Engineering Oriented Heme Protein Maquette Monolayers through Surface Residue Charge Distribution Patterns

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We have designed and synthesized four-α-helix-bundle proteins that accommodate heme groups to act as molecular “maquettes” of more complex natural electron-transfer proteins. These bundles can be oriented at an air—water interface and transferred onto solid surfaces to facilitate the exploration of the factors that govern biological electron transfer. We find that the orientation of these maquettes on an air—water interface can be controlled by choosing the distribution of charged amino acids along the sides of the helices exposed to water. The four α-helices were assembled either as two subunits, where each subunit consists of two α-helices linked by a terminal cysteine disulfide bond, or as a single, four-helix covalent unit consisting of two helix-loop-helix molecules linked by a terminal cysteine. In either case, when each α-helix contains both positively charged lysines and negatively charged glutamates, addition of the heme binding bundles to an air—water interface causes them to open up and lie on the surface with α-helical axes oriented parallel to the interface. In contrast, when the positive and negative charges are segregated on different helices (two negative, two positive) of the single covalent four-α-helix-bundle unit, the bundle preserved its integrity on transfer to the air—water interface. Moreover, the presence of heme dictates the orientation of the α-helical axes of the bundle with respect to the surface plane. The α-helices adopt a parallel orientation in the absence of heme and a perpendicular orientation in the presence of heme. Circular dichroism (CD) and ultraviolet—visible (UV—vis) spectroscopy supported by linear dichroism demonstrate that these molecular orientations are preserved in Langmuir—Blodgett monolayer films on solid substrate surfaces.

Introduction

A novel approach to the study of complex natural oxidoreductase enzymes is to design and synthesize proteins with the minimal requirements for the assembly and incorporation of active redox cofactors. These de novo designed1–3 minimalist redox proteins act as molecular “maquettes” to uncover the engineering behind the assembly of natural oxidoreductases that promote directed, long-range electron transfer.5,5 We choose sequences strongly favoring α-helical folding to present hydrophobic residues along one face of each helix and charged groups on the opposite face. Hydrophobic forces drive association of the helices into a four-α-helix bundle in solution, while histidines in the bundle core ligate added hemes. These heme protein maquettes have proven to possess the characteristic electrochemical and optical properties typical of natural h-type cytochromes.6–10

Maquettes can be engineered to organize on surfaces by exploiting the great freedom of de novo design and chemical synthesis to modify exposed residues and control the interactions between α-helices. Organizing maquettes on surfaces11,12 potentially not only facilitates electrochemical studies,13–15 which explore the factors that govern biological electron tunneling, but also forms the foundation for translating these heme protein maquettes into novel functional biomaterials.16–18 Furthermore, study of the surface assembly of these heme protein maquettes provides information on the nature of the various interactions between proteins and membranes or artificial surfaces.19

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In a previous study,12 we explored the Langmuir—Blodgett (LB) film properties of our prototype four-α-helix-bundle heme-binding protein maquettes. We showed that this homodimeric prototype heme protein (α-ss-α)2 shown in Figure 1A dissociates on the air—water interface into its monomeric subunits, α-ss-α, and orients its α-helical axes parallel to the surface, presumably with the hydrophobic interior of the original four-α-helix bundle exposed to the air. We also showed that the addition of hydrophobic tails, such as palmitic acid, to the N-termini of the (α-ss-α)2 heme protein induces reorientation at high surface pressure such that the α-helices adopt an angle perpendicular to the air—water interface. Encouraged by the observation that the polarity distribution on the exterior of the bundles can influence the structures on the air—water interface, we have focused on the effect of altering exterior charge distributions; three related synthetic heme protein maquettes are compared.

1. (α-ss-α)2. The (α-ss-α)2 heme protein has been described in the previous study.12 It is a highly water-soluble four-α-helix-bundle heme protein that is based on the prototype described by Robertson et al.,6 conservatively modified to improve packing of the bundle interior20 (see Materials and Methods). In the bundle, a pair of identical α-helices are linked by a disulfide bond at their cysteiny1 N-termini to make an α-ss-α subunit. A pair of α-ss-α subunits spontaneously assemble in aqueous solution to form a noncovalent four-α-helix bundle (α-ss-α)2. On one side of each α-helix there are seven glutamates, six lysines, and one arginine forming alternate positive and negative patches, which form a candy cane striped pattern on the exterior...
which the disulfide bonds are at the same or opposite ends of the four-helix bundle. 24,25

Figure 1. Design and construction of heme protein maquettes. Part A shows the construction of heme$_2$(α-ss-α)$_2$; the solid-phase peptide synthesis produced the 31-amino-acid peptide α, which is oxidized to form a dihelic unit α-ss-α. Two α-ss-α units self-assemble into a four-helix bundle (α-ss-α)$_2$. The four-helix bundle coordinates hemes by bis-histidyl ligation to form the heme protein maquette heme$_2$(α-ss-α)$_2$. Part B shows the construction of heme$_2$(α-l-α$_+$)$_2$; the solid-phase peptide synthesis produced the 63-amino-acid peptide α-l-α$_+$, which is oxidized to form the four-helix bundle (α-l-α$_-$)$_2$. The four-helix bundle coordinates hemes by bis-histidyl ligation to form the heme protein maquette heme$_2$(α-l-α$_-$)$_2$. Part C shows the construction of heme$_2$((α$^+$-l-α$^-$-s)$_2$); the solid-phase peptide synthesis produced the 63-amino-acid peptide α$^+$-l-α$^-$ (with lysines and glutamates segregated on different helices), which is oxidized to form the four-helix bundle (α$^+$-l-α$^-$-s)$_2$. The four-helix bundle coordinates hemes by bis-histidyl ligation to form the heme protein maquette, heme$_2$((α$^+$-l-α$^-$-s)$_2$).

of the assembled four-α-helix bundle. Figure 1A shows that the four-α-helix bundle can adopt a syn or anti topology in which the disulfide bonds are at the same or opposite ends of the bundle, depending on the sequence and cofactor binding.21-23 The structure of this four-α-helix bundle has been studied by NMR spectra.24,25

2. (α-l-α$_-$-s)$_2$. The amino acid sequence of each α-helix of the (α-l-α$_-$-s)$_2$ heme protein is identical to that of the (α-ss-α)$_2$ heme protein. However, as shown in Figure 1B, two repeating α-helix amino acid sequences with six glycines between are synthesized as a single sequence. A pair of single sequence α-l-α$_-$ are linked by a disulfide bond at their cysteiny1 N-termini to form a four-α-helix bundle (α-l-α$_-$-s)$_2$. The four-helix bundle coordinates hemes by bis-histidyl ligation to form the heme protein maquette, heme$_2$((α-l-α$_-$-s)$_2$).

3. (α$^+$-l-α$^-$-s)$_2$. The amino acid sequence of the (α$^+$-l-α$^-$-s)$_2$ heme protein is a variant of the (α-l-α$_-$-s)$_2$ heme protein with a major change in the distribution of the exterior charged residues compared with (α-l-α$_-$-s)$_2$ and (α-ss-α)$_2$. In (α$^+$-l-α$^-$-s)$_2$, 14 glutamates are placed in one helix and 14 lysines in the other helix to form a highly asymmetrically charged α$^+$-l-α$^-$, as shown in Figure 1C. Then, as with the (α-l-α$_-$-s)$_2$, a pair of α$^+$-l-α$^-$ are linked by a disulfide bond at their cysteiny1 N-termini to form a four-α-helix bundle (α$^+$-l-α$^-$-s)$_2$.

This report explores the properties of the three heme protein maquettes in a Langmuir–Blodgett film balance and after deposition on planar substrate surfaces. We show that the (α-ss-α)$_2$ and (α-l-α$_-$-s)$_2$ heme proteins display similar properties on the air–water interface while the (α$^+$-l-α$^-$-s)$_2$ heme protein is dramatically different. It is clear that the segregated charge distribution of (α$^+$-l-α$^-$-s)$_2$ increases the four-α-helix-bundle integrity through strong electrostatic interactions between the helices and provides a basic maquette design adapted for both solutions and interfaces.

Materials and Methods

Peptide Sequences. 1. (α-ss-α)$_2$. The amino acid sequence of each α-helix in the (α-ss-α)$_2$ heme protein is

Ac-CGGG-EIWKLH-EEFLKKF-EELKLF-EEFLKLF-KKKLKLK-CN

Ac-CGGG-EIWKLH-EEFLKKF-EELKLF-KKKLKLK-CN

Histidine at position 10 is for heme binding. A pair of identical α-helices are linked by the disulfide bond at their cysteiny1 N-termini to make an α-ss-α subunit. Two α-ss-α subunits spontaneously assemble in aqueous solution to form a four-α-helix bundle (α-ss-α)$_2$. This peptide differs from the prototype peptide6 H10A24 in the interior residues L6I, L13F, and A24F and, as shown by NMR, has a singular structure.25

2. (α-l-α$_-$-s)$_2$. The amino acid sequence of the α-l-α in the (α-l-α$_-$-s)$_2$ heme protein is

Ac-CGGG-EIWKLH-EEFLKKF-EELKLF-EEFLKLF-KKKLKLK-CN

A pair of single sequence α-l-α are linked by a disulfide bond at their cysteiny1 N-termini to form a four-α-helix bundle (α-l-α$_-$-s)$_2$.

Ac-CGGG-EIWKLH-EEFLKKF-EELKLF-EEFLKLF-KKKLKLK-CN

3. (α$^+$-l-α$^-$-s)$_2$. The amino acid sequence of the α$^+$-l-α$^-$ in the (α$^+$-l-α$^-$-s)$_2$ heme protein is

Ac-CGGG-EIWKLH-EEFLKKF-EELKLF-EEFLKLF-KKKLKLK-CN

A pair of α$^+$-l-α$^-$ are linked by a disulfide bond at their cysteiny1 N-termini to form a four-α-helix bundle (α$^+$-l-α$^-$-s)$_2$.

Chemicals and Solvents. Hemin, trifluoroacetic acid (TFA), dimethyl sulfoxide (DMSO), and octadecyltrichlorosilane were purchased from Aldrich Chemical Co. (Milwaukee, WI). Ethanedithiol and 1-hydroxybenzotriazole were purchased from Fluka (Ronkonkoma, NY). FMOC-protected amino acid perfluorophenyl esters were purchased from PerSeptive Biosystems (Framingham, MA) with the exception of FMOC-L-Arg(Pmc)-OPfp, which was obtained from Bachem (King of Prussia, PA). NovaSyn PR-500 resin was purchased from Calbiochem-Novabiochem (La Jolla, CA). All other chemicals and solvents were reagent grade. Water was purified using a Milli-Q water system from Millipore Corp. (Bedford, MA).

Peptide Synthesis. The peptides were synthesized on a continuous-flow Milligen 9050 solid-phase synthesizer using standard FMCOC-2Bu protection strategy with NovaSyn PR-500 resin at 0.2 mmol scale. The side chain protecting groups used are as follows: Lys(tBoc), Glu(OtBu), Cys(Trt), and Arg(Pmc). The peptides were cleaved from the resin and simultaneously deprotected using 90:8:2 trifluoroacetic acid/ethanedithiol/water for 2 h. Crude peptides were precipitated and triturated with cold ether, dissolved in water (0.1% TFA), lyophilized, and purified to homogeneity by reversed-phase C18 HPLC using aqueous acetonitrile gradients containing 0.1% (v/v) TFA. The resulting peptide identities were confirmed with laser desorption mass spectrometry.
Heme Protein Maquette Preparation. 1. Heme2-(α-ss-α)2. The 31-amino-acid peptides were dissolved in 100 mM ammonium hydrogen carbonate buffer (pH 9.2) in air to allow the cysteines to oxidize to form the 62-amino-acid peptide α-ss-α. After 4 h the solution was frozen and lyophilized. When needed, the α-ss-α peptides were dissolved in phosphate buffer (50 mM phosphate, 100 mM NaCl, pH 8.0) to yield a final concentration of 0.02 mM. Heme was incorporated into (α-ss-α)2 from a stock solution of 4 mM of Fe(III) protoporphyrin IX (heme) in DMSO by successive additions of 0.1 heme per binding site until 1 heme per bishistidine site was reached. During each addition, the solution was well stirred and then allowed to equilibrate for 5 min. The final concentration of DMSO in the aqueous solution was always lower than 1:100 (v/v). The heme incorporation was monitored by the increase of the Soret band maximum between 411 and 412 nm (ε = 120 000 M⁻¹ cm⁻¹).

2. Heme2-(α-l-α-s)2. The 63-amino-acid α-l-α-s peptides were dissolved in 100 mM ammonium hydrogen carbonate buffer (pH 9.2) in air to allow the cysteines to oxidize to form the 126-amino-acid peptide (α-l-α-s)2. After 4 h the solution was frozen and lyophilized. When needed, (α-l-α-s)2 peptides were dissolved in phosphate buffer (50 mM phosphate, 100 mM NaCl, pH 8.0) to yield a final concentration of 0.02 mM (α-l-α-s)2. Heme was incorporated into (α-l-α-s)2 to form heme2-(α-l-α-s)2 from a stock solution of 4 mM of Fe(III) protoporphyrin IX (heme) in DMSO by successive additions of 0.1 heme per binding site until 1 heme per bishistidine site was reached. During each addition, the solution was well stirred and then allowed to equilibrate for 5 min. The final concentration of DMSO in the aqueous solution was always lower than 1:100 (v/v). The heme incorporation was monitored by the increase of the Soret band maximum between 411 and 412 nm (ε = 120 000 M⁻¹ cm⁻¹).

3. Heme2-(α+-l-α-s)2. The 63-amino-acid α+-l-α-s peptides were dissolved in high ionic strength phosphate buffer (50 mM phosphate, 1.5 M NaCl) at pH 4.4 to yield a final concentration of 0.04 mM heme2-(α+-l-α-s)2. Heme was incorporated into (α+-l-α-s)2 to form heme2-(α+-l-α-s)2 following the same procedure described above.

Solution Molecular Weight Determination. Size exclusion chromatography was performed on a Beckman System Gold HPLC system with a diode array detector using a Supelco Sigmachrom GFC-100 column (300 mm × 57.5 mm) eluted with aqueous buffer. The column was standardized using aprotinin (6.5 kDa), ribonuclease A (13.7 kDa), chymotrypsinogen (25.0 kDa), ovalbumin (43.0 kDa), and bovine serum albumin (67.0 kDa).

Langmuir Film Preparation and Isotherm Measurement. A Langmuir—Blodgett film balance (Lauda Filmbalance FW2; Sybron/Brinkmann, Westbury, NY) was used to make the films according to a general procedure described elsewhere.26,27 The aqueous subphase contained 1 mM phosphate at pH 8.0. All films were made under an argon atmosphere at a subphase temperature of 22 °C. Proteins were spread at the air–water interface using the “glass-rod” method.28 This involved placing a 0.6 mm diameter glass rod at a small angle (5°) relative to the trough plane, just touching the water surface, to which 2 mL aliquots of material were added every 10–15 s. During this procedure the trough surface area was held constant. Surface pressure—area isotherms were measured using a barrier speed of 10 cm/min.

Substrate Preparation. Quartz slides, 1 mm in thickness (Esco Products, Oak Ridge, NJ), were sonicated first in detergent solution and then in water. This was followed by immersion in a freshly prepared solution of 4:1 (v/v) 95–98% sulfuric acid/30% hydrogen peroxide (CAUTION: This solution is highly oxidizing and very acidic and should be handled with extreme care) at room temperature for about 1 h. The slides were then thoroughly rinsed with water and finally stripped of visible surface water with a stream of argon. The cleaned slides were then rendered hydrophobic by silanation with octadecyltrichlorosilane. This was achieved by sonicating the slides for 20 min in a 0.1% octadecyltrichlorosilane in a solution of 80% hexadecane (99% pure), 12% carbon tetrachloride, and 8% chloroform. The slides were washed three times in chloroform for at least 5 min for each wash. The quality of the hydrophobic coating was checked by placing a 20 mL droplet of water on the slide. The coating was considered satisfactory if the droplet started moving freely and continuously on the slide when the tilt angle of the glass exceeded ~40°.

Langmuir—Blodgett Monolayer Film Preparation. Protein maquettes were spread as discussed above, compressed, and held at a constant surface pressure of ~20 mN/m. A hydrophobically modified quartz slide was passed vertically into the subphase at 3 mm/min. The submerged slide was then released into a vial resting at the bottom of the trough. After LB deposition, the slide was transferred from the vial to a quartz cuvette for spectroscopic characterization. The entire process was done in buffer so that the slide was never exposed to air.

Circular Dichroism Spectropolarimetry. Circular dichroism spectra were measured in an Aviv 62DS circular dichroism spectropolarimeter. Rectangular quartz cuvettes of 0.2 cm path length were used for measuring solution spectra. For film spectra, four quartz slides coated with LB film were positioned in a quartz cuvette of 1.0 cm path length using two Teflon blocks (each has four parallel slits and fits tightly on the bottom and top of the cuvette) holding the slides parallel to each other. The slides were immersed in buffer and oriented perpendicularly to the incident measuring beam. A baseline reference was recorded by using four blank, cleaned quartz slides positioned in the same way.

Ultraviolet—Visible Absorption Spectroscopy and Linear Dichroism. UV—vis absorption spectra were recorded with a Perkin-Elmer Lambda 2 UV—vis spectrophotometer. Rectangular quartz cuvettes of 0.2 and 1.0 cm path lengths were used for measuring solution spectra. For film spectra, quartz slides coated with LB film were positioned in a quartz cuvette of 1.0 cm path length using a Teflon block (which fits tightly on the bottom of the cuvette and has one slit at 0°, 30°, 45°, or 60° angle relative to the edges of the block) holding the slide at a selected angle with respect to the incident measuring beam. The slides were always immersed in buffer. For linear dichroism measurements, a UV dichroic polarizer from Oriel Corporation (Stratford, CT) sensitive from 230 to 770 nm was placed between the incident beam and the coated slide. In all cases a baseline reference was recorded by using a blank, cleaned quartz slide.

Results

1. Solution Characterization of Heme2-(α+-l-α-s)2 Maquette. The (α+-l-α-s)2 protein maquette was found to have very different solubility properties from the (α-ss-α)2 and (α-α-α-s)2 protein maquettes, evidently resulting from its highly asymmetric charge distribution. At ionic strengths up to 1 M, (α+-l-α-s)2 solutions are highly scattering in the pH range from 4.5 (pKa of glutamate) to 10.2 (pKa of lysine), as shown in parts A and B of Figure 2. However, solubility within this range increases with increasing ionic strength (parts C and D of Figure 2). Thus, at an ionic strength of 1.5 M, the protein can be readily solvated even at neutral pH, without causing precipitation. This is done by exposing the protein to pH 4.4
buffer with 1.5 M NaCl followed by raising the pH to 8.0 (Figure 2D). Heme can be incorporated into the maquette in the high ionic strength solution thus prepared. The apparent molecular weights at high ionic strength determined by gel filtration chromatography were 17.3 and 18.2 kDa before and after incorporation of hemes, which are very similar to the (αss-α2) and heme2-(αss-α2) maquettes, and consistent with a four-helix-bundle aggregation state for both (α1-α1-α1-α1)2 and heme2-(α1-α1-α1-α1)2. Figure 3A shows that heme2-(α1-α1-α1-α1)2 has a CD spectrum typical of a highly α-helical structure with minima at 208 and 222 nm, and the helical content (75%) is similar to those of heme2-(α1-α1-α1-α1)2, and heme2-(α1-α1-α1-α1)2 maquettes. Figure 3B shows that heme2-(α1-α1-α1-α1)2 was soluble in 200 mM NaCl buffer at pH = 4.4, and some aggregation was observed when the pH was adjusted to 8.0. Part D shows that (α1-α1-α1-α1)2 was soluble in 1.5 M NaCl buffer at pH = 4.4, and no aggregation was observed when the pH was adjusted to 8.0.

2. Characterization of the Maquettes on Air–Water Interface. The pressure–area isotherms of (αss-α2), (α1-α1-α1-α1)2, and (α1-α1-α1-α1)2 without heme are compared in Figure 4A, and the pressure–area isotherms of the helix-containing heme2-(αss-α2), heme2-(α1-α1-α1-α1)2, and heme2-(α1-α1-α1-α1)2 are compared in Figure 4B. The isotherms of all three maquettes share the characteristics that when the films were compressed to approximately 30 mN/m, a transition from gaseous state to liquid expanded state of the monolayer occurs; beyond the transition point, the surface pressure increases only to a small extent per unit decrease of surface area. The surface area per molecule at the transition point provides a measure of the molecular area for orientations found at low surface pressure, where π is the surface pressure, A the surface area per α-helix, A0 the limiting area per α-helix, k the Boltzmann constant, T the temperature in kelvin, and n the number of α-helices per independent particle. Figure 4D, kTπ/n values are plotted against A for the low-pressure region (from 1 to 4 mN/m) of each isotherm. The slopes of the fitted lines give the value of n, and the x-intercepts of the lines give the value of A0. We compare this area with the dimensions of a single helix derived from an NMR structure,24 which has a helical diameter of 12–13 Å, for a small cross section of 144–170 Å2, and a helical length of 50 Å, for a lengthwise cross section of about 600 Å2.

1. (αss-α2). Trace a in Figure 4A shows the isotherm of (αss-α2), and Figure 4D reveals that its n value is 2.09, which indicates that two α-helices are associated in the independent particles on the air–water interface. Figure 4D also reveals that the A0 for the (αss-α2) film is 593 Å2/α-helix, which indicates that the axes of the α-helices are parallel to the air–water interface. Thus, the four-α-helix bundle appears to be dissociated into a pair of αss-α subunits that are lying flat on the air–water interface, presumably with all polar sides in contact with the water surface and hydrophobic faces exposed to the air.

Trace b in Figure 4B shows the isotherm of heme2-(αss-α2), and Figure 4D reveals that its n value is 2.05 and A0 value is 602 Å2/α-helix, indicating that the maquette both with and without heme exhibits the same behavior on the air–water interface.

2. (α1-α1-α1-α1)2. Trace c in Figure 4A shows the isotherm of (α1-α1-α1-α1)2, and Figure 4D reveals that its n value is 4.06, which indicates as expected that the four α-helices are associated in

Figure 2. Solubility of (α1-α1-α1-α1)2 at different pH and ionic strength, demonstrated by UV spectra of pM peptide solution. A clean tryptophan spectrum with a flat baseline near 400 nm indicates good solubility, while a broadened tryptophan spectrum with a slanted baseline indicates aggregation in the solution (also visible by looking through the solution). Part A shows that (α1-α1-α1-α1)2 was soluble in 200 mM NaCl buffer at pH = 4.4 (where many glutamates are neutral) but aggregated when the pH was adjusted to 5.0. Part B shows that (α1-α1-α1-α1)2 was soluble in 200 mM NaCl buffer at pH = 10.5 (where many lysines are neutral) but aggregated when the pH was adjusted to 10.2. Part C shows that (α1-α1-α1-α1)2 was soluble in 1 M NaCl buffer at pH = 4.4, and some aggregation was observed when the pH was adjusted to 8.0. Part D shows that (α1-α1-α1-α1)2 was soluble in 1.5 M NaCl buffer at pH = 4.4, and no aggregation was observed when the pH was adjusted to 8.0.

Figure 3. Characterization of heme2-(α1-α1-α1-α1)2 in 10 mM phosphate, 1.5 M NaCl, pH 8.0 buffer. Part A shows a comparison of CD spectra of heme2-(α1-α1-α1-α1)2 (open circles) and heme2-(αss-α2) (solid line). For heme2-(α1-α1-α1-α1)2, the CD spectropolarimeter was saturated at wavelengths below 200 nm because of the high absorption of UV light by the high concentration salt. Part B shows the UV–vis absorption spectrum of heme2-(α1-α1-α1-α1)2.

In the low surface pressure gaseous state, a two-dimensional version of the ideal gas law applies:29

$$\pi(\Delta A_{0}) = kTln$$

where π is the surface pressure, A the surface area per α-helix, A0 the limiting area per α-helix, k the Boltzmann constant, T the temperature in kelvin, and n the number of α-helices per independent particle. In Figure 4D, kTπ/n values are plotted against A for the low-pressure region (from 1 to 4 mN/m) of each isotherm. The slopes of the fitted lines give the value of n, and the x-intercepts of the lines give the value of A0. We compare this area with the dimensions of a single helix derived from an NMR structure,24 which has a helical diameter of 12–13 Å, for a small cross section of 144–170 Å2, and a helical length of 50 Å, for a lengthwise cross section of about 600 Å2.

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2. (α1-α1-α1-α1)2. Trace c in Figure 4A shows the isotherm of (α1-α1-α1-α1)2, and Figure 4D reveals that its n value is 4.06, which indicates as expected that the four α-helices are associated in
faces down and hydrophobic faces to the air as discussed for \((\alpha ss-\alpha ss)\) maquettes.

Trace d in Figure 4B shows the isotherm of heme2-(\(\alpha l-\alpha l-\alpha s-\alpha s\)), and Figure 4D reveals that its \(n\) value is 3.89. It also indicates that this high surface pressure region does not lend itself to analysis by eq 1). The surface area per \(\alpha l\) helix when it stands perpendicular to the air–water interface. This analysis indicates that the four-\(\alpha l\)-helix-bundle structure of heme2-(\(\alpha l-\alpha l-\alpha s-\alpha s\)) remains integral and orient perpendicular to the air–water interface.

Traces g and h in Figure 4C show the isotherms of (palm2-\(\alpha s-\alpha s\)) and heme2-(palm2-\(\alpha s-\alpha s\)): characterized in our previous paper.12 The low surface pressure region of the isotherms of (palm2-\(\alpha s-\alpha s\)) and heme2-(palm2-\(\alpha s-\alpha s\)) is very similar to that of (\(\alpha s-\alpha s\)) and heme2-(\(\alpha s-\alpha s\)). Figure 4D reveals that the \(n\) value is 3.89 for (palm2-\(\alpha s-\alpha s\)) and 1.83 for heme2-(palm2-\(\alpha s-\alpha s\)), and the \(A_0\) value is 590 \(\AA^2/\alpha l\)-helix for heme2-(palm2-\(\alpha s-\alpha s\)) and 598 \(\AA^2/\alpha l\)-helix for heme2-(palm2-\(\alpha s-\alpha s\)). These indicate that at low surface pressure the number of helices associated as independent particles and helices orientation of (palm2-\(\alpha s-\alpha s\)) and heme2-(palm2-\(\alpha s-\alpha s\)) are similar to those of (\(\alpha s-\alpha s\)) and heme2-(\(\alpha s-\alpha s\)).

Remarkably, upon compression to high surface pressure, the isotherms of (palm2-\(\alpha s-\alpha s\)) and heme2-(palm2-\(\alpha s-\alpha s\)) display another phase transition that differs greatly from the isotherms of (\(\alpha s-\alpha s\)) and heme2-(\(\alpha s-\alpha s\)). This phase transition was suggested12 to correspond to the change of \(\alpha l\) helices orientation from parallel to perpendicular to the air–water interface (note that this high surface pressure region does not lend itself to analysis by eq 1). The surface area per \(\alpha l\)-helix at this second phase transition is similar to the surface area per \(\alpha l\)-helix of heme2-(\(\alpha l-\alpha l-\alpha s-\alpha s\)) at the first phase transition (trace f in Figure 4B). This match adds weight to the conclusion that both maquettes orient with \(\alpha l\)-helix axes perpendicular to the air–water interface. The difference is that high surface pressures are required to lift (palm2-\(\alpha s-\alpha s\)) and heme2-(palm2-\(\alpha s-\alpha s\)) out of the dissociated subunit stage with horizontal \(\alpha l\)-helices into the vertical orientation, while heme2-(\(\alpha l-\alpha l-\alpha s-\alpha s\)) adopts the vertical orientation spontaneously.

3. Characterization of the Heme Protein Maquettes on LB Films. The heme protein maquettes heme2-(\(\alpha s-\alpha s\)), heme2-(\(\alpha l-\alpha s-\alpha s\)), and heme2-(\(\alpha l-\alpha s-\alpha s\)) were transferred to quartz substrates and studied by UV–vis and CD spectra. UV–vis spectra revealed the heme ligation status, and CD spectra revealed the \(\alpha l\)-helical secondary structure composition. CD

**Figure 4.** Orientation of maquettes on the air–water interface characterized by surface pressure–area isotherms. The subphase is 1 mM phosphate buffer (with no salt), \(T = 22\,^\circ\text{C}\). Part A shows the \(\pi–A\) isotherms of the monolayer films derived from three maquettes without heme: (\(\alpha ss-\alpha ss\)), (\(\alpha l-\alpha s-\alpha s\)), and (\(\alpha l-\alpha ss-\alpha s\)). Part B shows the \(\pi–A\) isotherms of the monolayer films derived from three maquettes with heme: heme2-(\(\alpha ss-\alpha ss\)), heme2-(\(\alpha l-\alpha ss-\alpha s\)), and heme2-(\(\alpha l-\alpha l-\alpha ss-\alpha s\)). Part C shows the \(\pi–A\) isotherms of the monolayer films derived from (palm2-\(\alpha s-\alpha s\)), (palm2-\(\alpha s-\alpha s\)), (palm2-\(\alpha l-\alpha ss-\alpha s\)), and (palm2-\(\alpha l-\alpha l-\alpha ss-\alpha s\)). Part D shows plots of \(RT/\gamma\) versus \(A\) for \(\pi\) from 1 to 4 mN/m. The experimental points (open circles) are fitted with linear equations \(RT/\gamma = n(A – A_0)\) (solid lines): (a) (\(\alpha ss-\alpha ss\)), \(n = 2.09, A_0 = 593\,\AA^2/\alpha l\)-helix; (b) heme2-(\(\alpha s-\alpha ss\)), \(n = 2.05, A_0 = 602\,\AA^2/\alpha l\)-helix; (c) (\(\alpha l-\alpha ss-\alpha s\)), \(n = 4.06, A_0 = 573\,\AA^2/\alpha l\)-helix; (d) heme2-(\(\alpha l-\alpha ss-\alpha s\)), \(n = 3.91, A_0 = 580\,\AA^2/\alpha l\)-helix; (e) (\(\alpha l-\alpha l-\alpha ss-\alpha s\)), \(n = 3.89, A_0 = 322\,\AA^2/\alpha l\)-helix; (f) heme2-(\(\alpha l-\alpha l-\alpha ss-\alpha s\)), \(n = 4.19, A_0 = 194\,\AA^2/\alpha l\)-helix; (g) heme2-(\(\alpha s-\alpha ss-\alpha s\)), \(n = 1.81, A_0 = 590\,\AA^2/\alpha l\)-helix; (h) heme2-(palm2-\(\alpha ss-\alpha s\)), \(n = 1.83, A_0 = 598\,\AA^2/\alpha l\)-helix. Orientations of the maquettes on the air–water interface that are consistent with the observed \(n\) and \(A_0\) values are illustrated near the isotherms in Parts A–C.

* The independent particles on the air–water interface. Figure 4D also reveals that the \(A_0\) for (\(\alpha l-\alpha l-\alpha s-\alpha s\)) is 573 \(\AA^2/\alpha l\)-helix, which also approximately coincides with the area of each \(\alpha l\)-helix if they are parallel to the air–water interface. Thus, although the two \(\alpha l\)-\(\alpha ss\) units are still linked together by the disulfide bond, they do not form a fully associated bundle but rather lie flat side-by-side or end-to-end on the air–water interface with polar...
air-facing surface is not necessarily strongly hydrophobic, just hydrophobic alkylated substrate upon immersion. However, the relative to the substrate plane.

vis linear dichroism revealed the orientation of the heme plane
spectra also indicated the α-helical orientations, while UV—
vis linear dichroism revealed the orientation of the heme plane relative to the substrate plane.

Deposition Ratio. In the LB method, the relatively hydrophobic air side of the surface film, should adhere to a hydrophobic alkylated substrate upon immersion. However, the air-facing surface is not necessarily strongly hydrophobic, just less hydrophobic than the water-facing film surface; thus, deposition may not be perfect. The deposition ratio of heme2−(α-ss-α)2 and heme2−(α-l-α-s)2 on transfer from the LB trough to the quartz slides was found to be close to unity, while the deposition ratio of heme2−(α′-l-α′-s)2 was not found to be so high; values in the range 0.6—0.8 were typical.

Ultraviolet—Visible Spectra. The UV—vis spectra of heme2−(α-ss-α)2, heme2−(α-l-α-s)2, and heme2−(α′-l-α′-s)2 in solution all display the same absorption properties characteristic of intact bis-histidine heme ligation in both the ferric and ferrous forms. The spectra of heme2−(α-ss-α)2 and heme2−(α′-l-α′-s)2 as LB films on quartz also display the characteristic UV—vis absorption properties of bis-histidine hemes (Figure 5), and they are identical to the spectra of the maquettes in solution (Figure 3B). Hence, it is evident that the bis-histidine heme ligation is conserved on these LB films. In contrast, the spectrum of LB film of heme2−(α-l-α-s)2 differs from the others and shows absorption properties of unligated hemes (Figure 5), indicating loss of heme ligation on the film. These films were not analyzed further, and possible reasons for loss of heme will be discussed later.

The density of the bis-histidine ligated hemes of the heme2−(α-ss-α)2 and heme2−(α′-l-α′-s)2 in the LB films was estimated from the absorbance of the Soret band maximum between 411 and 412 nm, assuming the solution extinction coefficient (ε = 120 000 M−1 cm−1) applies and taking into account the deposition ratio. The heme density in the heme2−(α-ss-α)2 LB films (deposition ratio unity) was estimated to be 1230 Å2/bis-histidine ligated heme. This is equivalent to 615 Å2/α-helix and directly consistent with the 602 Å2/α-helix value found by the isotherm on the air—water interface. The heme density in the heme2−(α′-l-α′-s)2 LB films was estimated to be 570 Å2/bis-histidine ligated heme, which after correction for the deposition ratio (here measured to be 0.7; see earlier) is equivalent to 200 Å2/α-helix, again consistent with the 194 Å2/α-helix value determined by the isotherm.

CD Spectra. Figure 6 compares the CD spectra of heme2−(α-ss-α)2, (α′-l-α′-s)2, and heme2−(α′-l-α′-s)2 LB films and their corresponding solution spectra. The CD spectra of the LB films conserve the shape that is characteristic of a predominately α-helical structure. We note, however, that the ratio of two minima at 208 and 222 nm of the LB film spectra differs from that of the solution spectra. The CD band at 208 nm in the heme2−(α-ss-α)2 and (α′-l-α′-s)2 LB films is considerably more negative in comparison to their solution spectra, while the 208 nm band in the heme2−(α′-l-α′-s)2 LB film is considerably less negative in comparison to its solution spectrum.

The Moffit exciton theory predicts that the band at 208 nm is polarized parallel to α-helices. This prediction was proved by several experiments, including the work of Olah and Huang et al. and de Jongh et al. Since the incident light is normal to the substrate, a more negative 208 nm band corresponds to a predominantly parallel orientation of the helices with respect to the substrate surface and a less negative 208 nm band corresponds to a predominantly perpendicular orientation of the helices. Thus, the helices in heme2−(α-ss-α)2 and (α′-l-α′-s)2 LB films are predominantly parallel to the solid substrate while the helices in the heme2−(α′-l-α′-s)2 LB film are predominantly perpendicular to the substrate, results entirely consistent with their orientations identified by LB isotherms on the air—water interface.

Ultraviolet—Visible Linear Dichroism. The dichroic ratio of absorption of UV—vis light linearly polarized perpendicular versus parallel to the plane of incidence reveals the average tilt angle of the heme plane with respect to the substrate. It is generally assumed that the electronic transitions responsible for
the Soret and \( \alpha \)-bands in hemes are isotropically polarized in the plane of the heme.\(^{36} \) A relationship has been described\(^{37} \) to estimate the average orientation of the heme group in heme proteins.

\[
D = \cos^2 \theta + \frac{2 \sin^2 \psi}{1 + \cos^2 \psi} \sin^2 \theta
\]  

(2)

where \( \theta \) is the incident angle of the incoming light on the LB film and \( \psi \) is the angle between the normal of the heme plane and the normal of the substrate plane.

Figure 7 shows the UV--vis absorption spectra of a monolayer LB film of heme\(_2\)-(\(\alpha\)-ss-\(\alpha\))\(_2\) and heme\(_2\)-(\(\alpha\)^{-1}\(\alpha\)-ss-\(\alpha\))\(_2\) with parallel and perpendicular polarized incident light and with an incident angle of 60°. The dichroic ratio of the heme\(_2\)-(\(\alpha\)-ss-\(\alpha\))\(_2\) LB film is 0.60, corresponding to an average 38° tilt angle of the heme plane relative to the substrate plane. In contrast, the dichroic ratio of the heme\(_2\)-(\(\alpha\)^{-1}\(\alpha\)-ss-\(\alpha\))\(_2\) LB film is 1.14, corresponding to an average 60° tilt angle of the heme plane relative to the substrate plane.

As shown in our previous paper,\(^{12} \) if we assume that the heme planes are parallel to the \( \alpha \)-helix axis, an average 38° tilt angle of the heme plane implies that the average helix tilt angle is some value between 0° and 38°, and likewise, an average 60° tilt angle of the heme plane implies that the average helix tilt angle is between 0° and 60°. Thus, we cannot infer the helix tilt angle from these helix plane tilt angles. Nevertheless, these results further support the CD results that indicated that the helices in heme\(_2\)-(\(\alpha\)-ss-\(\alpha\))\(_2\) LB films are predominantly parallel to the solid substrate and those in heme\(_2\)-(\(\alpha\)^{-1}\(\alpha\)-ss-\(\alpha\))\(_2\) LB films are predominantly perpendicular to the substrate.

**Discussion**

Molecules at air--water interfaces are exposed to very strong surface tension forces quite different from the forces in bulk aqueous phase. These forces can have a profound effect on the molecular structure of four-\( \alpha \)-helix bundles originally designed for stability in the bulk aqueous phase. While the hydrophobic force will drive bundle assembly in solution by minimizing the hydrophobic surface area exposed to solvent, at the air--water interface this force will be asymmetric and bundle formation is not ensured. Careful redesign can create molecules with both desirable solubility in the bulk aqueous phase and controlled self-assembly at the interface.

In the case of the original (\( \alpha \)-ss-\( \alpha \))\(_2\) maquettes (with or without hemes), the hydrophobic forces and spiral distribution of the charged residues driving bundle association in solution (\( K_d \approx 10^{-12} \text{ M} \)) does not ensure bundle assembly at the air--water interface. The hydrophobic air phase appears to successfully compete with the hydrophobic interactions of the four-\( \alpha \)-helix-bundle core to simply open up the hydrophobic interface between two \( \alpha \)-ss-\( \alpha \) units and expose the hydrophobic face of each of the \( \alpha \)-ss-\( \alpha \) to the air, as shown in Figure 8 (the shaded regions in parts C and D of Figure 8 represent the hydrophobic parts of the helices). However, because added heme binds internally in the \( \alpha \)-ss-\( \alpha \) unit, the bis-histidine ligations of the hemes can be maintained on the air--water interface, as shown in parts B and D of Figure 8.

In the case of (\( \alpha \)-l-\( \alpha \)-ss-\( \alpha \))\(_2\) maquettes (with or without hemes), four-\( \alpha \)-helix-bundle formation is also overwhelmed on the air--water interface. However, the topological constraints of bundle association--dissociation in the covalently linked four-\( \alpha \)-helix bundle are much different than in the noncovalently linked (\( \alpha \)-ss-\( \alpha \))\(_2\). While the (\( \alpha \)-ss-\( \alpha \))\(_2\) four-\( \alpha \)-helix bundle can simply dissociate at the interface into \( \alpha \)-ss-\( \alpha \) units that maintain histidines in proximity for heme binding (Figure 8), the covalently linked bundle can partially dissociate in several ways, only some of which maintain histidines in proximity for heme binding (Figure 9). The configuration shown in parts A and B of Figure 9 in principle can maintain binding of two hemes, and the configuration shown in parts C and D of Figure 9 in principle can maintain binding of one heme between helices 2 and 3. It is clear that no significant binding of heme occurs (Figure 5). We suggest that interference from helices 1 and 4 force the histidine pair of helices 2 and 3 to point away from each other, causing the loss of the heme, as shown in parts E and F of Figure 9. An additional and possibly significant contributor to heme loss in a surface reorganization process is that a transient loss may be irreversible.

The dissociation of water-soluble bundles at the air--water interface can be conquered with several strategies. While the addition of hydrophobic tails to (\( \alpha \)-ss-\( \alpha \))\(_2\) maquettes, such as (palm\(_2\)-\( \alpha \)-ss-\( \alpha \))\(_2\) maquettes (with or without hemes), did not
The attractive forces between adjacent helices. These external forces, which will tend to rotate helices to maximize the attractive interaction between complementary charged surfaces, and hydrophobic forces, which will tend to minimize the exposure of the hydrophobic residues of these helices to the aqueous subphase. Introducing the largely hydrophobic heme cofactor may significantly shift this balance to favor the geometry of Figure 11B, with helices oriented more perpendicular to the film surface. In another view of this balance, protein repacking upon heme binding may shift the distribution of partly exposed hydrophobic residues of the core to favor an orientation with one end of the bundle exposed to the air.

Segregation of like charges onto different helices is in a sense an extreme design because the attractive forces between the helices are so strong that under low ionic strength conditions they cause the bundles to aggregate. However, charge segregation is a common design in nature where relatively large patches of like charges are distributed on one side of a protein, such as cytochrome c and plastocyanin. This kind of charge distribution usually promotes the binding of proteins to an oppositely charged surface. The modification of the charged residues distribution in de novo designed proteins offers an engineering technique to control the assembly of proteins not only at the air–water interface but on surfaces in general.

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References and Notes

Heme Protein Maquette Monolayers
