X-ray Structure of a Maquette Scaffold

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Maquettes are de novo designed mimics of nature used to test the construction and engineering criteria of oxidoreductases. One type of scaffold used in maquette construction is a four-α-helical bundle. The sequence of the four-α-helix bundle maquettes follows a heptad repeat pattern typical of left-handed coiled-coils. Initial designs were molten globular due partly to the minimalist approach taken by the designers. Subsequent iterative redesign generated several structured scaffolds with similar heme binding properties. Variant [L6F13], a structured scaffold, was partially resolved with NMR spectroscopy and found to have a set of mobile inter-helical packing interfaces. Here, the X-ray structure of a similar peptide ([I6F13M31]2 i.e. [CGGG EIWKL HEEFLKK FEELLKL HEERLKKM]2 which we call L31M), has been solved using MAD phasing and refined to 2.8 Å resolution. The structure shows that the maquette scaffold is an anti-parallel four-helix bundle with “up–up–down–down” topology. No pre-formed heme-binding pocket exists in the protein scaffold. We report unexpected inter-helical crossing angles, residue positions and translations between the helices. The crossing angles between the parallel helices are −5° rather than the expected +20° for typical left-handed coiled-coils. Deviation of the scaffold from the design is likely due to the distribution and size of hydrophobic residues. The structure of L31M points out that four identical helices may interact differently in a bundle and heptad repeats with an alternating [HPPHHPP]/[HPPHHPH] (H: hydrophobic, P: polar) pattern are not a sufficient design criterion to generate left-hand coiled-coils.

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Keywords: four-helix bundle; X-ray crystallography; de novo protein design; MAD phasing

Introduction

In the last decade, de novo protein design has emerged as a constructive methodology for testing our understanding of the relationship between protein sequence, structure and function.1–6 Four-helix-bundle proteins are ubiquitous in nature and serve as convenient scaffolds for the construction of simplified functional versions of complex natural enzymes or molecular maquettes. We have designed a series of maquettes to allow us to study how large multi-heme proteins function in respiratory complexes.7 The prototypical heme maquette, H10H24, is a de novo designed four-helix bundle derived from the cytochrome bc1 respiratory complex and α28 designed by DeGrado and co-workers.7–9 These molecules are excellent models of b-type cytochromes and allow us to probe the engineering tolerances of heme binding and to delineate the role of the various factors in modulating heme electrochemical function.

H10H24 and related maquettes are four-helix bundles consisting of a non-covalent dimer of disulfide linked δ-α-helical subunits, (α-SS-α)2. These sequences are composed mostly of three amino acid types (Glu, Lys and Leu) laid out approximately in an alternating [HPPHHPP]/[HPPHHPH] (H: hydrophobic, P: polar) binary pattern.7,9 Replacement of Leu in the a position by His10, Phe17 and His24 and in the d position by Arg27, was inspired by the sequence of the cytochrome b subunit of the cytochrome bc1 complex.
The incorporation of histidine residues provides for heme binding and the arginine residues were used to modulate the reduction potentials of the heme groups.

Structural studies of the maquettes were initially hampered by the molten globule-like folded states of both the apo and the holo forms. Many of the early de novo designed peptides that simply relied on hydrophobic sequestration to drive the folding and assembly lacked a well-defined tertiary structure. Although contemporary design strategies that can potentially promote structural uniqueness were devised, the early maquettes contained few of those features that would contribute to a native-like structure. The maquette sequences had uniformly large hydrophobic residues (L, I, F) instead of alternating large and small residues in the core, which would promote complementary packing. Constraints, such as carefully placed disulfide bonds, were also absent in the maquette design. The maquettes do have N-terminal disulfide bonds, but they do not influence the aggregation state, packing or conformational specificity of apo maquettes (B.R.G., unpublished results). Metal binding sites have been used to define the structure of bundles. The coordination of histidine residues to heme iron in maquettes constrains modifications to the sequence: L6I and L13F. The incorporation of histidine residues provides for heme binding and the arginine residues were used to modulate the reduction potentials of the heme groups.

Gibney et al. introduced structural specificity into H10H24 with two conservative hydrophobic modifications to the sequence: L6I and L13F. The resulting four-helix bundle scaffold has similar heme binding and redox properties as the prototyple H10H24 but possesses a structured hydrophobic core in the apo state. Although changing Leu to Phe at position 13 of H10H24 resulted in a structured peptide, maquette design does not generally support the notion that the addition of aromatic residues promotes structural uniqueness. Both H10H24 and a maquette variant, [I 6F13] (H10H24, L6I, L13A), have four Phe and four His in the hydrophobic core, but only [I 6F13] is structured.

Dynamic Interfaces Immobilized in Crystal

The structure of the covalently linked (α-SS-α) subunit was solved in solution using NMR spectroscopic methods. The structure of the di-α-helical monomer, (α-SS-α), was well defined using the intra-(α-SS-α) nuclear Overhauser effects (NOEs) with clearly delineated hydrophobic and hydrophilic faces. The lack of observable inter-(α-SS-α) NOEs left the precise hydrophobic core packing scheme undetermined. However, a variety of analytical methods including sedimentation equilibrium analytical ultracentrifugation and 15N relaxation measurements of the molecular re-orientation correlation time indicate that [L,F] is a four-helix bundle in solution. Furthermore, the relative orientation of the two subunits, parallel (syn) or anti-parallel (anti), was ambiguous and was only recently determined to be anti by the chemical attachment of fluorescence probes and rational redesign (B.R.G., unpublished results).

We are uncertain about the range of relative motion between the di-helices at the mobile interface with the available data. However, given that all four helices of the bundle have defined main-chains and side-chains in solution, the relative movements are likely rigid-body motions. The maquette scaffold in solution is thus best described as a four-helix bundle consisting of two uniquely structured di-helices with a "slippery" interface. In the context of the crystalline state, this interface becomes fixed in a unique anti-parallel position. Herein, we present the complete four-helix bundle structure of a maquette scaffold, L31M, solved by X-ray crystallography at 2.8 Å resolution. L31M is a variant of [L,F] with a single C-terminal modification per helix. This leucine to methionine change has minimal affect on the overall structure. The 13C-heteronuclear single quantum coherence (HSQC) spectrum of L31M is nearly identical to the 13C-HSQC of [L,F] (data not shown.) Furthermore, both L31M and [L,F] can be crystallized under the same conditions in the same space groups and unit cells.

Symmetry and Topology of the Four-helix Bundle Maquette Scaffold

The most common topology of four-helix bundles found in the RCSB database is up–down–up–down, in which all helices orient opposite to their adjacent helices. Proteins such as ROP, PDI, Mnt repressor and di-zinc DF all exhibit up–down–up–down topology. In principle, helix dipole and charge–charge interactions can be used to promote anti-parallel up–down–up–down topology. In principle, helix dipole and charge–charge interactions can be used to promote anti-parallel up–down–up–down topology and discourage formation of parallel four-helix bundles. The maquette design did not purposely promote a specific topology. Nonetheless, placing Gln in the b and e positions of the maquette sequence and placing Lys in the c and g positions theoretically destabilizes up–up–down–down conformation and favors the “all up” topology. The arrangement of helices of L31M maquette scaffold is best described as up–up–down–down (Figure 1). A31P ROP mutant and α20 have similar topologies. However, both structures are made of two identical helix-loop-helix units assembled in bisecting-U conformation. The loops connecting the helices in A31P ROP and α20 cross the hydrophobic core diagonally instead connecting adjacent helices in an anti-parallel conformation. In a bisecting-U arrangement, the anti-parallel helices are closer together near the loop and farther apart at the termini. The anti-parallel helices in the maquette scaffold, on the other
hand, do not exhibit the prominent bifurcation of the bisecting-U structures.

L31M has $C_2$ symmetry; helices labeled A and C are symmetrically equivalent as are helices B and D. The three unique interfaces appear in the structure between helix A-helix C, helix B-helix D and helix A-helix B (or its symmetry equivalent helix C-helix D). For clarity, only helices A and B will be discussed, since the same applies to helices C and D. No point group symmetry operator describes the relationship between the two covalently linked di-helices within the monomer subunit. However, the backbones of the A and B helices are approximately related to each other via a 94° rotation plus a 1.96 Å translation about an axis perpendicular to the $C_2$ axis (rmsd = 0.4 Å from residues 6 to 28). Helix A is translated toward the C termini relative to helix B.

The symmetry and unusual topological arrangement of the helices is not a consequence of the disulfide bonding. The methyl region of the $^{13}$C-HSQC of [L$_{13}$F$_{13}$], (four unlinked helices) without the N-terminal cysteine or disulfide is the same as the complete [L$_{13}$F$_{13}$], (two $\alpha$-SS-$\alpha$) suggesting that the disulfide bond is not the determining factor for the up–up–down–down arrangement. Thus, four identical helices with the sequence of L31M or [L$_{13}$F$_{13}$], form four-helix bundles with $C_2$ symmetry and three distinct helix–helix interfaces.

The unusual crossing angles between the helices, which will be discussed later, do not cause the reduction in symmetry either, since the right-handed coiled coil RH$_4$, which has similar helical crossing angles, possesses $C_4$ symmetry. The reason why four identical helices can form non-identical interfaces is that the helices A and B pack with a 1.96 Å relative axial translation. The fact that there is no rotational symmetry axis between the helices A and B excludes the possibility that the interface between the $(\alpha$-$\text{SS}$-$\alpha$) subunits is identical to the interface within the intra $(\alpha$-$\text{SS}$-$\alpha$) subunit.

**Comparison of Crystal Di-helix Unit with the NMR Structure**

In addition to the fact that the mobile interface observed in NMR experiments is fixed in the crystal structure, there are some differences between the NMR structure of [L$_{13}$F$_{13}$], $(\alpha$-$\text{SS}$-$\alpha$) and the X-ray structure of L31M. The differences may be related to the presence of mercury in the crystal structure. The backbone rmsd between L31M and the NMR solution structure of [L$_{13}$F$_{13}$], is 1.23 Å for residues 6–28. The axial translational offset between the helix A and helix B is similar in the crystal and in solution. A majority of the hydrophobic core side-chain conformations are nearly identical between the two structures. However, the core residue side-chain conformations $(\chi$-1) differ at Phe13A, Leu14A, Ile6B, Leu9B, Phe13B, Phe17B, Leu20B and Leu28B. The Phe13 residues are uniformly gauche$(+$) in the X-ray structure while trans in the NMR structure. There are two significant main-chain differences of note between the X-ray and NMR maquette scaffold structures. First, the C terminus in the NMR structure’s helix A turns toward helix B, whereas helix A in the X-ray structure remains straight relative to helix B. The divergence is most noticeable C-terminal to His24A in the NMR structure. Second, the two structures are different between residues six and ten of helix A. The NMR structure evinces a tighter turn than a standard $\alpha$-helix, reminiscent of a $3_{10}$-helix, while the crystal structure remains $\alpha$-helical to Ile6 of helix A.

**The Four-helix Bundle Maquette Scaffold is not a Left-handed Coiled-coil**

There are several structures available of designed proteins with left-handed coiled-coil architectures that were based on the heptad repeat sequence pattern. All of these helical bundles have crossing angles near those observed in the natural left-handed coiled-coils ColE1 ROP and yeast GCN4.

As designed, residues 5–31 are helical as previously shown by NMR spectroscopy. Table 1 summarizes the helical bundle parameters resulting
from fitting to the Crick parameterization with the addition of two parameters that account for the helix–helix translation (values not shown). The average helix–helix separation within the (α-SS-α) subunit is about 10.8 Å, which is slightly larger than those observed in a coiled-coil such as ROP: 9.2 Å intramonomer and 10.1–10.3 Å intermonomeric distance. The crossing angle between the two disulfide-linked helices in the (α-SS-α) subunit is −5°, while the crossing angle between the two symmetrically related subunits is +178°. The crossing angle between helix A and helix B is similar to that of right-handed coiled-coil RH4, designed with a 11-fold or undecadecadal helix B is similar to that of right-handed coiled-coil. The number of residues/turn is 3.6, which is consistent with α-helix and not the 3.5 of a left-handed coiled-coil. The helical backbone, including the CB atoms, matches closely to that of a straight polyalanine helix (rmsd 0.33 Å for helix A, and 0.4 Å for helix B). The difference between a 3.6 residues/turn pitch of the α-helix and the 3.5 residues/turn pitch of a coiled-coil is apparent when attempts are made to describe L31M as a coiled-coil with a heptad repeat sequence. The positions of the CB atom within the context of the bundle, deviate markedly from the positions expected. Based on the heptad repeat design strategy, H10, F17 and H24 are expected to be in the heptad α position. However, in the structure H10 is in a heptad d position, F17 occupies a location between the heptad a and d positions and H24 occupies a d position. Also, expected heptad d position residues F13, L20 and R27 are situated near theoretical heptad g positions. Clearly, as one progresses down the helix the difference between a straight α-helix and the expected coiled-coil become more apparent. In addition to the larger number of residues per turn, the mismatch of residue positions contributes to the clockwise crossing between helix A and helix B and near-parallel arrangement between the symmetry related helices (A and C; B and D).

Table 1. L31M mercury MAD experiment statistics

<table>
<thead>
<tr>
<th>Unit cell</th>
<th>26.9 Å 48.8 Å 46.7 Å</th>
</tr>
</thead>
<tbody>
<tr>
<td>Space group</td>
<td>P2₁(1)</td>
</tr>
<tr>
<td>Wavelength (Å)</td>
<td>1.0019 1.0098 0.9824</td>
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<tr>
<td>Resolution (Å)</td>
<td>20–2.7 20–2.7 20–2.7</td>
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<tr>
<td>Redundancy</td>
<td>1.76 1.7 1.7</td>
</tr>
<tr>
<td>I/σ</td>
<td>28.4 27.2 25.9</td>
</tr>
<tr>
<td>Completeness* (%)</td>
<td>97.1 (88.6) 96.4 (85) 96.1 (80.6)</td>
</tr>
<tr>
<td>Rmerge* (%)</td>
<td>2.8 (7.2) 2.9 (7.2) 3.1 (8.5)</td>
</tr>
<tr>
<td>Figure of merit*</td>
<td>0.71 (0.54)</td>
</tr>
</tbody>
</table>

* Numbers in parentheses are that of the last resolution ring.

The Maquette Scaffold Does not Exhibit Knob-in-hole Packing

Left handed coiled-coils employing the heptad repeat sequence pattern are expected to exhibit knob-in-hole packing. The −5° interhelical crossing angle found in the L31M structure cannot support systematic knob-in-hole packing which is optimal at a crossing angle of +20°. Helices packed with crossing angles near 0° have overlapping Cα positions between many residues in a helical-net diagram. At the crossing angles observed in L31M, the Cα positions of residues from different coiled-coil helices overlap, preventing formation of a regular packing lattice. With certain amino acid sequences, residues from the same helix may pack with each other instead of with residues of another helix. This creates the non-interdigitated packing at the helix B–helix D interface of L31M as shown in Figure 3. Some side-chain interdigitation is observed at the A–B (and symmetry-related C–D) interface (Figure 2) and A–C interfaces (Figure 3) of L31M. The B–D interface of L31M shows little interdigitation. In contrast, knob-in-hole packing, like that observed in GCN4, shows extensive interdigitation at the packing interface. A non-interdigitated packing interface may lack structural specificity due to shortage of mutual lateral restrictions, which would have been provided by the interdigitated side-chains. It is possible that inadequate interlocking of residues may be one of the causes of the lack of structural specificity seen in early maquette scaffold designs. The lack of interdigitation between
the helices B and D of L31M may also cause the interface to be slippery, leading to the absence of NOEs at the B–D interfaces.

Concluding Remarks

Structural determination of L31M completes the first design and assessment cycle of a maquette scaffold. We learned that helices with crossing angles near 0° must be considered in the design especially when the sequence contains many large hydrophobic residues. We also realize that symmetry in the design clearly depends on whether or not the most stable conformation preserves the structural identity among helices. Identical helices need not generate identical interfaces. Furthermore, not all interfaces need to be extensively interdigitated for the structure to be unique in crystalline state. Careful adjustment of the residues at the interface can alter considerably the association at the interfaces. The design history of [I6F13] and L31M suggests that one may iteratively fine-tune the packing at each interface. The L6I and L13F amino acid changes that are responsible for the conformational specificity of [I6F13] are found at the A–B (C–D) and A–C interfaces but not the B–D interface; the A–B, C–D and A–C interfaces have more observed interdigitation (Figs. 2 and 3) than the B–D interface (Figure 3). With modifications at positions such as 14, 21 and 28, which are at the B–D interface, the amount of interdigitation between the helices may increase, leading to the achievement of structural specificity between the di-helices.

Material and Methods

Crystallization and phasing

We replaced L31 in [I6F13] with methionine early in the study to gain the option of selenomethionine multiple anomalous dispersion (MAD) studies. The sequence of L31M is (Ac-CGGG EI 6WKL-H10EEF13LKK-FEELLKL-FEERLKK-M31-CONH2). The N termini are acetylated and the C-termini are amidated. Synthesis and purification of the peptides are detailed elsewhere. Both L31M and [I6F13] can be crystallized under the same conditions in the same unit cell of a = 26.9 Å, b = 48.8 Å, c = 46.7 Å, 

![Figure 2](image2.png)

**Figure 2.** A slab through the center of packing interfaces between the covalently linked di-helices (RASMOL). The helices are represented by the color coding scheme in Figure 1. The red circles are mercury atoms. The diagram on the right shows a scheme of packing interface with interdigitation. An interdigitated interface in this cross-sectional representation has juxtaposition of different colors. Note, the lighter blue and orange represent the “sliced atoms” of the blue and orange helices at the cross-section.

![Figure 3](image3.png)

**Figure 3.** A slab through the center of packing interfaces between the symmetry-related helices (RASMOL). The interface between the symmetry-related B–D interface, shown in orange and blue, has fewer interdigitations than the interface between helix A and helix C. The scheme on the right shows a simplified packing arrangement between helix B and helix D illustrating the lack of interdigitation between the helices when compared to the scheme in Figure 2. Note, the lighter gray and blue represent the “sliced atoms” of the gray and blue helices at the cross-section.

<table>
<thead>
<tr>
<th>Regions of fitting</th>
<th>Superhelical residues/turn (2π/ω0)</th>
<th>Superhelix radius (R0) (Å)</th>
<th>R1 α-helix radius (R1) (Å)</th>
<th>Superhelix crossing angle (deg.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NMR 17–27</td>
<td>235</td>
<td>5.287</td>
<td>2.32</td>
<td>5.5</td>
</tr>
<tr>
<td>X-ray 17–27</td>
<td>285</td>
<td>5.407</td>
<td>2.25</td>
<td>−4.5</td>
</tr>
<tr>
<td>X-ray 6–17</td>
<td>222</td>
<td>5.25</td>
<td>2.27</td>
<td>−5.6</td>
</tr>
<tr>
<td>Rise per residue (D)</td>
<td>Residues/helical turn (2π/ω1)</td>
<td>RMSD of fitting (Å)</td>
<td></td>
<td></td>
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<tr>
<td>NMR 17–27</td>
<td>1.47</td>
<td>3.69</td>
<td>0.46</td>
<td></td>
</tr>
<tr>
<td>X-ray 17–27</td>
<td>1.50</td>
<td>3.58</td>
<td>0.25</td>
<td></td>
</tr>
<tr>
<td>X-ray 6–17</td>
<td>1.513</td>
<td>3.59</td>
<td>0.27</td>
<td></td>
</tr>
</tbody>
</table>

Crossing angle = ArcSin(R0/ω0/d)
α = 90.0°, β = 104.7°, γ = 90.0° in P2₁, space group. The optimal crystallization samples are HPLC pure L31M (α-S-Sα) peptides dialyzed against deionized water and concentrated to 0.6 mM four-helix bundle. The crystals were formed from hanging-drop over a reservoir containing 30% (w/v) PEG2000 monomethyl ether and 0.1 M Hepes at pH 7.4. The crystallization drop itself contained a mixture of 2.5 peptide/reservoir. Crystals emerged within one to three days at 28 °C and were grown to 200 μm × 50 μm over an additional 24 hours. The crystals were conditioned with cryoprotectant (30% PEG550 buffered with 30 mM Hepes at pH 7.4) before they were frozen in a liquid nitrogen stream for storage or for data collection. For heavy atom derivitization, the crystals adapted to the cryoprotectant were soaked in a solution consisting of 30 mM ethylmercuryphosphate in 30% PEG550 for 12 hours before freezing. A MAD data set was collected at Beamline X12C at the National Synchrotron Light Source on a Brandeis CCD detector at the mercury LIII edge (1.0019 Å), the inflection point (0.9824 Å), and a remote wavelength (1.0098 Å). Data integration and reduction were performed with DENZO.47 The initial phase was calculated with SOLVE39 and solvent flattened with RESOLVE.44,45 Structural analysis and verification were performed with PROCHECK.46

Refinement

The backbone of the four-helix bundle is visible from residues 3–31 with the Ca positions of these residues clearly defined. However, many solvent-exposed amino acid side-chains, particularly Glu and Lys, are not visible beyond their Cβs. Mercury density is located within the hydrophobic core near the His24s. Model building used O47 followed by refinement with CNS.46 Refinements that minimize the standard crystallographic residual did not converge because the data (unique reflections) to variable number of non-hydrogen atoms multiplied by four) ratio is only about 0.7 at 2.8 Å. However, refinements were successful when phase restraints and non-crystallographic symmetry (NCS) restraints were applied simultaneously. The phase restraint is applied by using the “mixed” target for refinement. The final R = 23.4% and Rp = 27.8%. The native data sets of L31M, [L31M], and other maquette crystals (S.S.H., unpublished results) have not been refined to satisfaction because there is no additional phase information at this time due to lack of high resolution data and phase restraints. Table 2 summarizes the MAD experiment and the refinement statistics for L31M.

Within the crystal structure there is a questionable χ-1 torsion angle: Phe17 of the B helix. The density to which the structure was fitted is incomplete, and the phenyl ring has high temperature factors (B > 40 Å²). The χ-1 angle of Phe17β in the structure is 230°, which is mid-way between two favorable angles (183° and −66°), suggesting that Phe17β exists in two equally favorable conformations. The fitted Phe17β conformation is likely the average of positions of Phe17β. Note that Phe17 of the helix B in the NMR structure is in the allowed region of trans conformation with χ-1 approximately −160° (or 200°).

Atomic coordinates

The coordinates have been deposited into the RCSB database with accession code PDB ID 1M3W.

Acknowledgements

This work was supported by grants from the National Institutes of Health (GM 48130 and GM 41048 to P.L.D.; Cell and Molecular Biology Training Grant to S.S.H.). The MAD data set was collected during the RapiData 2000 course at Brookhaven National Laboratory. Mass spectroscopic analyses were performed by the Protein Chemistry Laboratory of the University of Pennsylvania.

References


Edited by D. Rees

(Received 5 August 2002; received in revised form 2 December 2002; accepted 5 December 2002)