NMR investigation of ferricytochrome c unfolding: Detection of an equilibrium unfolding intermediate and residual structure in the denatured state

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Communicated by Harry B. Gray, California Institute of Technology, Pasadena, CA, May 24, 2000 (received for review February 23, 2000)

Horse ferricytochrome c (cyt c) undergoes exchange of one of its axial heme ligands (Met-80) for one or more non-native ligands under denaturing conditions. We have used $^1$H NMR spectroscopy to detect two conformations of paramagnetic cyt c with non-native heme ligation through a range of urea concentrations. One non-native form is an equilibrium unfolding intermediate observed under partially denaturing conditions and is attributed to replacement of Met-80 with one or more Lys side chains. The second non-native form, in which the native Met ligand is replaced by a His, is observed under strongly denaturing conditions. Thermodynamic analysis of these data indicates a relatively small $\Delta G (17 \text{ kJ/mol})$ for the transition from native to the Lys-ligated intermediate and a significantly larger $\Delta G (47 \text{ kJ/mol})$ for the transition from native to the His-ligated species. Although CD and fluorescence data indicate that the equilibrium unfolding of cyt c is a two-state process, these NMR results implicate an intermediate with His-Lys ligation.

The folding and unfolding of ferricytochrome c (cyt c) is strongly influenced by the binding of protein-donated ligands to its covalently attached heme. Denaturation of cyt c by concentrated guanidine hydrochloride or urea at neutral pH leads to replacement of the native Met heme axial ligand with a His, is observed under strongly denaturing conditions. Thermodynamic analysis of these data indicates a relatively small $\Delta G (17 \text{ kJ/mol})$ for the transition from native to the Lys-ligated intermediate and a significantly larger $\Delta G (47 \text{ kJ/mol})$ for the transition from native to the His-ligated species. Although CD and fluorescence data indicate that the equilibrium unfolding of cyt c is a two-state process, these NMR results implicate an intermediate with His-Lys ligation.

Materials and Methods
Preparation of Cyt c and Urea Solutions. Horse heart cyt c was purchased from Sigma, oxidized with KCo(EDTA) by a literature method (20), and purified by FPLC using a Mono S column (Amersham Pharamacia). Cyt c concentration was determined by using an extinction coefficient of 106,100 liters/mol-cm at 410 nm (21). Solutions of urea and urea-d$_4$ (highest quality available, Aldrich) were prepared in 50 mM sodium phosphate in H$_2$O (with 10% D$_2$O added for lock) or D$_2$O, stored at 4°C, and used within 12 h. The uncorrected pH (pH*) of urea solutions was adjusted to 7.0, and the concentration of denaturant was determined by refractive index measurement (22). Urea and protein solutions were combined to give the desired final concentrations, followed by adjustment of pH* to 7.0. Data collection commenced within 6 h of sample preparation. Under the conditions used in these experiments, the amount of degradation of urea to give ammonium cyanate under the conditions used is expected to be negligible (23).

The cyanide derivative of horse heart myoglobin (CN-myoglobin) was prepared by addition of a slight excess of sodium cyanide to 3 mM oxidized myoglobin (Sigma) in 50 mM sodium phosphate in H$_2$O (with 10% D$_2$O added for lock) or D$_2$O, stored at 4°C, and used within 12 h. The uncorrected pH (pH*) of urea solutions was adjusted to 7.0, and the concentration of denaturant was determined by refractive index measurement (22). Urea and protein solutions were combined to give the desired final concentrations, followed by adjustment of pH* to 7.0. Data collection commenced within 6 h of sample preparation. Under the conditions used in these experiments, the amount of degradation of urea to give ammonium cyanate under the conditions used is expected to be negligible (23).

The cyanide derivative of horse heart myoglobin (CN-myoglobin) was prepared by addition of a slight excess of sodium cyanide to 3 mM myoglobin (Sigma) in 50 mM phosphate buffer in D$_2$O, pH* 7.0. The pH* was readjusted to 7.0 after mixing, and the Soret band shift to 422 nm confirmed complete cyanide binding. The concentration was measured by using an extinction coefficient of 116,000 liters/mol-cm at 422 nm (24).

Abbreviations: cyt c, ferricytochrome c; Har, homoarginine; (Lys$_{19}$→Har$_{19}$) cyt c, guanidinated cyt c; NDE, nuclear Overhauser effect; pH*, uncorrected pH; 1D, one-dimensional.

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According to a standard literature preparation (26). A sample of terpyridine platinum(II) chloride ([Pt(trpy)Cl]Cl) was prepared and used in NMR experiments.

Chemical Modification of Cyt c. To convert all Lys residues in cyt c to homoarginine, a 0.8 mM cyt c solution was incubated with 0.5 M o-methylisourea hydrogen sulfate (Aldrich) for 120 h at pH 11.0, 4°C, according to a literature preparation (25). Modified protein was purified by gel filtration on Sephadex G-25 (Amersham Pharmacia). Amino acid analysis (University of Rochester Microchemical Protein Peptide Core Facility) confirmed that all 19 Lys residues were modified. To modify His residues, chloro(2,2',6',2'-terpyridine)platinum(II) chloride ([Pt(trpy)Cl]Cl) was prepared according to a standard literature preparation (26). A sample of 2 mM cyt c was incubated with equimolar [Pt(trpy)Cl]Cl for 72 h in 0.1 M sodium acetate, pH 5.0, 22°C, followed by chromatography on CM Sepharose (Amersham Pharmacia) as described in detail (27). UV-vis spectra of fractions collected from the CM Sepharose column indicated that unmodified, singly modified, and doubly modified products eluted as expected (27). The presence of a 342-nm band indicated modification by the reagent, and its ratio to the protein Soret band indicated whether the fraction collected contained singly or doubly modified protein (27). The fraction containing the doubly modified protein, bis-(2,2':6',2'-terpyridinyl)platinum(II)-ferricytochrome c ([Pt(trpy)]cyt c), was collected and used in NMR experiments.

Collection and Analysis of 1H NMR Spectra. 1H NMR spectra were collected on a Varian 500-MHz spectrometer at 30°C, 45°C, or 55°C with presaturation of the residual solvent residue and repetition rates of 4 s⁻¹ or 0.5 s⁻¹. One-dimensional (1D) NOE difference spectra were collected by using a 3 s⁻¹ repetition rate at 30°C and 45°C. Difference spectra were obtained by alternately collecting eight scans with the decoupler applied to the peak of interest and a reference point in the spectrum, followed by subtraction of the resulting free induction decays (FIDs). Magnetization transfer experiments were performed with the same pulse sequence as the 1D NOE but at 55°C. Data were processed by using FELIX 97 (Molecular Simulations). Exponential multiplication (20–50 Hz) was applied to FIDs before Fourier transform to enhance detection of broad resonances. Chemical shifts were referenced to 3-(trimethylsilyl)-1-propanesulfonic acid (DSS) via the residual ¹H F2O or H2O signal. All samples used for integration had a 1.5 mM protein concentration. Integration was performed by using 3 mM CN-myoglobin as an external standard in a coaxial insert. Integrals of native and non-native heme methyl peaks in varying concentrations of urea-d₄ were related to the integrals of the native heme methyl peaks in the absence of urea via the resolved CN-myoglobin heme 5-CH3 peak (28). Measurement at each urea concentration was repeated at least three times, and the results were averaged.

Results and Discussion

Detection of Non-Native Heme Resonances in the Presence of Urea. The chemical shifts of the substituents of low-spin ferric hemes (S = 1/2) are extremely sensitive to heme axial ligation. In native cyt c, the interaction of the axial ligands (His-18 and Met-80) with the iron is the primary factor that determines the asymmetry of unpaired electron spin density on the heme, resulting in the characteristic chemical shift pattern observed for the four heme methyl groups (29–32). Replacing Met-80 with other ligands such as a lysine (as in the alkaline form of cyt c, ref. 33), cyanide (34), imidazole (35), or pyridine (36) causes the heme resonance shift pattern to change entirely (Table 1). The relationship between heme substituent shifts and heme ligation is used to probe cyt c heme ligation in the presence of denaturant.

Upon subjecting horse cyt c to partially denaturing conditions (3–8 M urea-d₄, 50 mM sodium phosphate, D₂O, pH* of 7.0, 30°C (the midpoint for unfolding using CD spectroscopy at this temperature is 7.2 M urea, R.M., unpublished results)] well-

![Structure of c-type heme group and Fischer nomenclature used in the text. The covalent thioether linkages are shown at positions 2 and 4. P denotes propionate group.](Image)

**Table 1. Chemical shifts for selected heme substituents of cyt c derivatives**

<table>
<thead>
<tr>
<th>Cyt c derivative</th>
<th>8-CH₃</th>
<th>5-CH₃</th>
<th>3-CH₃</th>
<th>1-CH₃</th>
<th>δ-meso</th>
<th>Temp, °C</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Native cyt c *</td>
<td>34.5</td>
<td>10.2</td>
<td>31.7</td>
<td>7.2</td>
<td>2.7</td>
<td>30</td>
<td>(32)</td>
</tr>
<tr>
<td>Species L</td>
<td>23.9</td>
<td>22.2</td>
<td>11.9</td>
<td>13.2</td>
<td>-5.6</td>
<td>30</td>
<td>twv</td>
</tr>
<tr>
<td>Alkaline horse cyt c †</td>
<td>23.4</td>
<td>22.1</td>
<td>12.9</td>
<td>13.2</td>
<td>-5.4⁴</td>
<td>30</td>
<td>(33)</td>
</tr>
<tr>
<td>CN-Met80Ala cyt c</td>
<td>22.5</td>
<td>19.5</td>
<td>11.3</td>
<td>15.4</td>
<td>-4.3</td>
<td>30</td>
<td>(34)</td>
</tr>
<tr>
<td>Species H</td>
<td>25.7</td>
<td>18.6⁵</td>
<td>13.5</td>
<td>nd</td>
<td>nd</td>
<td>30</td>
<td>twv</td>
</tr>
<tr>
<td>Im-cyt c</td>
<td>24.2</td>
<td>14.1</td>
<td>16.7</td>
<td>10.7</td>
<td>nr</td>
<td>46</td>
<td></td>
</tr>
<tr>
<td>py-cyt c</td>
<td>28.6</td>
<td>20.5</td>
<td>17.2</td>
<td>14.7</td>
<td>nr</td>
<td>32</td>
<td>(36)</td>
</tr>
</tbody>
</table>

*Shifts measured at 30°C. The shifts of 1-CH₃ and δ-meso were measured from cross peaks between δ-meso/8-CH₃ and δ-meso/1-CH₃ in a two-dimensional NOE spectroscopy spectrum (R.M. and K.L.B., unpublished work).

†Assignments for one of the two conformers. Chemical shifts measured for horse cyt c in 50 mM sodium phosphate, pH 10.5.

⁴δ-meso for the alkaline form was assigned by 1D NOE (R.M. and K.L.B., unpublished work).

⁵Tentative assignment.
resolved non-native resonances appear at the expense of those caused by the native protein (Fig. 2B–D). Upon fully denaturing the protein by exposure to >9 M urea-d4, only selected non-native peaks (i.e., at 25.7 and 18.6 ppm) remain whereas other non-native (i.e., at 23.9 and 22.2 ppm) and native resonances are no longer observed (Fig. 2D). The species that gives rise to the non-native resonances present in moderate but not in high concentrations of urea-d4 will be referred to as species L. At a lower urea concentration and higher temperature (5.4 M urea at 55°C; Fig. 3A) species L is highly populated, with well-resolved resonances. Four of these resonances (23.9, 22.2, 13.2, and 11.9 ppm at 30°C) have intensities of three relative to well-resolved resonances. Four of these resonances (23.9, 22.2, 21.0, and 19.6 ppm) of the non-native form L, allowing assignment of LI and LIV in species L to heme 8-CH3 and 3-CH3, respectively. (D) Reference spectrum, 90% D2O, 40°C, 5.5 M urea-d4. (E) 1D NOE difference spectrum (90% D2O/10% H2O, 40°C, 5.5 M urea-d4) showing the NOEs observed upon saturating the δ-meso resonance (lII).

Assignment of Non-Native Heme Resonances. The resonances attributed to non-native species L are detected under partially denaturing conditions but not under strongly denaturing conditions. This observation indicates that species L is an equilibrium unfolding intermediate. The significant change in hyperfine shifts from native suggests that species L has non-native heme ligation. To aid in determining its heme ligation state, assignments of the heme methyl resonances for species L were made through 1D NOE and magnetization transfer experiments (Fig. 3). Magnetization transfer

from the well-resolved heme 8-CH3 and 3-CH3 resonances of native cyt c at 55°C allows assignment of resonances L1 and L1 and HII to 8-CH3 and 3-CH3 of the non-native form L, respectively (Fig. 3B and C). Irradiation of an upfield one-proton non-native resonance (lII, at ~5.6 ppm at 30°C) results in strong NOEs of similar intensity for two of the heme methyls, L1 and LIV, allowing assignment of LI and LIV in species L to heme 8-CH3 and 3-CH3, respectively. Upon fully denaturing, which improved quality of saturation transfer data [preparation and further studies of (Lys19→Har19)-cyt c are discussed below]. Irradiation of resonance H1 in this derivative results in saturation transfer to native heme 8-CH3, allowing assignment of H1. Assignment of H3 was made through observation of weak saturation transfer upon irradiating native methyl 3-CH3. The saturation transfer experiment aided in the detection of H3 at 13.5 ppm (3-CH3 for species H) which is not well resolved in 1D spectra. Irradiating resonance H3 did not yield reliable saturation transfer results because of overlap with the native heme propionate resonance (7-Hα). The remaining heme methyl group (HIV) is not resolved for species H. Although resonance HIV could not be assigned by saturation transfer, the observation that heme 5-CH3 shifts are always larger than heme 1-CH3 shifts for S = 1/2 ferric heme proteins (30) allows its tentative assignment to 5-CH3.

Relationship Between Heme Resonance Assignments and Heme Ligation. Low-spin ferric heme proteins with similar type and orientation of axial ligands have similar heme methyl shift patterns...
(Table 1). This is attributed to the dominant role played by the heme axial ligands in determining the unpaired electron spin density pattern on the heme and, thus, heme substituent contact shifts (29–31). In cyt c, the proximity of the native His-18 ligand to the covalent Cys thioether linkages to the heme (Cys-14 and Cys-17) causes it to remain strongly bound to the heme iron upon denaturation at neutral pH. In the cyt c heme octapeptide (MP-8), which consists of the heme and residues 14–21, it has been determined by NMR that the native His-18 remains bound at an angle nearly the same as in native cyt c despite the absence of most of the polypeptide (37). Changes in heme substituent shifts for cyt c, therefore, are attributed to replacement of Met-80 with non-native ligands, whereas the His-18 maintains its native orientation. The pattern of heme methyl shifts for species L bears a striking resemblance to that observed for the alkaline form of cyt c, known to have two conformers in which the axial Met ligand is replaced with Lys side chains (Table 1) (33, 38). It also bears a strong similarity to the cyanide adducts of Met-80–Ala cyt c (34) and MP-8 (37). Species L, alkaline cyt c, and these cyt c derivatives all have heme methyl shifts ranging from 25 to 10 ppm with an order of 8-CH3 > 5-CH3 > 1-CH3 > 3-CH3, as well as an upfield-shifted heme δ-meso resonance (Table 1) (32–37). Its similarities with the NMR properties of these characterized paramagnetic proteins indicate that the cyt c unfolding intermediate in urea is ligated by the native His-18 and a second ligand with cylindrical symmetry. The only such available donor in horse cyt c that would give a low-spin heme is an amino group. Because the N terminus of horse cyt c is acetylated (39), the remaining candidate is a deprotonated Lys side chain. The heme in species L thus is proposed to be ligated by the native His-18 and a deprotonated Lys.

As bis-His heme ligation has previously been established in denatured cyt c (2), His is expected to be the non-native heme ligand in species H, which persists under strongly denaturing conditions. The assignments (H and HIII to 8-CH3 and 3-CH3) and proposed assignment (HII to 5-CH3) for the heme methyl resonances are indeed similar to those observed for the pyridine (36) and imidazole (35) derivatives of cyt c (Table 1). Note that the ordering of 5-CH3 and 3-CH3 shifts in imidazole-cyt c is reversed from that observed for the pyridine derivative and proposed for species H. This is consistent with small differences in the angle at which these planar ligands bind with respect to the heme in-plane axes (30).

Investigations of Chemically Modified Derivatives. To test the proposal that species H and L observed by NMR have bis-His and His-Lys heme ligation, respectively, derivatives of cyt c were prepared in which His or Lys residues were modified to render them incapable of binding the heme. To make a variant in which non-native His are prevented from acting as heme ligands, His-26 and His-33 both were modified with chloro(2,2'-6',2'-terpyridine)platinum(II) chloride ([Pt(trpy)Cl2]Cl) to give [Pt(trpy)Cl2]cyt c (bis-(2,2'-6',2'-terpyridine)platinum(II)-cyt c) (27). The 1H NMR chemical shifts of the heme substituents for [Pt(trpy)Cl2]cyt c are similar to those observed for native, indicating that the protein’s structure is not significantly perturbed by modification. The platinum-modified protein, when partially denatured by urea, shows hyperfine-shifted peaks L1 and L2 (Fig. 4B), supporting the proposal that species L does not have bis-His ligation. In addition, resonances attributed to species H are absent for partially denatured [Pt(trpy)Cl2]cyt c, supporting the assignment of species H to a bis-His species.

To prepare a cyt c variant lacking Lys side chains, all 19 Lys residues in cyt c were converted to Har by chemical guanidination to give (Lys19→Har19)-cyt c (25). Native cyt c and (Lys19→Har19)-cyt c have nearly the same heme resonance shifts, verifying that guanidination does not significantly perturb the protein fold. Upon partial urea denaturation, this variant gives rise to resonances attributed to species H but not to species L, supporting the assignment of His-Lys heme ligation to species L (Fig. 4 C and D). The NMR spectra of both [Pt(trpy)Cl2]cyt c and (Lys19→Har19)-cyt c were examined over a range of urea concentrations (5.0–7.4 M) and temperatures (20–55°C) to verify that the inability to observe species H for [Pt(trpy)Cl2]cyt c or species L for (Lys19→Har19)-cyt c is not a result of a perturbation of the energetics of the conformational change for the variant as compared with native.

Thermodynamic Analysis. The fractional populations of each of the observable conformations (native, species L, and species H) were measured over a range of urea-d4 concentrations by integration of resolved heme methyl resonances (Fig. 5; for details, see Materials and Methods). Note that the sum of the measured fractional populations is clearly less than one at high

![Fig. 4. 500-MHz ·H NMR spectra (50 mM sodium phosphate, 90% D2O/10% H2O, pH* 7.0) of (A) cyt c in 7.0 M urea-d4, 30°C. (B) Bis-(2,2'-6',2'-terpyridine)platinum(II)-ferriocytochrome c ([Pt(trpy)Cl2]cyt c) in 7.2 M urea-d4, 30°C. (C) (Lys19→Har19)-cyt c in 7.0 M urea-d4, 30°C. (D) (Lys19→Har19)-cyt c in 7.0 M urea-d4, 45°C. Raising the temperature to 45°C in D increases population of species H relative to native.

![Fig. 5. Plot of fractional populations of native protein, species L, and species H, as a function of urea-d4 concentration.]
urea concentrations. This indicates that there is at least one unobservable species (species O, for “other”) present that is substantially populated under denaturing conditions. As denatured cyt c previously has been shown to be low-spin with bis-His heme ligation (1–3) and no resonances indicative of high-spin species were observed in these experiments, we propose that species O is a bis-His-ligated species in which there is substantial disorder in the angle at which the non-native His binds the heme, leading to lack of observable hyperfine-shifted resonances.

The possibility that species O and the other species present here consist of protein dimers or oligomers was considered. The shifts of native protein, species L, and species H were found to be independent of protein concentration (0.2–4.0 mM protein in 7 or 8 M urea-d_4). In addition, the fractional populations of species H and L do not increase with increasing protein concentration. These observations indicate that species H and L are monomeric. However, the fractional population of species O does increase with increasing protein concentration (for concentrations above 1.5 mM), indicating that it consists at least in part of protein dimers or oligomers. For the protein concentrations used in NMR studies (1.5 mM), small angle x-ray scattering results suggest that the amount of aggregated protein present should be minimal (40). In addition, our small angle x-ray scattering results suggest that the amount of unfolding that it consists at least in part of protein dimers or oligomers. However, the fractional population of species O does increase with increasing protein concentration (for concentrations above 1.5 mM), indicating that it consists at least in part of protein dimers or oligomers.

The fractional populations determined through integration of NMR resonances (native, species L, and species H) were fit to a four-state thermodynamic model. The population of species O was defined as f_O = 1 - f_N - f_L - f_M (f_N, f_L, and f_M are the measured populations of native, species L, and species H, respectively). The fractional populations can be expressed in terms of the free energies of the transitions from the native state to each of the others (L, H, and O), where ΔG^D_O is the free energy change for a transition from the native state to state X at a given denaturant concentration. The dependence of the free energy on urea concentration for each state is assumed to be linear (ΔG^D_O = ΔG^D_O(urea) - m_X [urea]) (41), and the data were fit by linear least squares to extract m_X values and ΔG^D_O(urea) in the absence of denaturant. The resulting values of ΔG^D_O and m_X are 17 kJ/mol and 2.2 kJ/(mol-M), and the values obtained for ΔG^D_H and m_H are 47 kJ/mol and 6.5 kJ/(mol-M).

Examination of Fig. 5 clearly reveals that species L is an equilibrium unfolding intermediate. The detection of an intermediate is consistent with previous studies using small angle x-ray scattering and UV-vis spectroscopy. At least one intermediate state for equilibrium cyt c unfolding by guanidine hydrochloride was reported by researchers using small angle x-ray scattering (40). Others have noted that loss of the 695-nm band, attributed to a Met-80 sulfur → Fe(III) charge transfer, slightly precedes loss of CD signal upon guanidine hydrochloride denaturation and that this may indicate deviation from two-state unfolding because of ligand replacement (42). Species L also may serve as an intermediate on the kinetic folding pathway; recent results implicate a Lys-ligated intermediate in the slow folding of cobalt-substituted cyt c (7). The small values of ΔG^D_O and m_X suggest that species L has a structure quite similar to that of the native protein (41); this is consistent with the lack of evidence for an intermediate in CD studies. Replacement of Met-80 by one or more Lys (Lys-72, -73, or -79) could be achieved without substantial disruption to the overall protein fold. Interestingly, the value for ΔG^D_O is similar in magnitude to that measured for a low-energy local unfolding of cyt c in low concentrations of guanidine hydrochloride by isotope exchange studies (25.1 kJ/mol) (17). This local unfolding was attributed to unfolding of the small loop consisting of residues 70–85 (i.e., including Met-80 and three Lys). Experiments designed to identify sequence specifically the Lys ligand(s) in species L are in progress.

The high intrinsic pKa of Lys (10.8) makes it an unlikely heme ligand at neutral pH. The Lys residues proposed as candidates for ligation to the heme iron in species L are the same as those proposed to bind heme in the alkaline form of the protein, which forms with a pKa of 9 (43). In the alkaline form of Saccharomycetes cerevisiae iso-1-cytochrome c, Lys-73 and -79 have been identified as the non-native heme ligands (38). By analogy, Lys-73 and -79 (as well as Lys-72, which is trimethylated in S. cerevisiae cyt c but not in horse cyt c) are candidates for heme ligands in the alkaline form of horse cyt c. Given that the pKa for the formation of the alkaline form is substantially lower than that for the side chain of a Lys, it has been proposed that deprotonation of another group (yet to be identified) acts to trigger a conformational change that leads to loss of Met ligation and introduction of the Lys into the heme vicinity. The formation of the strong Lys-Fe bond may then drive Lys deprotonation (43). The lowering of the pKa of the alkaline transition to values as low as 7.2 by mutating the heme pocket residue Phe-82 indicates that perturbation of heme pocket structure may drive displacement of Met by Lys, even near neutral pH (44). We propose that a similar conformational change occurs for cyt c under mildly denaturing conditions.

NMR analysis of denatured cyt c has allowed the direct detection of a form of cyt c with bis-His heme ligation. The ΔG^D_H measured here (47 kJ/mol) is similar to the ΔG^D determined for the global unfolding reaction by isotope exchange (53.6 kJ/mol) (17). The presence of observable hyperfine-shifted peaks for species H indicates that the heme ligation geometry is well defined in this species. In other words, species H represents a denatured form of cyt c with a well-defined non-native structural (heme-ligand) contact. NMR analysis of the urea denaturation of cyt c thus has shown that it involves at least four states, three of which (native, species L, and species H) have well-defined heme ligation geometry.

We thank Andrew Vetter for his assistance with CD and NMR experiments and James McGarrah for assistance in preparing [Pt(trpy)Cl]Cl. This work was supported by National Science Foundation-Research Experience for Undergraduates Grant CHE-9619935 and the University of Rochester.