Computational de novo Design, and Characterization of an A$_2$B$_2$ Diiron Protein

Christopher M. Summa†, Michael M. Rosenblatt†, Jae-Kyoung Hong, James D. Lear and William F. DeGrado*

Department of Biochemistry and Biophysics
School of Medicine
The University of Pennsylvania
1010 Stellar-Chance Bldg
421 Curie Blvd, Philadelphia, PA 19104-6059, USA

Diiron proteins are found throughout nature and have a diverse range of functions; proteins in this class include methane monooxygenase, ribonucleotide reductase, Δ$^9$-acyl carrier protein desaturase, ruberythrin, hemerythrin, and the ferritins. Although each of these proteins has a very different overall fold, in every case the diiron active site is situated within a four-helix bundle. Additionally, nearly all of these proteins have a conserved Glu-Xxx-Xxx-His motif on two of the four helices with the Glu and His residues ligating the iron atoms. Intriguingly, subtle differences in the active site can result in a wide variety of functions. To probe the structural basis for this diversity, we designed an A$_2$B$_2$ heterotetrameric four-helix bundle with an active site similar to those found in the naturally occurring diiron proteins. A novel computational approach was developed for the design, which considers the energy of not only the desired fold but also alternatively folded structures. Circular dichroism spectroscopy, analytical ultracentrifugation, and thermal unfolding studies indicate that the A and B peptides specifically associate to form an A$_2$B$_2$ heterotetramer. Further, the protein binds Zn(II) and Co(II) in the expected manner and shows ferroxidase activity under single turnover conditions.

Introduction

The design of proteins de novo$^{1,2}$ has progressed significantly within the last decade. While the design of native-like proteins is by no means a routine endeavor, sufficient progress has been made that attention is beginning to focus on the design of functional proteins. One aspect of function currently being explored involves the design of a binding interaction between a de novo protein and a cofactor$^{3–13}$ or another protein.$^{14,15}$ A far more complex and subtle design objective is the rational design of de novo proteins with catalytic properties.

With the exception of some elegant demonstrations of the generation of novel enzyme activities through the mutagenesis of a naturally occurring protein,$^{6,16–26}$ most research seeking novel enzymatic activities has focused on combinatorial approaches such as in vitro evolution,$^{27–29}$ domain shuffling,$^{30}$ incremental truncation,$^{31}$ and phage display methods.$^{32–46}$ Recently, catalytic activities have been reported in some de novo designed systems.$^{47–49}$

We have focused on the design of diiron-binding proteins because of their extraordinary functional diversity. Representative members of this class of proteins include methane monooxygenase (MMO)$^{50–57}$ ribonucleotide reductase (RR)$^{50,58}$ Δ$^9$-acyl carrier protein desaturase$^{59,60}$ hemerythrin$^{61–64}$ ruberythrin$^{65}$ and the ferritins.$^{66}$ These proteins function as monooxygenases, radical generators, oxidoreductases, desaturases, ferroxidases, and oxygen and iron storage proteins. While the overall folds of the proteins differ, the structural subunits responsible for forming the diiron sites are remarkably similar four-helix bundles.$^{67,68}$ One of the goals of this project is to identify the determinants of metal binding and substrate activation through the study of simplified
dimetal-binding systems. Interestingly, recent protein engineering studies of the diiron site in natural ribonucleotide reductase R2 subunits have resulted in the production of new catalytic intermediates.69 – 71

In previous work, antiparallel homo-dimeric four-helix bundle proteins were designed using a helix-turn-helix motif that binds metal ions including Zn, Fe, and Co ions.47,72 At the active site of these proteins are four glutamate and two histidine residues, including two conserved Glu-Xxx-Xxx-His motifs found in nearly all diiron proteins of this class. These proteins were produced using previously established methods employing symmetry operators to produce $C_2$ and $D_2$-symmetric backbone geometries.68 Here, we have taken the conserved metal binding site of our DueFerro1 protein (DF1) and modified it to form a heterotetrameric assembly, DFtet.

DFtet was designed to self-assemble into a

**Figure 1.** Sequence, helical wheel diagram, and active site of DFtet. (a) The sequence of DF1 is shown along with DFtet. DF1 has a helix-turn-helix topology; a KL link between the helices has been omitted from (a). The ligating residues (green), those residues that form second-shell hydrogen bonds to the ligating residues (black), and a residue that is important for solvent access (brown) are shown in the linear sequence and the helical wheel diagram (b). The position of the 2-fold axis running through the dimetal site is shown at the left oval. The positions of the additional quasi 2-fold axis relating the backbone atoms are shown at the center and bottom of the diagram. (c) Two views of the dimetal active site of di-Zn(II) DF1.
heterotetrameric $\text{A}_2\text{B}_2$ bundle from four isolated peptide units. The protein assembles with a precise topology allowing for the generation of a dimetal binding site by utilizing residues from each of the four independent peptides. Thus, a heterotetrameric system can bridge the gap between “rational design” and combinatorial approaches. However, while a heterotetrameric system is uniquely suited to the production of a library of proteins from a significantly smaller number of peptides, the design of such a system presents significant challenges for protein design. Specifying a unique topology is a more complex problem for a heterotetrameric system, as in DFtet, than it was in a homodimeric system, such as DF1 or DF2. In particular, it was important to destabilize both homoooligomeric folds as well as undesired heterotetrameric topologies. Therefore, new computational design methods were used to solve this problem. While the concept of “negative design” (or designing against alternative structures) has been considered in the past and see DeGrado et al., and references therein), this idea has not been explicitly codified into a computational design algorithm that was tested experimentally.

**Results**

**Computational design of DFtet**

DueFerro tetramer, or DFtet, is a set of two 33-residue peptides that have been designed to self-associate into a dimetal-binding $\text{A}_2\text{B}_2$ heterotetrameric protein. Because the target was a heterotetramer, we used negative design, to prevent alternate topologies from occurring. Also, we used a “minimalist” approach in order to minimize extraneous structural variables, which might complicate the design and interpretation of results. The protein sequence was intended to be as simple as possible, while retaining the ability to tetramerize specifically into the desired topology and bind two metal ions.

The first step in the design process involved the specification of the backbone. Although there are two different chains in DFtet, the backbone structure shows approximate $D_2$ symmetry with three orthogonal 2-fold rotational axes as shown in Figure 1. The parameters used to generate the four-helix bundle backbone of DFtet were obtained from an analysis of DF1, as described in Materials and Methods. The helices of DFtet were extended relative to those of DF1 (33 residues in DFtet versus 24 residues in DF1) to increase the stability of the DFtet system by increasing the size of the hydrophobic core.

While the backbone conformation of DFtet is approximately $D_2$ symmetric, the metal-binding site shows lower symmetry ($C_2$), because only two of the four helices donate His side-chains to the binding site (Figure 1(b) and (c)). Thus, in the second stage of the design, side-chains were asymmetrically placed onto the helices, resulting in an $\text{A}_2\text{B}_2$ heterotetramer with $C_2$ symmetry. The designation A was assigned to the peptide containing only the Glu ligands, and the designation B was assigned to the peptide containing the EXXH metal-binding motif. The residues within the region of the binding site are the same as the corresponding residues of DF1. Specifically, residues A12–A23 of DFtet are identical with residues 6–17 of DF1, and residues B12–B23 of DFtet correspond to residues 32–43 of DF1, with two exceptions. Residue 22 in DFtet-A was mutated from Leu (residue 13 in DF1) to Ala because Leu19 blocks access to the active site in the crystal structure of DF1. Second, residue 18 in DFtet-B (corresponding to residue 38 of DF1) was mutated from Lys to Arg, Arg at this position might form a hydrogen bond with Asp15, which itself hydrogen bonds directly to the coordinating histidine. This Asp-Glu-Xxx-Arg-His motif is seen in diiron proteins such as MMO and $\Delta^2$-ACP desaturase; the arg might position the liganding His residue and possibly also tune the redox potential of the metal site.

The ligand-binding site, together with a requirement for low-energy side-chain rotamers entirely defined the backbone conformation. Residues at the remaining $a$ and $d$ positions were modeled as leucine because this side-chain effectively filled the interior volume of the bundle. This side-chain also has a high intrinsic helical propensity and is known to stabilize antiparallel four-helix bundles, such as ROP. Residues at $f$ positions outside the vicinity of the binding site were chosen to maximize the overall helical propensity of the sequence, while minimizing hydrophobicity and electrostatic interactions with the interfacial residues. These positions were modeled as Ser or Ala in an alternating manner along the chain.

The nature of the residues at the remaining $e, g, b, c$ positions was chosen to specifically stabilize only one of the possible topologies for an antiparallel $\text{A}_2\text{B}_2$ heterotetramer (Figure 2). The desired topology is shown in Figure 2(a) with two alternate topologies (shown in Figure 2(b) and (c)) that may be considered. Clearly, there are other possible configurations for the protein than those shown, such as parallel structures or antiparallel structures with frame shifts along the central axis of the bundle. We made the assumption that these frame shifts would not occur, because of the energetic cost of exposing excess non-polar surface area. Also, parallel tetramers are possible, although modeling suggests that they would result in distorted metal-binding sites, which lacked some of the liganding interactions. In Figure 2(a), the $b/e$ interface lies at the helix $A/helix$ B boundary and the $g/c$ interface lies between both the $A/A$ and the $B/B$ boundaries. The reverse situation holds for the other topology (Figure 2(b)); in Figure 2(c) both the $b/e$ and $g/c$ interfaces lie between helices $A$ and $B$.

Interacting interfacial positions were allowed to
Figure 2. Possible topologies for an antiparallel tetrameric coiled-coil. The upper panel shows three different possibilities for an A₂B₂ heterotetramer. Helices of like color are identical (red denotes an A-helix and yellow denotes a B-helix) and the directionality of the chain is denoted by the markings denoting the N and C termini. The topology in (a) is the desired topology, having g/c interfaces between helices of identical sequence, and b/e interfaces between helices of different sequence. The topologies shown in (b) and (c) are undesirable in the present study, and a computational design algorithm was implemented to prevent these topologies from occurring in the design of DFtet. In (b), the g/c interface now lies between helices of different sequence, and the b/e interface lies between helices with identical sequence, opposite to what is found in (a). The topology shown in (c) is an intermediate between the previous two examples; both g/c and b/e interfaces occur between helices which differ in sequence. Configurations (d) and (e) show what would occur if a homotetramer were allowed to form. (f) The different interfaces of the intended antiparallel heterotetramer in detail.
assume an identity of either $+$ or $\pm$. These designations would ultimately correspond to residue identities of Lys and Glu, respectively, in the amino acid sequence. We chose not to allow neutral or non-polar residues at these positions in order to maximize solubility in aqueous solution. Interacting pairs were scored as follows:

$$E_{\text{contact}} = \left[ \begin{array}{c} +2, \quad +/ + \text{ interaction} \\ +3, \quad -/ - \text{ interaction} \\ -1, \quad +/ - \text{ interaction} \end{array} \right]$$

A larger unfavorable “energy” for $-/-$ over $+/+$ interactions was used because Glu has shorter side-chains than Lys, and hence is less able to maximize the distance between one another. A pair of residues was labeled interacting if the distance between their C$^\alpha$ atoms was less than 8.5 Å. These scores are a simplified representation of experimentally measured interhelical electrostatic interaction energies. Since the energies were intended only as an approximation to find an optimal pattern of charge, and not to calculate binding energies, this simplification was reasonable.

The desired topology and a single alternate topology (Figure 2(b)) were explicitly considered in the calculation of the energy function. The energy function used was:

$$E_{\text{tot}} = E_{\text{desired}} - E_{\text{undesired}}$$

Figure 3. CD spectra of DFtet are shown at various pH values. (a) Spectra at pH 7.0 in the absence of exogenous metal, (b) spectra at pH 7.0 in the presence of one equivalent of Zn(II) and (c) spectra at pH 4.0 in the absence of exogenous metal. In each panel, red diamonds denote spectra of DFtet-A alone, blue triangles denote spectra of DFtet-B alone, and filled black circles show spectra for a 1:1 mixture of DFtet-A and DFtet-B.
\[ E_x = \sum_{i=1}^{N_{\text{contacts}}} E_{\text{contact}}(i) \] (3)

where \( x \) designates a particular three-dimensional conformation for the sequence. The identities of all the variable positions were initially set to 0, representing no charge. At each step in the optimization protocol, a residue was chosen at random, and its identity was flipped to that of the opposite sign (or, in initial cases where the identity was unsigned, to +). At this point, the interaction energies for the new sequence were calculated. If the new sequence was lower than the lowest energy sequence yet encountered, the program would retain the sequence and its energy value. This process was run through 700,000 iterations for the DFtet sequence, and three-dimensional models were built for the sequences with the top four scores. A sequence was chosen (ranked second among all the top scorers) which showed the minimum number of unfavorable contacts in the desired topology (among the top four considered) and is shown in Figure 1. Visual inspection also showed that these sequences would show multiple unfavorable electrostatic interactions as a parallel homo or heterotetramer.

**Solution characteristics of the protein**

**CD spectroscopy**

At pH 7.0, the circular dichroic (CD) spectra of DFtet showed strong bands at 222 nm and 208 nm, indicating a high content of \( \alpha \)-helical secondary structure (Figure 3(a)). By contrast, the individual DFtet-A and DFtet-B peptides showed a spectrum typical of a random-coil. This finding suggested that the DFtet-A and DFtet-B peptides are unstructured in the absence of their partners, and were able to form stable secondary structure only when mixed together at this pH. In the case of peptide DFtet-A, formation of a homooligomer at pH 7.0 would require the burial of one active-site glutamic acid residue per peptide. This is clearly an energetically unfavorable process, and likely inhibited association. A homooligomer of peptide DFtet-B at this pH would involve the burial of one His and one Glu per peptide.

To confirm that DFtet-A and DFtet-B peptides associate with a 1:1 stoichiometry, different molar ratios of the peptides were mixed and their signal at 222 nm was evaluated. A minimum (signifying maximal helical content) occurred at a molar ratio of precisely 0.5, indicating that the stoichiometry of the peptides was 1:1 (Figure 4).91–93

The next series of experiments were undertaken to determine the effects of transition metal ions on the secondary structure of the DFtet at neutral pH. Zn(II) was chosen for these experiments, because it binds strongly to the active sites of other diiron proteins.47,72,94 The secondary structure of DFtet was not significantly altered by the addition of Zn(II) (Figure 3(b)). Similarly, the addition of Zn(II) had little effect on the secondary structure of the DFtet-A peptide, presumably because Zn(II) ions are rarely coordinated by carboxylate groups in the absence of other liganding groups in protein structures. DFtet-B, on the other hand, did appear to form significant secondary structure in the presence of the Zn(II) at pH 7.0, which was presumably due to coordination of the EXXH motifs. Histidine is commonly found to be an active-site ligand in Zn(II) proteins95 and Zn(II)-assisted protein folding has been reviewed recently.96

Protonation of the active-site glutamate residues has been shown to stabilize the homodimeric DF2,47 an analogous designed protein. Thus, we were interested to determine the effect of low pH on the individual components of the DFtet system. At pH 4.0, below the unperturbed pK, of the Glu side-chain, DFtet-A, DFtet-B and a 1:1 molar ratio of these two peptides showed a spectrum typical of the \( \alpha \)-helix (Figure 3(c)).

**Sedimentation equilibrium ultracentrifugation**

Equilibrium analytical ultracentrifugation was used to determine the molecular mass of DFtet. The proteins were centrifuged at 40,000, 45,000, and 48,000 rpm, and the data were globally analyzed to determine the mass averaged molecular mass. Analysis of a 1:1 mixture of DFtet-A and DFtet-B, at pH 7.0 indicated the formation of a tetramer with an apparent molecular mass of 15,400 Da (Figure 5(a)). Addition of one equivalent of Zn(II) to the 1:1 mixture resulted in an improved fit to the data (Figure 5(b)), and an observed
Figure 5. Equilibrium ultracentrifugation analysis of (a) DFtet-A + DFtet-B at pH 7.0 in the apo state, (b) in the presence of Zn(II) at pH 7.0, and (c) at pH 4.0 in the apo state. Data from 40,000, 45,000, and 48,000 rpm were used in a simultaneous fit of a fixed molecular mass model.
molecular mass (16,500 Da) in excellent agreement with that expected for the tetramer 16,400 Da.

In contrast, DFtet-A was monomeric even at concentrations tenfold above the CD measurements (data not shown). The average molecular mass for DFtet-B was between that expected for a dimer and trimer, suggesting an intermediate degree of aggregation (data not shown).

The pH-dependence of the oligomerization state was also assessed using ultracentrifugation. Data collected for DFtet-A alone at pH 4.0 were well described by a fit to a single species with an apparent molecular mass of 11,500 Da, close to that expected for a trimer (data not shown). Numerous coiled-coil peptides have previously been reported to form trimers. 97 A monomer–tetramer equilibrium model provided the best fit to the sedimentation data collected for DFtet-B; attempts to use a single-species model resulted in poor fits. DFtet (a 1:1 mixture of DFtet-A and DFtet-B) data were best described by a fit to a single tetrameric species at pH 4.0 (Figure 5(c)), with an apparent molecular mass of 15,000 Da (theoretical mass is 16,292 Da).

**Thermal unfolding**

After confirming that the A₂B₂ system was indeed helical and tetrameric at neutral pH, the thermal stability of the designed protein was investigated. Figure 6 illustrates the thermal stability of the apo-protein at 5 μM and 50 μM total concentration near pH 7.5. As expected for a self-associating system, the protein was more stable at a higher concentration. Even at 5 μM, however, the protein was very stable and unfolded only at elevated temperatures, with a midpoint near 75 °C. Addition of Zn(II) led to a dramatic increase in stability, with the protein becoming entirely stable up 95 °C. A comparison of the unfolding curves measured at pH 7.4 versus 6.0 (5 μM protein concentration in each case), also showed the anticipated pH dependence. At pH 6.0 the energetic cost associated with proton condensation at the active site 52 was less unfavorable, and the stability of the protein was increased.

**Co(II) binding**

Co(II) provided an especially convenient spectroscopic probe since it has d–d transitions in the visible region (between 500 nm and 700 nm), 98 whose molar extinction coefficient depend on the coordination environment of the Co(II). The extinction coefficient increases from 10 M⁻¹ cm⁻¹ to ~150 M⁻¹ cm⁻¹ to ~400–600 M⁻¹ cm⁻¹ as the coordination proceeds from octahedral to pentacoordinate to tetrahedral. Addition of CoCl₂ to a solution of the DFtet (36 μM in tetramer) produced a species with an absorption spectrum that is characteristic of Co(II) in a pentacoordinate environment. This result agrees with the design in which each Co(II) ion is ligated by a total of four carboxylate ligands (two from the two bridging glutamate residues and two from a chelating glutamate) and a single His ligand. Furthermore, the shape of the Co(II) spectrum (Figure 7 inset) was identical with that of DF2 47 and the extinction coefficients were the same within experimental error (DF2: λ = 520 nm, ε = 140 M⁻¹ cm⁻¹; λ = 550 nm, ε = 155 M⁻¹ cm⁻¹; λ = 600 nm, ε = 90 M⁻¹ cm⁻¹; DFtet: λ = 520 nm, ε = 125 M⁻¹ cm⁻¹; λ = 550 nm, ε = 140 M⁻¹ cm⁻¹; λ = 600 nm, ε = 80 M⁻¹ cm⁻¹). The spectrum also showed a very close correspondence with that of bacterioferritin (λ = 520 nm, ε = 125 M⁻¹ cm⁻¹; λ = 555 nm, ε = 155 M⁻¹ cm⁻¹; λ = 600 nm, ε = 107 M⁻¹ cm⁻¹) 94,99 whose active site resembles our intended design. 100 The correspondence of the spectra strongly suggests that the metal-binding site of DFtet adopts the intended geometry.

The stoichiometry of binding was probed by measuring the absorbance at 550 nm as a function of added Co(II) at a constant tetramer concentration of 36 μM (Figure 7). A linear increase in absorbance was observed, until a stoichiometry of two Co(II) per tetramer was reached. Beyond this point, no further significant increase in absorbance beyond that expected for hexa-aqua Co(II) was observed. These data indicate that the binding was tight and stoichiometric, and that the dissociation constant for binding to both sites was significantly lower than the total concentration of binding sites.

**Ferroxidase reaction**

A number of diiron proteins, such as bacterioferritin and ruberythrin, catalyze the oxidation of Fe(II) to Fe(III). In these proteins, this ferroxidase reaction 65,101–104 results in the formation of a diferric oxo-bridged cluster characterized by a broad charge transfer transition centered near 300 nm. 105,106 The time-course of Fe(II) oxidation in the presence of DFtet (50 μM in tetramer, Figure...
Under single turnover conditions, the reactivity of the DFtet was of the same order of magnitude as DF2;47 both were significantly more rapid than the uncatalyzed reaction, but nevertheless were about three orders of magnitude slower than the reaction catalyzed by bacterioferritin.

The ferroxidase reaction catalyzed by DF2 was first-order in both protein and Fe(II). Similarly, the initial rate of iron oxidation by DFtet depended linearly on the concentration of Fe(II) (Figure 8(b)). Furthermore, the individual time-courses at various iron concentrations successfully globally fit to an integrated rate equation which is first-order in both iron and protein (Figure 8(c)). The rate constant determined in this manner was 11.8 M⁻¹ s⁻¹, and the extinction coefficient at 320 nm for the dimeric diiron center was 3730 M⁻¹ cm⁻¹, which was within the range of extinction coefficients observed for diiron proteins including bacterioferritin¹⁰⁶ (3380 M⁻¹ cm⁻¹ at 300 nm), H-chain ferritin¹⁰⁴ (2990 M⁻¹ cm⁻¹ at 300 nm), horse spleen ferritin¹⁰⁴ (3540 M⁻¹ cm⁻¹ at 300 nm), ribonucleotide reductase¹⁰⁷ (4700 M⁻¹ cm⁻¹ at 325 nm), and stearoyl-ACP Δ⁹ desaturase¹⁰⁸ (2080 M⁻¹ cm⁻¹ at 325 nm).

Discussion
Backbone design
The method of retrostructural analysis allows the construction of a consensus backbone from a group of related protein structures. More importantly, it provides data with which to set ranges of backbone parameters for a protein within that class. With this method it should be possible to access global parameters (such as helical tilt angle, interhelical distance) and test their effect on function in a way that was heretofore impossible with single mutation-based approaches. These design techniques will be used in future work on this system.

Sequence design
Three criteria must be satisfied before a design can be successful. First, the target structure must be the lowest-energy structure for the designed sequence. Second, there should exist a large energy gap between the sequence folded into the desired structure and that of any other, undesired structure. Finally, the ground state configuration should be non-degenerate. The last two criteria ensure that the desired structure will be significantly populated relative to any alternative conformations.

Historically, computational protein design approaches have focused on satisfaction of the first criterion alone.¹⁰⁹,¹¹⁰ This method has, in general, proven successful in many real-world protein designs. Guttin & Shakhnovich¹¹¹ have also shown that for a lattice model of a heteropolymer (based on the HP-model designed by Dill and co-workers) the probability of finding a degenerate
ground state decreases exponentially with a decrease in the ground state energy. Therefore, the lower the energy of the design, the more likely that the designed structure will be non-degenerate. However, it appeared unlikely that the present problem could be solved by simply optimizing the energetics of the intended structure. Shakhnovich & Gutin have suggested that explicit consideration of alternative configurations was an essential part of the optimization process for designing a uniquely folded protein.\textsuperscript{111,112} Since exhaustively searching configurational space is impossible for all but the shortest peptide sequences, implementing such a design protocol in real-world protein design applications is non-trivial. In our case, we had a specific alternative configuration in mind; this may not be true of all protein design applications.

In our method, there were 28 unique residues which underwent optimization and each was allowed an identity of $+$ or $-$. This represents $2^{28}$ or roughly 268 million possible sequences. While it would have been impossible to enumerate each possible sequence for comparison, our goal was to

Figure 8 (a) and (b) (legend opposite)
develop methodology that could be used in other design projects; this energy function was optimized with a random search algorithm. A simulated annealing algorithm, a genetic algorithm, or other optimization protocol could also be used successfully.

Here, a single alternative conformation was used as the “unwanted” configuration. Homooligomeric species of DFtet-A and DFtet-B were not considered in the sequence design process. As can be seen from the CD and sedimentation results on the isolated peptides, DFtet-B is capable of forming a homooligomer in the presence of Zn(II) in solution. Ideally, both peptides could be designed while taking the homooligomeric states into account as undesired alternate topologies. While this optimization alone is unlikely to completely abolish homooligomer formation in the isolated peptides in solution (due to the favorable energetics of hydrophobic core burial), it may be possible to significantly destabilize these states through the use of carefully designed interhelical electrostatic interactions.

Solution characteristics

The design of DFtet was successful: DFtet-A and DFtet-B are unfolded at neutral pH; however, when mixed in a 1:1 ratio they assemble into a helical bundle with the expected stoichiometry. Furthermore the peptide is tetrameric and stable over a wide temperature range, being thermally unfolded only at low concentrations and at an elevated pH. The main transitions of the thermal unfolding curves are steep, as would be expected for a native protein. Only at low pH do the individual peptides associate into helical oligomers, as expected. DFtet also binds metal ions in the proper stoichiometry, and the spectroscopic properties of the Co(II) and Fe(III) derivatives strongly suggest that the ligand environment is precisely as intended. The match of the Co(II) spectrum between DFtet and DF2 (whose crystal structure has been determined, unpublished results), and the identical kinetic scheme for Fe(II) oxidation provide good evidence that the metal-binding sites of these proteins are very similar. Thus, the intended topology appears to have been achieved.

Materials and Methods

Automated design

The parameters used to define the structure of DF1 were determined by fitting to the crystal structure of DF1 using a genetic algorithm and equations similar to those employed previously. The best-fit Ds symmetric coiled-coil parameters (Figure 9, legend) were used to produce a model whose backbone trace was within 1.1 Å r.m.s.d. over 68 C atoms of the template backbone. These parameters were then used to create an elongated bundle as follows: because the intended product was a coiled-coil model, the α-helix was slightly overwound to give a precise seven residue repeat; next the bundle was built using the parameters described above, and the entire bundle was then

Figure 8. DFtet-catalyzed oxidation of Fe(II) to Fe(III). (a) The time-course of the ferroxidase reaction is followed by measuring the absorbance of the solution at 320 nm. (b) The initial rates are shown as a function of the initial Fe(II) concentrations at a total concentration of 50 μM tetramer. (c) Time-courses for the reaction (50 μM tetramer); the data were globally fit to a second-order rate equation \( \frac{A_{320}}{A_{320}} = \frac{1}{1 + \frac{[P]}{[Fe^{2+}]}} \ln \left( \frac{[Fe^{3+}]}{[Fe^{2+}]} \right) = k t \) in which \([P]\) is the initial protein concentration, \([Fe^{3+}]\) is the initial concentration of Fe(II), and \([Fe^{2+}] = (A_{320} - A_{320, \text{initial}})/\Delta A_{320}\). The rate constant was globally fit and found to be 11.8(± 0.8) M⁻¹ s⁻¹. The extinction coefficient at 320 nm was allowed to vary from run to run, but was nevertheless found to be approximately constant with a value of 1862(± 153) M⁻¹ cm⁻¹ (3729 M⁻¹ for the dimeric diiron center). Concentrations of initial [Fe²⁺] are listed above each curve and are color-coded.
an Applied Biosystems model 433A solid-phase peptide synthesizer. Standard coupling conditions were used. In order to minimize failure sequences, “capping” with acetic anhydride (100-fold excess) and DIEA (N,N-diisopropylethylamine) in NMP (N-methylpyrrolidinone) was performed after each coupling reaction. All peptides were acetylated at the amino terminus.

DFtet-A was cleaved using a mixture of 95.0% TFA (trifluoroacetic acid), 2.5% water, 2.5% TIS (trisopropylsilane) (by vol.). DFTet-B was cleaved with 81.5% TFA, 5% thioanisole, 5% phenol, 5% water, 2.5% EDT (ethanedithiol), 1% TIS. Both reactions were run at room temperature, under nitrogen, for two hours. The TFA was evaporated and the resins were precipitated with cold diethyl ether, centrifuged, and washed three times with ether. The mixture was extracted with water (80%)/acetonitrile (19.9%/TFA(0.1%)), centrifuged, and the supernatant lyophilized. The peptides were then purified by reverse-phase HPLC on a preparative C4 column (Vydac, 2.2 cm × 25 cm; 10 μm particle size) and were determined to be at least 95% pure by analytical HPLC and MALDI-TOF (matrix-assisted laser desorption/ionization time-of-flight) mass spectrometry.

CD spectroscopy

All CD measurements were performed on an AVIV 62A DS spectropolarimeter. Samples containing 5–30 μM of peptide were prepared in either 25 mM sodium acetate (pH 4.0) or 25 mM Mops (pH 7.0) and 100 mM NaCl. Stoichiometric amounts of ZnCl2 were also added to selected samples and allowed to equilibrate overnight. All spectra were measured at 25 °C and were an average of six to eight scans. Thermal denaturation studies were performed by averaging the signal for 30 seconds, after five minutes of equilibration, and monitoring the signal at 222 nm between 2 °C and 94 °C.

Sedimentation analysis

Sedimentation equilibrium data were collected at 25 °C on a Beckman XLI analytical ultracentrifuge equipped with both adsorption and interference optics. Peptide concentrations were determined from either the tryptophan (ε280 = 5500 M⁻¹ cm⁻¹) or the tyrosine absorbance (ε277 = 1490 M⁻¹ cm⁻¹). Five samples of isolated A and B peptides and a 1:1 mixture (apo; Fe(II); Zn(II)) were prepared at 100 μM at pH 4.0 and 7.0 in the same buffers as used for the CD experiment. Samples were spun at three speeds (40,000, 45,000, and 48,000 rpm) and the data were analyzed using an in-house fitting routine in IGOR Pro (WaveMetrics, Inc.). Partial specific volumes were estimated from amino acid composition.

Metal ion-binding studies

Co(II) binding was conducted at 36 μM tetramer concentration using a Hewlett Packard model 8453 diode array spectrometer as described. CoCl2 was added in 0.2 eq (Co(II)/protein) increments to a solution of DFTet, and spectral data ranging from 400 to 800 nm were collected. To ensure proper equilibration, individual samples were allowed to equilibrate at room temperature for at least 12 hours.
**Fe(II) oxidation studies**

Kinetic studies of Fe(II) oxidation were conducted at 50 μM protein concentration as described.²

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Computational Design of an A$_2$B$_2$ Diiron Protein


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