein. Nevertheless, models by Link, Berry, Rich, Trumper, and Crofts, based on the X-ray structures, kinetics, and thermodynamics studies,^{16,39,42,43,45} agreed that the catalytic mechanism involves the formation of two hydrogen bonds as shown in Figure 7 (see the product of step A).

This study shows that His154 of oxidized *TtRp* is at least partially deprotonated at physiological pH. Hydrogen bonding to hydroquinone should help to stabilize the deprotonated state. The large increase in the pK_a of His154 upon reduction confirms that it is the proton carrier between the *b*- and *c*-sites. Evidence also is presented here that reduction does not involve a large protein conformational change. The large conformational change in the Rieske protein observed by X-ray crystallography¹² apparently occurs as part of the subsequent translocation to the *c*-site. These results favor the direct electron/proton transfer mechanism as the first step of the oxidation of the hydro(mena)quinone. Owing to the high degree of conservation of residues in the cluster binding domains, our results with the bacterial Rieske protein (*TtRp*) probably will hold for Rieske proteins of the respiratory *bc*₁ complexes in the mitochondria of other organisms and probably also for the Rieske proteins in the photosynthetic *b*₆*f* complexes.

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Supporting Information Available: NMR related technical details (Materials and Methods, Figure 2, and Figure 3), data from controls related to the double resonance decoupling experiment (Figure S1), 1D ¹³C-SW spectra of oxidized [U-¹³C, ¹⁵N]-His, [¹⁵N]-Leu *TtRp* collected under conditions optimized for paramagnetic and diamagnetic signals (Figure S2), ¹⁵N pH titration curves at 298 K (Figure S3), and the pH dependence of the chemical shifts of the backbone nitrogens of His134, Leu135, and His154 (Figure S4). This material is available free of charge via the Internet at <http://pubs.acs.org>.

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