

signed to the rotational diffusion of the proteins [19,20], gives an excellent measure of dimer or higher aggregate formation. The identical relaxation times before and after heme binding, $\tau_{\text{rHSA}} \sim 52\text{--}58$ ns, provides identical effective molar volume for rHSA and rHSA-heme, $V^{\text{eff}} = 4.7 \times 10^4 \text{ cm}^3 \text{ mol}^{-1}$ ($c \rightarrow 0$), according to the Stokes-Einstein-Debye equation, which is very close to the anticipated value of $4.9 \times 10^4 \text{ cm}^3 \text{ mol}^{-1}$ from the molecular mass and specific volume of HSA. This reveals that the freedom of the rotational diffusive motion of the protein is not significantly affected by the heme incorporation.

We point out that in contrast to HSA, well-investigated aqueous lysozyme solutions [11–14,21,22] are already in aggregation regime; even at very low ionic strength, most of lysozyme molecules stick together due to its highly adhesive nature, which is demonstrated by the appearance of the low- q subpeak in $S^{\text{eff}}(q)$ and d^* coinciding with the diameter of the protein molecule. Although there arose a controversy as to whether the low- q rise for lysozyme solutions is due to a long-range attraction (LRA) [13,21] or large aggregate formation [22], as for rHSA-heme, the simultaneous observations of the low- q rise in $S^{\text{eff}}(q)$ with a host of evidence for monodispersity of the protein, such as d^* far exceeding the contact distance, no additional frictional force on the rotational diffusive motions, identical molecular volume with rHSA, and unbiased COP, provide a plausible argument for the emergence of a LRA. Note that any kind of protein aggregation requires the direct contact between the monomers. Generally, when the repulsion is so strong as to make the particles apart, the low- q rise can be explained only by a LRA [12].

The theoretical $S(q)$ analysis based on a two Yukawa potential model [13] has shown that the relatively longer attraction range than the repulsion one is necessary to produce the so-called zero- q peak in $S(q)$. The more pronounced low- q decrease in $S^{\text{eff}}(q)$ for aqueous rHSA is clearly taken over by the deeper dip in $S^{\text{eff}}(q)$ for aqueous rHSA-heme, which indicates that the electrostatic repulsion is still active in rHSA-heme solutions and the attraction range is greater than the range of the weakly screened electrostatic repulsion.

For further quantitative description, we tested a two Yukawa model [12,13] for $S^{\text{eff}}(q)$ of rHSA-heme. When the attraction range is very long, the model produces a downward convex low- q rise and a huge zero- q intensity reaching more than ~ 1000 even at small c . However, $S(q)$ starts to rise at $q \leq 0.1 \text{ nm}^{-1}$, whereas we observed the onset around $q = 0.2 \text{ nm}^{-1}$. The real expression for a LRA and its potential shape are still not very clear, but the formalism of the potential should significantly affect the low- q shape of $S(q)$. If the system exhibits more slowly decaying attractive potential than the Yukawa decay at small- r , the onset of the low- q rise is expected to shift to higher- q values than that predicted by the Yukawa LRA model.

It is important to recognize that isotropic interaction is not self-evident for any kind of protein system because

proteins have irregular shape and inhomogeneously distributed patches by nature. However, until now, almost all experimental and theoretical works on the interactions proteins have been performed based on mean spherical approximation (MSA) and an isotropic interaction assumption [10–15,23]. The recent theoretical work of Bianchi *et al.* [24] revealed that particles interacting with an anisotropic attractive potential can enhance anomalous density fluctuations or gel-network formation even at very low volume fractions. This also implies that anisotropic potentials caused by a site-specific interaction or inhomogeneous distributions of charge or hydrophobic patches may generate unexpectedly drastic effects on the spatial correlations of proteins, where the inherent limitation of small-angle scattering technique lies in the fact that such anisotropic interactions are reduced into one-dimensional $S(q)$. Nevertheless, carefully confirmed monodispersity of rHSA-heme leads us to conclude that our present interpretation based on a LRA is still broadly correct, even if the actual situation is much more complicated, where anisotropic interactions may affect the spatial correlation of the proteins.

Compared to the well-developed short-range attraction and long-range electrostatic repulsion [11,13,23], the general understanding of a LRA of proteins in solution

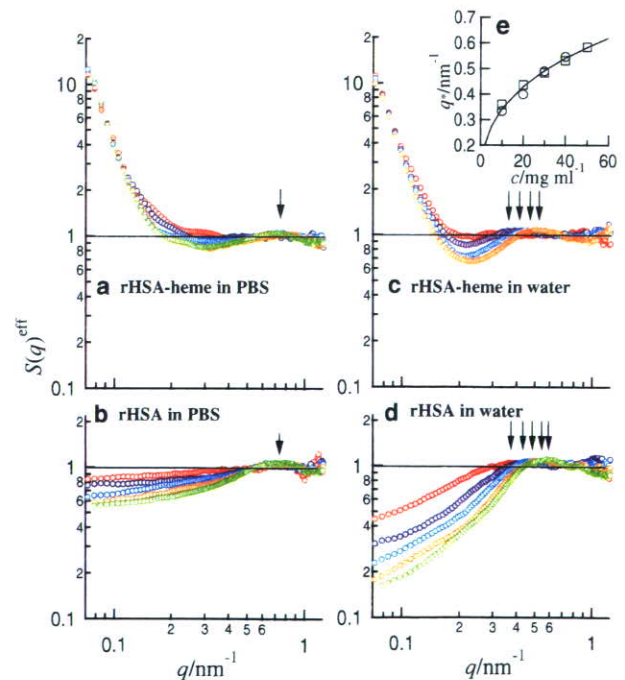


FIG. 3 (color online). Effects of concentration, ionic strength, and heme binding on $S^{\text{eff}}(q)$. (a) rHSA-heme and (b) rHSA in 0.15M PBS solutions and (c) rHSA-heme and (d) rHSA in aqueous solutions in $10 \leq c/\text{mg ml}^{-1} \leq 50$ (an increment of 10 mg ml^{-1}). (e) The protein-protein correlation peak position q^* in $S^{\text{eff}}(q)$ for aqueous rHSA (\square) and rHSA-heme (\circ) as a function of c .

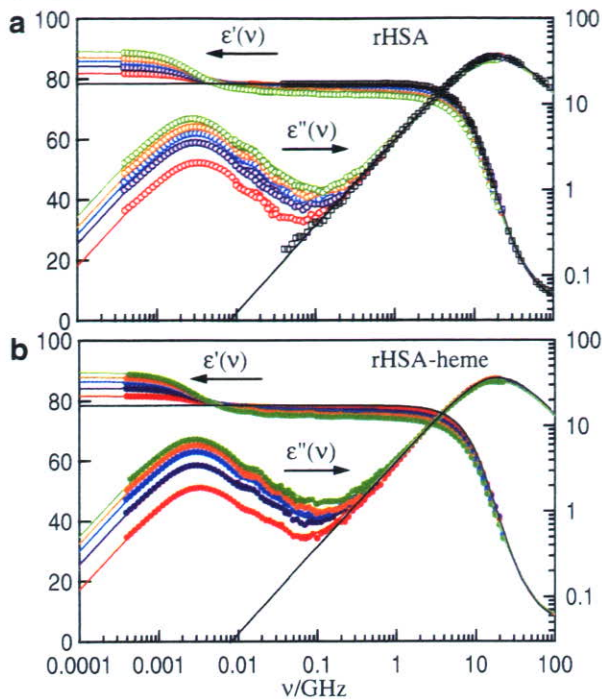


FIG. 4 (color online). Complex dielectric spectra of aqueous solutions of (a) rHSA and (b) rHSA-heme in $10 \leq c/\text{mg ml}^{-1} \leq 50$ (an increment of 10 mg ml^{-1} from the bottom) at 25°C .

[12,13,15] is at an incipient stage. Judging from unbiased $pI(= 4.9)$ and polarization fluctuation amplitudes for rHSA-heme, net charges and their distributions are unlikely to be modified by heme binding, whereas how the occupation of the ligand-binding site allosterically affects the electrostatic interaction of the protein is unclear. The physical origin of a LRA might be entropic driven, possibly due to modulated hydrophobic patches and their inhomogeneous distributions.

HSA binds heme; the rHSA-heme hybrid takes advantage of this naturally occurring process. In the human body, heme released from methemoglobin is immediately captured by hemopexin or HSA acting as scavengers, and it is efficiently transported to the liver for metabolism [25,26]. Since the emergence of the collective nature of HSAs while preserving the monodispersity could be an efficient way to give the ligand-filled HSA molecules a sort of marker, our data suggest that the heme-bound or -unbound HSAs may be recognized in the bloodstream in terms of the presence and absence of the LRA. Therefore, the optimization of the interparticle potential will be a key to

the control of distribution, circulation persistence, and metabolism of functional ligands for medical applications.

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O₂-Binding Albumin Thin Films: Solid Membranes of Poly(ethylene glycol)-Conjugated Human Serum Albumin Incorporating Iron Porphyrin

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Poly(ethylene glycol) (PEG)-conjugated human serum albumin (HSA) incorporating the tetrakis($\alpha,\alpha,\alpha,\alpha$ -amidophenyl)porphyrinatoiron(II) derivative (FeP) [PEG(HSA-FeP)] is a unique plasma protein-based O₂ carrier as a red blood cell substitute. The aqueous solution of PEG(HSA-FeP) [mw of PEG: 2-kDa (PEG₂) or 5-kDa (PEG₅)] was evaporated on a glass surface to produce a red-colored solid membrane. Scanning electron microscopy observations revealed that the PEG₂(HSA-FeP) membrane consisted of two parts: (i) a surface layer made of a fibrous component (10 μ m thickness), and (ii) a bottom layer of an amorphous phase (5 μ m thickness). The condensed solution provided a thick membrane (70 μ m), which also has the amorphous bottom layer. On the other hand, the PEG₅(HSA-FeP) produced homogeneous membrane made of the fibrous component. The FeP active sites in the solid membrane formed very stable O₂-adduct complexes at 37 °C with a half-lifetime of 40 h. The O₂-binding affinity of the PEG₂(HSA-FeP) membrane ($P_{1/2}$ = 40 Torr, 25 °C) was 4-fold lower than that in aqueous solution, which is kinetically due to the low association rate constant. The membrane was soluble again in water and organic solvents (ethanol and chloroform) without deformation of the secondary structure of the protein. The addition of hyaluronic acid gave a free-standing flexible thin film, and it can also bind and release O₂ as well. These O₂-carrying albumin membranes with a micrometer-thickness would be of significant medical importance for a variety of clinical treatments.

INTRODUCTION

Covalently surface-modified proteins with poly(ethylene glycol) (PEG) show a number of unique properties that make them of interest in a range of practical applications (1–6). The most beneficial effect of the PEG conjugation from a biological aspect is to confer a nonimmunogenicity to the proteins, rendering them invisible in the body (7–10). Several classes of protein drugs, such as enzymes and cytokines, have already been approved by the FDA (4, 5), and one of the expected compounds is a PEG-conjugated hemoglobin (PEG-Hb) for use as artificial blood (11, 12). The optimized product has completed a phase I trial and is currently undergoing a phase II safety study (13). Second, the PEG modification allows the proteins to be soluble in nonaqueous solvents (benzene, ethanol, and chloroform, etc.) (14–16). In organic solutions, the proteins exhibited new characteristics, for example, a high thermal stability and a different substrate selectivity. Third, the aqueous PEGylated proteins can be dried on a flat surface to produce solid membranes without any loss of their original activities. However, only limited characterization has been performed on the PEG-conjugated proteins in the solid state.

Recombinant human serum albumin (HSA) incorporating the tetrakis{ $\alpha,\alpha,\alpha,\alpha$ -(1-methylcyclohexanamido)phenyl}porphyrinatoiron(II) derivative (FeP, Figure 1) (HSA-FeP) is a unique plasma protein-based O₂ carrier (17, 18). This entirely synthetic hemoprotein can reversibly bind and release O₂ under physiological conditions (pH 7.4, 37 °C) in a fashion similar to

Hb. It has also been demonstrated that the surface decoration of HSA-FeP by PEG improved its circulation persistence in the bloodstream (19). The PEG(HSA-FeP) is now the most promising material for an entirely synthetic red blood cell substitute.

We have recently found that the aqueous PEG(HSA-FeP) solution cast on the glass surface produces a very smooth solid membrane. The spectroscopic measurements revealed that a reversible O₂ binding to FeP took place in the red-colored membrane. We now report our new findings on the layered film structures of the PEG-conjugated artificial hemoprotein and their O₂-binding behavior.

EXPERIMENTAL PROCEDURES

Materials and Apparatus. All reagents were purchased from commercial sources as special grades and used without further purification. 2-{8-(2-Methylimidazolyl-1-yl)octanoyloxymethyl}-5,10,15,20-tetrakis{ $\alpha,\alpha,\alpha,\alpha$ -o-(1-methylcyclohexanamido)phenyl}porphyrinatoiron(II) (FeP) was synthesized according to our previously reported procedures (17, 20). Recombinant HSA was provided by the NIPRO Corp. (Osaka, Japan). 2-Iminothiolane hydrochloride (IMT) was purchased from Wako Pure Chemical Industries, Ltd. (Osaka, Japan). α -[3-(3-Maleimido-1-oxopropyl)amino]propyl- ω -methoxy PEG [averaged mw: 2333, Sunbright ME-020MA, PEG₂, and averaged mw: 5207, Sunbright MEMAL-50H, PEG₅] was purchased from the NOF Corp. (Tokyo, Japan). Hyaluronic acid sodium salt (HA, mw: 1.9–2.7 \times 10³ kDa) was a gift from Shiseido Co., Ltd. (Tokyo, Japan). The water was deionized using Millipore Elix and Simpli Lab-UV. The UV-vis absorption spectra were recorded using an Agilent 8453 UV-visible spectrophotometer fitted with an Agilent 89090A temperature control unit. The circular dichroism (CD) spectra were obtained using a JASCO J-820 spectropolarimeter over the range of 200–250 nm.

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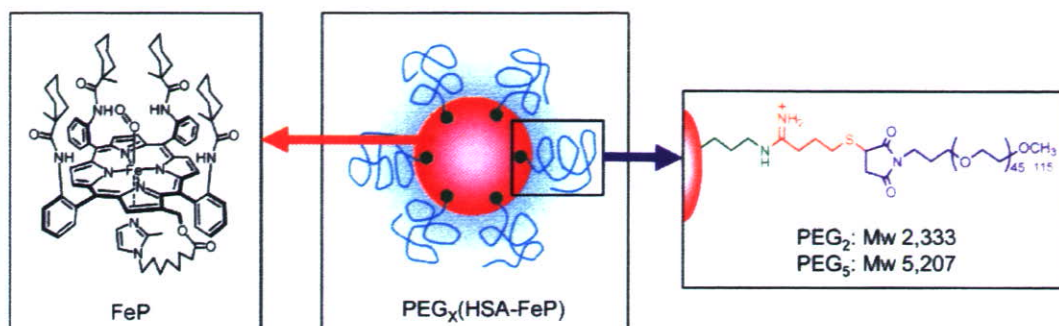


Figure 1. Structure of the $\text{PEG}_x(\text{HSA}-\text{FeP})$ molecule.

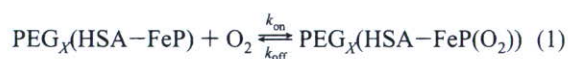
Preparation of $\text{PEG}_x(\text{HSA}-\text{FeP})$ Solid Membranes. The phosphate-buffered saline (PBS) solutions of $\text{PEG}_x(\text{HSA}-\text{FeP})$ ($X = 2$ or 5) were prepared with IMT and PEG_x (pH 7.4, $[\text{HSA}] = 5$ wt %, $\text{FeP}/\text{HSA} = 4/1$ (mol/mol)) according to our previously reported methods (19). The binding number of the PEG_x chain on the $\text{HSA}-\text{FeP}$ surface was determined by the MALDI-TOF MS spectra (19). The PBS solution was dialyzed using a Spectra/Por 1 regenerated cellulose dialysis membrane (MWCO: 6–8 kDa, Spectrum Laboratories, Inc.) against the pure water to remove any included electrolytes. The obtained aqueous solution of $\text{PEG}_x(\text{HSA}-\text{FeP})$ (CO adduct complex, 0.5 mL) was spread on a glass plate [30 × 40 mm, thickness: 0.12–0.17 mm, Matsunami Glass Ind., Ltd. (Osaka, Japan)] and evaporated at an ambient temperature in the dark for 12 h, which produced a red-colored transparent solid membrane.

For the scanning electron microscopy measurements, the samples on the glass plate were sputtered with Pd–Pt using a Hitachi E-1030 ion sputter. The SEM observations were performed using a Hitachi S-4500S field emitted scanning electron microscope.

Furthermore, 0.5 mL of an aqueous solution of hyaluronic acid (HA, 0.2, 0.4, 0.6, 0.8, 1.0 wt %) was added to the $\text{PEG}_2(\text{HSA}-\text{FeP})$ solution (CO adduct complex, 0.5 mL, $[\text{HSA}] = 5$ wt %). The mixture was poured into a poly(styrene) balance dish (44 × 44 × 15 mm) and dried at room temperature in the dark. After 12 h, the formed membrane was gently peeled off and characterized.

The water content of the solid membrane was measured by thermogravimetric-differential thermal analysis using a Rigaku TG8120 instrument at a heating rate of 10 °C/min.

O_2 -Binding Parameters. The O_2 binding to $\text{PEG}_x(\text{HSA}-\text{FeP})$ is expressed by eq 1,



where the O_2 binding equilibrium constant $K = k_{\text{on}}/k_{\text{off}}$.

The $\text{PEG}_x(\text{HSA}-\text{FeP})$ membrane or $\text{PEG}_2(\text{HSA}-\text{FeP})/\text{HA}$ hybrid membrane on the glass plate was placed in a 1-cm quartz cuvette for the spectral measurements, which was sealed tightly with a rubber septa. The O_2 -binding affinity (gaseous partial pressure at which 50% of FeP was dioxygenated, $P_{1/2} = 1/K$) was determined by the spectral changes at various O_2/N_2 pressures (17, 19, 20). The UV–vis absorption spectra were recorded within the range of 350–700 nm. The half-lifetime of the autooxidation of the $\text{FeP}(\text{O}_2)$ complex was determined by the time-course of the absorption change at 426 nm. The association and dissociation rate constants for O_2 (k_{on} , k_{off}) to the $\text{PEG}_x(\text{HSA}-\text{FeP})$ membrane or $\text{PEG}_2(\text{HSA}-\text{FeP})/\text{HA}$ hybrid membrane were measured by a competitive rebinding technique using a Unisoku TSP-1000WK laser flash photolysis instrument (17, 19, 20).



Figure 2. Photograph of the $\text{PEG}_2(\text{HSA}-\text{FeP})$ solid membrane on the glass plate.

RESULTS AND DISCUSSION

Structure of $\text{PEG}_x(\text{HSA}-\text{FeP})$ Membrane. The PEGylation of $\text{HSA}-\text{FeP}$ ($\text{FeP}/\text{HSA} = 4/1$ (mol/mol)) was readily accomplished under mild conditions using the commercially available IMT and maleimide-terminated PEG (19). The average number of the PEG chain on the $\text{HSA}-\text{FeP}$ surface was modulated to 6.0; the viscosity and colloid osmotic pressure of the PBS solution of this molecule ($[\text{HSA}] = 5$ wt %) satisfied the clinical requirements of a red blood cell substitute (19). The solution was subsequently desalted by dialysis against pure water and spread on the glass plate. After drying overnight at room temperature, a red-colored transparent solid membrane was formed (Figure 2). The $\text{HSA}-\text{FeP}$ without the PEG modification did not produce such a homogeneous thin film, only affording a brittle membrane with many cracks.

Scanning electron microscopy observations of the $\text{PEG}_2(\text{HSA}-\text{FeP})$ membrane showed a uniform thickness of 15 μm and a very smooth surface [Figure 3a,b]. From a careful inspection of the side-view, we found that the membrane consists of two parts: (i) the surface layer with a thickness of 10 μm made of a highly oriented fibrous component, and (ii) the bottom layer with a thickness of 5 μm made of an amorphous phase (Figure 3b). Cutting the glass plate with a lateral force, the $\text{PEG}_2(\text{HSA}-\text{FeP})$ membrane was extended and produced long fibers with a width of 350 nm (Figure 3c). The condensed solution ($[\text{HSA}] = 15$ wt %) provided a thick membrane with a thickness of 70 μm , which also has the amorphous bottom layer of 5 μm (Figure 3d). Interestingly, the membrane prepared on the poly(styrene) dish was mainly composed of the amorphous layer; the surface fibrous phase was less than 20% (Figure 3e). On the other hand, water evaporation of the $\text{PEG}_5(\text{HSA}-\text{FeP})$ solution ($[\text{HSA}] = 5$ wt %) on the glass surface provided a homogeneous membrane, which is made of the fibrous component. The precise mechanism of the fiber formation during the water evaporation process is still not clear, but our results show that the structure of the $\text{PEG}_x(\text{HSA}-\text{FeP})$ membrane is very dependent on the surface feature of the substrate and the chain length of the PEG.

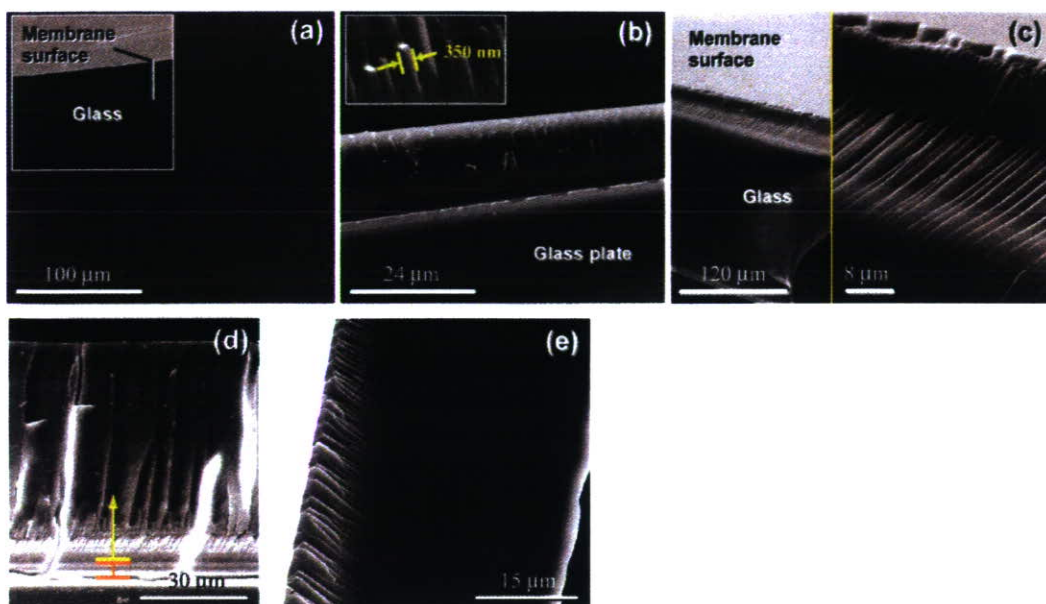


Figure 3. SEM images of the PEG₂(HSA-FeP) membrane on the glass plate. (a) Top-view of the very smooth surface without any crack. (b) Side-view of the membrane section, showing a surface layer made of fibrous component and a bottom layer made of an amorphous phase. (c) Extended nanofiber structure of the surface layer. (d) Thick membrane with a 5 μm amorphous bottom layer (indicated by orange bars) on the glass surface. (e) Membrane prepared on the poly(styrene) surface.

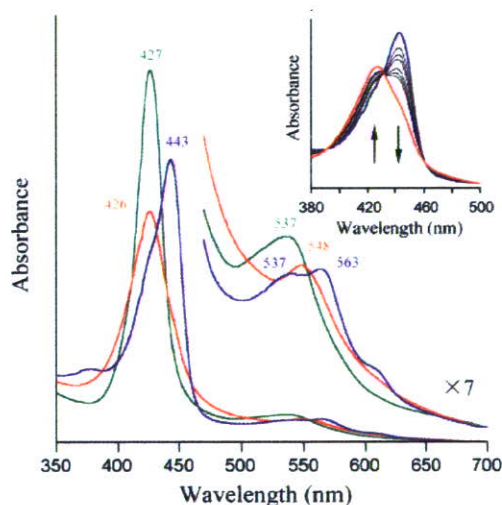


Figure 4. UV-vis absorption spectral changes of the PEG₂(HSA-FeP) membrane on the glass plate at 25 °C (blue line: under N₂, red line: under O₂, green line: under CO). The inset shows the spectral changes at various O₂-partial pressures (P_{O_2} : 0, 10, 19, 30, 76, 114, 152, 760 Torr from blue to red line).

O₂-Binding Properties. The UV-vis absorption spectrum of the PEG_x(HSA-FeP) membrane showed an absorption maxima (λ_{max}) at 427 and 537 nm, indicating the formation of the CO-coordinate low-spin state of FeP (Figure 4) (17, 19–21). This suggested that the ferrous carbonyl complex was retained during the water evaporation for 12 h at an ambient temperature. Light irradiation to the membrane using a 500-W halogen lamp under a 100% O₂ for 20 min led to the CO dissociation and O₂-adduct complex formation of FeP (λ_{max} : 426, 548 nm). By flowing N₂ to the membrane in the quartz cuvette, the UV-vis absorption spectrum shifted to that of a high-spin ferrous complex with an intramolecularly coordinated 2-methylimidazolyl group (λ_{max} : 443, 537, 563 nm). These spectral changes were repeatedly observed and found dependent on the O₂-partial pressure (0 Torr \leftrightarrow 760 Torr), which

demonstrated that the reversible dioxygenation of FeP took place in the membrane. The half-lifetime of the autoxidation ($\tau_{1/2}$) of the FeP(O₂) to Fe³⁺+P was 40 h at 37 °C, which is 3-fold longer than the value in the PBS solution.

The O₂-binding affinity of the PEG₂(HSA-FeP) membrane ($P_{1/2}$ = 40 Torr, 25 °C) determined from the UV-vis absorption spectral changes at various O₂/N₂ pressures (Figure 4 inset) was 4-fold lower (high $P_{1/2}$) than that of the monomeric PEG₂(HSA-FeP) in water (Table 1). Since the O₂ coordination to the Fe porphyrin is an exothermic reaction, the O₂-binding affinity of the membrane decreased at 37 °C ($P_{1/2}$ = 61 Torr). In contrast, PEG₅(HSA-FeP) membrane showed the identical O₂-binding affinity to that in aqueous PBS solution ($P_{1/2}$ = 11 Torr, 25 °C).

In order to elucidate the O₂-binding kinetics of the PEG_x(HSA-FeP) membrane, flash photolysis experiments were carried out. The time course of the absorption change after the laser pulse irradiation to the PEG₂(HSA-FeP) membrane in the quartz cuvette exhibited two first-order kinetics. We previously reported that the binding processes of O₂ to HSA-FeP and PEG_x(HSA-FeP) in aqueous media were fitted to a double-exponential expression, giving two different association rate constants for the fast and slow reactions (k_{on} and k'_{on}) (19, 22). It has been interpreted that the O₂ recombination to FeP in the protein could be affected by the each nanoscopic environment around the accommodation site, for example, a steric hindrance of the amino acid residue and difference in polarity. The k'_{on} was used to be approximately one-third of the k_{on} (17, 19, 22). Nevertheless, the ratio of k'_{on}/k_{on} observed in the PEG₂(HSA-FeP) membrane was less than one-tenth (Table 1). We then theorized that this can be attributed to the O₂ diffusion in the two parts of the membrane. In the major surface layer of the fibrous component, the diffusion of the O₂ could be slower than in water, and it could be much slower in the amorphous bottom phase. The following results support our assumption. (i) The thickness ratio of the two layers (10 μm/5 μm = 2.0) corresponds well to the molar ratio of the k_{on} and k'_{on} (= 2.1). (ii) The absorption decay accompanied with the O₂ rebinding

Table 1. O₂-Binding Parameters of Solid Membranes of PEG_X(HSA-FeP) at 25 °C

system	k_{on} (Torr ⁻¹ s ⁻¹)	k'_{on} (Torr ⁻¹ s ⁻¹)	k_{off} (s ⁻¹)	k'_{off} (s ⁻¹)	$P_{1/2}$ (Torr) ^e
PEG ₂ (HSA-FeP) solution ^b	20	7.5	1.7×10^2	70	11 [32]
PEG ₅ (HSA-FeP) solution ^b	20	10	1.7×10^2	90	11 [31]
PEG ₂ (HSA-FeP) membrane	5.7	0.54	2.3×10^2	22	40 [61]
PEG ₅ (HSA-FeP) membrane	3.1		34		11 [33]
PEG ₂ (HSA-FeP)/HA membrane	4.3	0.22	1.4×10^2	7	32 [60]
Hb solution (T-state) ^c	4.8		1.8×10^2		40
Hb/maltose membrane ^d	1.2×10^{-13}		8.3×10^{-11}		760

^a At 37 °C in brackets []. ^b In PBS solution (pH 7.4), ref 19. ^c In 50 mM potassium phosphate buffer (pH 7.0) at 20 °C, ref 24. ^d Reference 23.

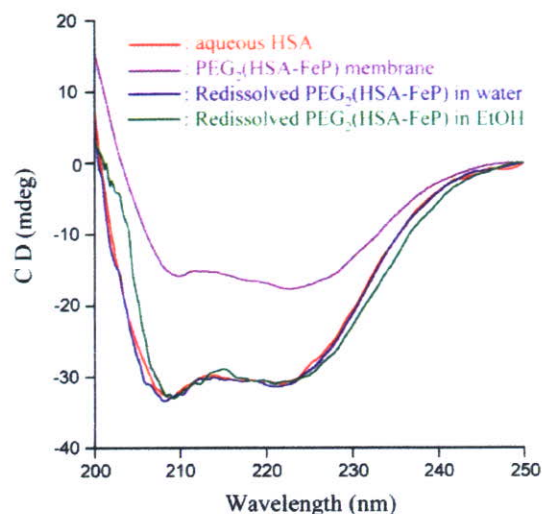


Figure 5. CD spectra of PEG₂(HSA-FeP) at various conditions at 25 °C.

to the PEG₅(HSA-FeP) membrane, which consists of the homogeneous fibrous layer, was fitted by single exponential, giving only one association rate constant k_{on} .

The low O₂-binding affinity of the solid membrane of PEG₂(HSA-FeP) compared to that in the PBS solution is kinetically due to the low association rate constants (Table 1). Even though this result was in significant contrast to the fact that the similar solid membrane of human Hb with maltose (Hb/maltose = 6/4, wt/wt) could not bind O₂ under the same conditions, its O₂-binding affinity was extremely low ($P_{1/2} = 760$ Torr) (23). Since the quaternary structure of Hb was restrained in the solid form, its tense-state conformation cannot change to a relaxed-state with a high O₂-binding affinity (24). In the PEG_X(HSA-FeP) membrane, the protein scaffold was solidified by a bulk water evaporation; however, it would not dramatically influence the dioxygenation of FeP, because the O₂-binding equilibrium is not synchronized with the quaternary structure of the HSA matrix.

PEG_X(HSA-FeP) in Organic Solutions. We have found that the PEG modification enables HSA-FeP to dissolve in

organic solvents. The red-colored PEG_X(HSA-FeP) membrane was homogeneously soluble not only in water but also in ethanol and chloroform. The CD spectrum of the membrane showed a different pattern compared to that in aqueous media (Figure 5). The intensity ratio of the double minimum peaks at 208 and 222 nm (I_{208}/I_{222}) of rHSA or PEG_X(HSA-FeP) is normally 1.1 (25, 26), while the membrane showed 0.9. Interestingly, the redissolved aqueous and ethanol solutions both showed the same spectra as the original aqueous HSA. This result implies that the change of the secondary structure in the protein in the solid and liquid states is reversible. The ethanolic PEG_X(HSA-FeP) also exhibited the same absorption changes upon exposure to O₂ and N₂. These organic solutions can also be cast on a glass surface to form identical PEG_X(HSA-FeP) membranes.

PEG₂(HSA-FeP) Membrane with Hyaluronic Acid. An attempt to isolate the PEG_X(HSA-FeP) membrane from the glass surface unfortunately failed, because it was rather fragile, when peeled off. We then added a supporting polymer to the protein solution and prepared the solid membrane on a glass plate or a poly(styrene) dish. Hyaluronic acid (HA), which is known as a glycosaminoglycan component of connective tissues, hyaline body and extracellular matrix, was selected as the biocompatible polymer support (27). We also expected that the high water retention capability of HA may have a positive effect on retarding the proton-driven oxidation of the FeP(O₂) complex. Water evaporation of the PEG₂(HSA-FeP)/HA mixture [0.5 mL/0.5 mL, total [HSA] = 2.5 wt %, total [HA] = 0.2 wt %] on the glass plate or poly(styrene) dish produced the red-colored uniform solid membranes. It could not be isolated from the glass plate but was easily peeled off from the poly(styrene) surface, providing a free-standing thin film of the PEG₂(HSA-FeP)/HA hybrid [Figure 6a]. The formation of the flexible film was quite dependent on the HA content. When the total concentration of HA is 0.2–0.5 wt %, we could readily obtain the membrane from the plastic surface. The SEM observations of the film showed that it was made of a homogeneous layer with a relatively coarse structure (Figure 6b). In contrast, the film cast on the glass plate consists of the two-layered form: a major coarse phase and bottom dense phase (Figure 6c). This implies

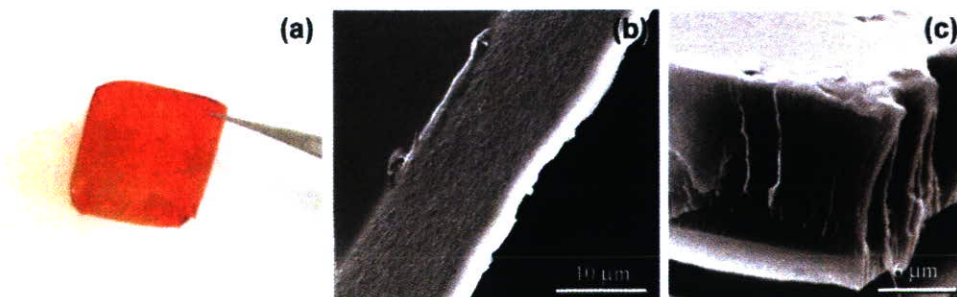


Figure 6. PEG₂(HSA-FeP)/HA membrane. (a) Photograph of the free-standing flexible film, which is prepared on the poly(styrene) surface. (b) SEM of the isolated film. (c) SEM of the membrane prepared on the glass surface.

that the membrane structure of the PEG₂(HSA-FeP)/HA is also influenced by the surface properties of the substrate.

The UV-vis absorption spectral pattern reversibly changed from that of the five-N-coordinate high-spin ferrous complex under an N₂ atmosphere (λ_{max} : 445, 540, 566 nm) to that of the O₂-adduct complex under an O₂ atmosphere (λ_{max} : 426, 545 nm). The O₂-binding parameters of the PEG₂(HSA-FeP)/HA membrane showed a tendency similar to the PEG₂(HSA-FeP) film; the low O₂-binding affinity ($P_{1/2} = 32$ Torr, 25 °C) is mainly due to the slow association rate constant (Table 1). The half-life of the O₂-adduct complex was 40 h at 37 °C; we could not see any water retention effect of HA that could prolong the stability of FeP(O₂). The water content of the PEG₂(HSA-FeP) membrane and PEG₂(HSA-FeP)/HA membrane were determined by the differential thermal analyses to be 6.5 wt % and 6.6 wt %, respectively.

CONCLUSIONS

The stable solid membranes of a PEGylated artificial hemoprotein have been prepared, and their O₂-binding properties were physicochemically characterized in relation to the layer morphology. The red-colored thin film is soluble again in water and organic solvents (ethanol, chloroform) without any deformation of the secondary structure of the protein. The addition of hyaluronic acid as the polymer support gave the free-standing flexible film. These O₂-binding albumin membranes are a red blood cell substitute with a micrometer-thickness that can be preserved anywhere (e.g., on a shelf and in an ambulance) and reproduced as a saline solution at anytime (e.g., at the scene of a disaster). Furthermore, it would be of great medical importance for a variety of clinical treatments, such as O₂-enriched coating agents for medical devices or artificial organs and an O₂-transporting adhesive plaster for wound healing.

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Genetic Engineering of the Heme Pocket in Human Serum Albumin: Modulation of O₂ Binding of Iron Protoporphyrin IX by Variation of Distal Amino Acids

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Abstract: Complexing an iron protoporphyrin IX into a genetically engineered heme pocket of recombinant human serum albumin (rHSA) generates an artificial hemoprotein, which can bind O₂ in much the same way as hemoglobin (Hb). We previously demonstrated a pair of mutations that are required to enable the prosthetic heme group to bind O₂ reversibly: (i) Ile-142 → His, which is axially coordinated to the central Fe²⁺ ion of the heme, and (ii) Tyr-161 → Phe or Leu, which makes the sixth coordinate position available for ligand interactions [I142H/Y161F (HF) or I142H/Y161L (HL)]. Here we describe additional new mutations designed to manipulate the architecture of the heme pocket in rHSA–heme complexes by specifically altering distal amino acids. We show that introduction of a third mutation on the distal side of the heme (at position Leu-185, Leu-182, or Arg-186) can modulate the O₂ binding equilibrium. The coordination structures and ligand (O₂ and CO) binding properties of nine rHSA(triple mutant)–heme complexes have been physicochemically and kinetically characterized. Several substitutions were severely detrimental to O₂ binding: for example, Gln-185, His-185, and His-182 all generated a weak six-coordinate heme, while the rHSA(HF/R186H)–heme complex possessed a typical bis-histidyl hemochrome that was immediately autoxidized by O₂. In marked contrast, HSA(HL/L185N)–heme showed very high O₂ binding affinity ($P_{1/2}^{O_2}$ 1 Torr, 22 °C), which is 18-fold greater than that of the original double mutant rHSA(HL)–heme and very close to the affinities exhibited by myoglobin and the high-affinity form of Hb. Introduction of Asn at position 185 enhances O₂ binding primarily by reducing the O₂ dissociation rate constant. Replacement of polar Arg-186 with Leu or Phe increased the hydrophobicity of the distal environment, yielded a complex with reduced O₂ binding affinity ($P_{1/2}^{O_2}$ 9–10 Torr, 22 °C), which nevertheless is almost the same as that of human red blood cells and therefore better tuned to a role in O₂ transport.

Introduction

In the human circulatory system, iron(III) protoporphyrin IX (hemin) released from methemoglobin (metHb) is captured by a specific glycoprotein, hemopexin (Hpx, 60 kDa), which binds it with very high affinity ($> 10^{13} \text{ M}^{-1}$).^{1,2} Nevertheless, due to the extremely low abundance of Hpx in the blood stream ($\sim 17 \mu\text{M}$), human serum albumin (HSA, 66.5 kDa, 640 μM) acts as a depot of hemin under pathological conditions of trauma and severe hemolysis.³ HSA is the most prominent plasma protein and has a remarkable ability to bind a broad range of insoluble endogenous and exogenous compounds, such as fatty acids,

hemin, bilirubin, bile acids, thyroxine, and a wide variety of drugs.^{4,5} This heart-shaped carrier protein is composed of three structurally similar domains (I–III), each of which contains two subdomains (A and B).^{6,7} Recent crystallographic studies revealed that the hemin is bound within a narrow D-shaped hydrophobic cavity in subdomain IB (Figure 1a).^{8,9} The central iron atom is weakly coordinated by Tyr-161 and the porphyrin propionate side chains interact with a triad of basic amino acid

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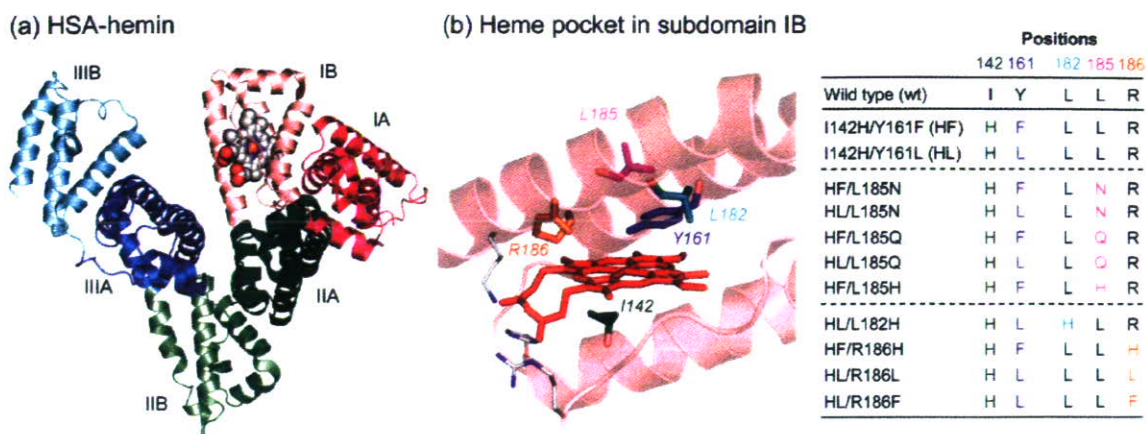


Figure 1. (a) Crystal structure of HSA-hemin complex (1O9X) from ref 9. Hemin is shown in a space-filling representation. (b) Heme pocket structure in subdomain IB and positions of amino acids where site-specific mutations were introduced. The essential double mutations to confer O₂ binding capability to the heme group are I142H and Y161F (or Y161L). Abbreviations of the triple mutants are shown in the table.

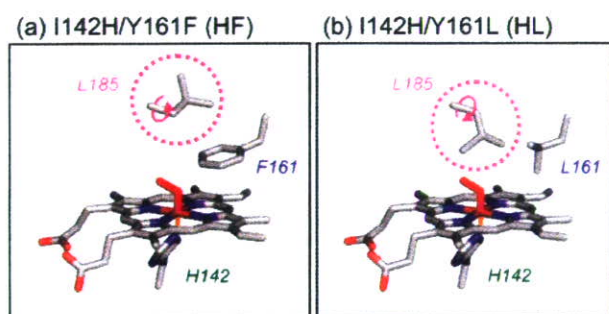


Figure 2. Structural models of the heme pocket in dioxxygenated (a) rHSA(HF)-heme and (b) rHSA(HL)-heme: distal-side steric effect of Leu-185 on O₂ and CO association.²⁰

residues at the entrance (Arg-114, His-146, and Lys-190) (Figure 1b). In terms of the general hydrophobicity of the α -helical heme pocket, subdomain IB of HSA has broadly similar features to the globin-wrapping heme in Hb and myoglobin (Mb). If the HSA-based O₂ carrier is realized, it has the potential of acting not only as a red blood cell (RBC) substitute but also as an O₂-providing therapeutic reagent. However, the reduced ferrous HSA-heme would be immediately autoxidized by O₂, because HSA lacks the proximal histidine that in Hb and Mb allows the prosthetic heme group to bind O₂.¹⁰ On the basis of the detailed structure of the heme binding site of HSA, we introduced a His into the Leu-142 position by site-directed mutagenesis that provides axial coordination to the central Fe²⁺ atom of the heme, and we replaced the coordinated Tyr-161 by Phe or Leu, neither of which can interact with the Fe²⁺ ion (Figure 2).¹¹ This mutagenic approach produced the recombinant HSA(I142H/Y161F)-heme [rHSA(HF)-heme] and HSA(I142H/Y161L)-heme [rHSA(HL)-heme] complexes; these artificial hemoproteins can bind and release O₂ at room temperature, although the O₂ binding affinity of rHSA-heme is at least an order of magnitude lower than that of Hb(α) (R-state).¹¹ To develop this promising O₂-carrying plasma protein as a blood substitute,

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further work is required to regulate the O₂ binding affinity suitable for Hb, Mb, and human RBC.

In Hb and Mb, His-64 on the distal side of the heme has been conserved during evolution and plays an important role for tuning their ligand affinities. A neutron diffraction study of MbO₂ clearly showed that the N–H bond of the distal His-64 is restrained from optimal alignment for strong hydrogen bonding with the coordinated O₂.¹² Olson et al.^{13a} reported that the substitution of Gly for His-64 in Mb and Hb(α) caused a significant decrease in the O₂ binding affinity due to an \sim 100-fold increase in the O₂ dissociation rate constant. A number of systematic investigations of site-directed mutants of Hb and Mb have shown that the overall polarity and packing of the distal residues are key factors in regulating the rate and equilibrium constants for ligand bindings.¹³

In addition to mutagenic analyses of heme binding sites on proteins, the value of using synthetic iron porphyrins as Hb and Mb active-site models has also been amply demonstrated.^{14,15} Tetrakis($\alpha,\alpha,\alpha,\alpha$ -pivalamido)phenylporphyrato-iron(II) “picket-fence porphyrin” of Collman et al.¹⁶ was a pioneering molecule, which forms an O₂ adduct complex at room temperature that is quite stable and shows a high O₂ binding affinity. The polar secondary amide groups in the four fences were believed to contribute to the high O₂ affinity. Moementeau and Lavalette¹⁷ first demonstrated the distal polarity effect on the O₂ binding to the “hanging-base porphyrins”. The presence of the amide groups in the strapped handle over the porphyrin macrocycle yielded a 9-fold higher O₂ binding affinity compared to the ether-bond analogue; it was due to an 8-fold reduction in the dissociation rate constant. This polarity effect of the substituent

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was also well illustrated by our “double-sided porphyrins” having ester fences with a 23-fold lower O₂ binding affinity relative to the picket-fence porphyrin.¹⁸

In view of these investigations, we reasoned that systematic variation of the steric hindrance and local polarity of the heme pocket in subdomain IB of HSA would allow us to modulate the O₂ binding affinity of rHSA–heme. In this study, we designed and generated nine rHSA(triple mutant)–heme complexes, in which the specific third mutation was introduced into three different positions near the O₂ binding site. The effects of the engineered distal amino acids on the O₂ and CO binding properties of the prosthetic heme group have been physico-chemically and kinetically characterized. We now present a new chemistry of albumin-based artificial hemoproteins that would serve as an entirely synthetic O₂ carrier with a controllable ligand binding affinity.

Experimental Section

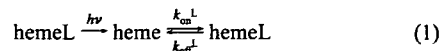
Materials and Apparatus. All materials were reagent-grade and were used as purchased without further purification. Iron(III) protoporphyrin IX (hemin) chloride was purchased from Fluka. UV–vis absorption spectra were obtained on an Agilent 8453 UV–visible spectrophotometer equipped with an Agilent 89090A temperature control unit. Kinetic measurements for the O₂ and CO bindings were carried out on a Unisoku TSP-1000WK time-resolved spectrophotometer with a Spectron Laser Systems SL803G-10 Q-switched Nd:YAG laser, which generated a second-harmonic (532 nm) pulse of 6-ns duration (10 Hz).^{11b} A 150 W xenon arc lamp was used as the probe light source. The gas mixture with the desired partial pressure of O₂/CO/N₂ was prepared by a Kofloc Gasblender GB-3C. MCD spectra were measured by a Jasco J-820 circular dichrometer.

Preparations of rHSA Triple Mutants and Their Heme Complexes. The designed rHSA triple mutants were prepared according to our previously reported techniques.¹¹ The third mutation (L185N, L185Q, L185H, L182H, R186H, R186L, or R186F) was introduced into the HSA coding region in a plasmid vector encoding the double mutants [rHSA(I142H/Y161F) [rHSA(HF)] or rHSA(I142H/Y161L) [rHSA(HL)]]¹¹ by use of the Stratagene QuikChange mutagenesis kit. All mutations were confirmed by DNA sequencing. The plasmid was then digested by *NorI* and introduced into yeast (*Pichia pastoris* GS115) by electroporation. The expression protocols and media formulations were as previously described.^{11b} Briefly, the clones were grown in BMGY medium and transferred to BMMY medium for induction with methanol in baffled shaking flasks at 30 °C, 200 rpm. The obtained proteins were harvested from the growth medium by precipitation with ammonium sulfate and purified by a Cibacron Blue column of Blue Sepharose 6 Fast Flow (Amersham Pharmacia Biotech). The rHSA triple mutants were finally subjected to gel filtration on an ÄKTA Prime Plus FPLC system with a Superdex 75 preparative-grade column (Amersham Pharmacia Biotech). The protein concentration was assayed by measuring the absorbance at 280 nm ($\epsilon_{280} = 3.4 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$) and by SDS–PAGE.

The ferric rHSA(mutant)–hemin complexes [hemin:rHSA(mutant) molar ratio of 1:1] were prepared by established procedures.^{9,11} The resulting samples were analyzed by SDS–PAGE to confirm a pure preparation. The 50 mM phosphate buffered solution (pH 7.0, 3 mL) of rHSA(mutant)–hemin ([hemin] = 10 μM) in a 10-mm path length optical quartz cuvette sealed with a rubber septum was purged with Ar for 40 min. A small excess amount of degassed aqueous sodium dithionate was added by a microsyringe to the sample under an Ar

atmosphere to reduce the central ferric ion of the hemin, to give the ferrous rHSA(mutant)–heme complexes.

O₂ and CO Binding Parameters. The O₂ or CO recombination with the heme after nanosecond laser flash photolysis of hemeO₂ or hemeCO occurs according to eq 1 with the association rate constant (k_{on}^{L}) and dissociation rate constant ($k_{\text{off}}^{\text{L}}$):^{11,16c,19}



where L = O₂ or CO. The CO association rate ($k_{\text{on}}^{\text{CO}}$) was simply measured by following the absorption at 425 nm after laser pulse irradiation to rHSA(mutant)–hemeCO at 22 °C.¹¹ The O₂ association rate constant ($k_{\text{on}}^{\text{O}_2}$) and O₂ binding equilibrium constant K^{O_2} [$= (P_{1/2}^{\text{O}_2})^{-1}$] can be determined by a competitive rebinding technique by use of gas mixtures with different partial pressures of O₂/CO/N₂ at 22 °C.^{11,16c,19} The relaxation curves that accompanied the O₂ or CO recombination were analyzed by single- or double-exponential profiles with Unisoku Spectroscopy & Kinetics software. The O₂ dissociation rate ($k_{\text{off}}^{\text{O}_2}$) was calculated from $k_{\text{on}}^{\text{O}_2}/K^{\text{O}_2}$.

The CO dissociation rate constant ($k_{\text{off}}^{\text{CO}}$) was measured by displacement with NO for rHSA(mutant)–hemeCO at 22 °C.^{11b} The time course of the UV–vis absorption change that accompanied the CO dissociation was fitted to two single exponentials. The CO binding constants [$K^{\text{CO}} = (P_{1/2}^{\text{CO}})^{-1}$] were calculated from $k_{\text{on}}^{\text{CO}}/k_{\text{off}}^{\text{CO}}$. Fresh solutions of rHSA–(mutant)–heme were normally made up for each set of experiments.

Magnetic Circular Dichroism Spectroscopy. MCD for the 50 mM potassium phosphate buffered solutions (pH 7.0) of rHSA(mutant)–hemin or –heme complex (10 μM) were measured under Ar and CO atmospheres with a 1.5 or 1.65 T electromagnet at 22 °C.

Results and Discussion

Design of Distal Pocket with Asn, Gln, and His. We recently compared the O₂ and CO binding properties of the rHSA(double mutant)–heme complexes [rHSA(HF)–heme and rHSA(HL)–heme] and found evidence for a noteworthy distal-side steric effect on ligand binding.^{11b} The rHSA(HF)–heme complex binds O₂ and CO about 4–6 times more tightly than rHSA(HL)–heme, primarily because of enhanced association rate constants. Structurally, this affect appears to be due to the concerted effects of the residues at positions 161 and 185 on ligand binding. In the rHSA(HF)–heme complex, the bulky aromatic side chain of Phe-161 is presumed to prevent rotation of the neighboring Leu-185, thereby providing easy access to the O₂ binding site in the distal pocket (Figure 2a). In contrast, the substitution of Phe-161 by the smaller Leu-161 may allow rotation of the isopropyl group of Leu-185, which reduces the volume of the distal side (Figure 2b) and hinders association of O₂ and CO ligands with the heme iron atom. On the basis of these findings, we reasoned that other modifications of the heme pocket architecture would allow us to further modulate its O₂ binding properties.

One approach to enhancing the O₂ binding affinities of rHSA–(HF)–heme and rHSA–(HL)–heme would be to introduce a histidine into an appropriate position on the distal side of the heme. The N_ε atom of His may act as a proton donor to form an H-bond with the coordinated O₂. However, another important requirement in this molecular design is to prevent the formation

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(20) The pictures were produced on the basis of crystal structure coordinates of rHSA(wt)–hemin (1O9X, ref 9) by use of PyMOL: DeLano, W. L. The PyMOL Molecular Graphics System; DeLano Scientific: San Carlos, CA, 2002.

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of a six-coordinate low-spin ferrous complex. The bis-histidyl hemochromes are normally autoxidized by O_2 via an outer-sphere mechanism as well as by inner-sphere pathways involving the metal-coordinated O_2 .^{21–23} The distal amino acid must therefore be located relatively far ($>4 \text{ \AA}$) from the central iron.

Our modeling experiments suggested that the favorable position for the distal His insertion was Leu-185, which is in the final helix in subdomain IB and forms part of the top of the distal pocket (Figure 1). Leu-182 and Arg-186 were also considered likely to be good candidate positions for the introduction of an amide-containing side chain designed to stabilize O_2 binding (see below). In elegant studies on Mb, Rohlfs and co-workers showed that Gln, which has a primary amide group potential to form an H-bond, was able to substitute effectively for the stabilizing role of the distal histidine (His-64).^{13b,c} Thus, we decided to vary the polarity of the distal side of the heme in rHSA(HF) and rHSA(HL) by replacing Leu-185 with Asn, Gln, and His by site-directed mutagenesis. The Asn residue should behave similarly to Gln, although a rMb-(H64N) mutant has never been reported. The His-185 mutation was only done for rHSA(HF), because His-185 could be long enough to bind to the sixth coordinate position of the heme if allowed the greater freedom of movement that would occur in the rHSA(HL) background. As a result, five triple mutants [rHSA(HF/L185N), rHSA(HL/L185N), rHSA(HF/L185Q), rHSA(HL/L185Q), and rHSA(HF/L185H)] were cloned and their hemin complexes were prepared.

Ferric States of L185N, L185Q, and L185H Mutants. The site-specific mutations with Asn, Gln, and His were successfully introduced into the Leu-185 position of rHSA(HF) or rHSA(HL), and the proteins were purified to homogeneity as determined by SDS-PAGE. The rHSA(mutant)-hemin complexes produced from these proteins were stable for several months at $4 \text{ }^\circ\text{C}$ without precipitation.

The UV-vis absorption spectra of the five rHSA(triple mutant)-hemin complexes are essentially the same regarding their general features (Figure 3, Table 1). When analyzed by MCD spectroscopy to evaluate the redox state, spin state, and axial ligand environment, all the ferric rHSA(triple mutant)-hemins showed a characteristic MCD with similar S-shaped patterns in the Soret band region, though their intensities were dependent on the nature of the distal amino acid (Figure 4). Vickery et al.²⁴ previously reported that the Soret MCD intensity of the ferric Mb with different anions at the sixth coordinate position was correlated with the amount of low-spin component. rHSA(HL)-hemin showed almost the same band as ferric Mb, in which one water axially coordinates to the sixth position of the heme to produce the aquo complex.^{11,24,25} In contrast, rHSA(HF/L185H)-hemin showed 3-fold greater intensity at 405 nm. This is probably caused by the coexistence of a low-spin six-coordinate heme. Introduction of Asn or Gln at position 185

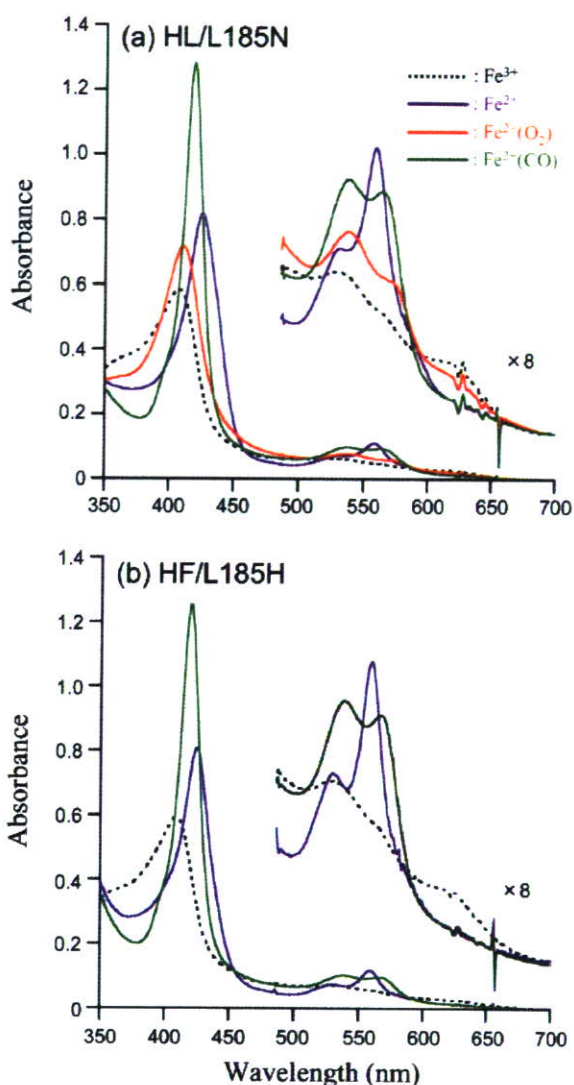


Figure 3. UV-vis absorption spectral changes of (a) rHSA(HL/L185N)-heme and (b) rHSA(HF/L185H)-heme complexes in 50 mM potassium phosphate buffered solution (pH 7.0, $22 \text{ }^\circ\text{C}$).

gave intermediate effects, though mutants with Q185 yielding a slightly more intense peak at 405 nm. Overall, our MCD results for these five rHSA(triple mutant)-hemin complexes imply that the introduction of the distal nitrogenous residue at the 185 position tends to increase the ferric low-spin nature.

Ferrous States of L185N, L185Q, and L185H Mutants and O_2 Binding. rHSA(triple mutant)-hemins were easily reduced to form the ferrous complexes by adding a small excess of aqueous sodium dithionite under an Ar atmosphere. rHSA(HF/L185N)-heme, rHSA(HL/L185N)-heme, and rHSA(HL/L185Q)-heme each showed a visible absorption band at 558–559 nm with a small shoulder at 530 nm (Figure 3a; Figure S1, Supporting Information), that was similar to the spectra observed for rHSA(HF)-heme, rHSA(HL)-heme,^{11b} deoxyMb,²⁷ and synthetic chelated protoheme.²⁶ The spectral patterns clearly indicated the formation of a five-N-coordinate high-spin complex. In contrast, in the spectra of rHSA(HF/L185Q)-heme and

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Table 1. UV-vis Absorption Spectral Data of rHSA(mutant)-Heme Complexes^a

hemoproteins	λ_{max} (nm)			
	Fe ³⁺	Fe ²⁺	Fe ²⁺ O ₂	Fe ²⁺ CO
rHSA(HF)-heme ^b	402, 533, 620	425, 532(sh), 559	411, 538, 576	419, 538, 565
rHSA(HL)-heme ^b	402, 533, 620	426, 531(sh), 559	412, 537, 573	419, 538, 565
rHSA(HF/L185N)-heme	406, 528, 618	425, 530(sh), 559	411, 540, 575	419, 539, 566
rHSA(HL/L185N)-heme	407, 530, 620	425, 530(sh), 559	411, 537, 575	419, 537, 564
rHSA(HF/L185Q)-heme	406, 530, 620	424, 528, 558		419, 538, 566
rHSA(HL/L185Q)-heme	406, 530, 620	425, 530(sh), 558	411, 537, 574 ^c	419, 537, 566
rHSA(HF/L185H)-heme	407, 528, 620	424, 528, 558		419, 538, 566
rHSA(HL/L182H)-heme	410, 532, 624	425, 530, 559		419, 539, 567
rHSA(HF/R186H)-heme	411, 533, 565	424, 529, 560		420, 539, 568
rHSA(HL/R186L)-heme	406, 530, 620	426, 531(sh), 559	411, 539, 576	419, 539, 567
rHSA(HL/R186F)-heme	405, 532, 621	426, 531(sh), 559	410, 535, 571	419, 538, 568
Mb ^d	409, 503, 548(sh), 632	434, 557	418, 544, 581	423, 541, 579
chelated heme ^e	408, 540, 565	427, 530, 558	414, 543, 575	420, 540, 569

^a In 50 mM potassium phosphate buffered solution (pH 7.0) at 22 °C. ^b Reference 11b. ^c At 5 °C. ^d Horse muscle myoglobin (Sigma); ref 11b. ^e In DMF/H₂O (7/3) at 15 °C; ref 26.

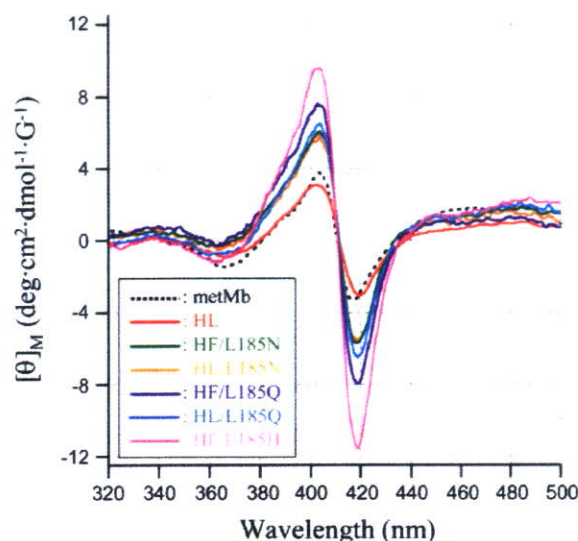


Figure 4. MCD spectra of rHSA(185-mutant)-hemin complexes in 50 mM potassium phosphate buffered solution (pH 7.0, 22 °C).

rHSA(HF/L185H)-heme, the β band at 528 nm appeared relatively sharp (Figure 3b), which suggests partial formation of a six-coordinate heme complex. This is consistent with the finding that the ferric state of these two mutant complexes had the highest peaks in the MCD.

The Soret MCD spectra of ferrous rHSA(HF/L185N)-heme, rHSA(HL/L185N)-heme, and rHSA(HL/L185Q)-heme under Ar atmosphere are dominated by an intense positive peak at 433 nm and a small trough at 402 nm as expected for the Faraday *C*-terms for high-spin ferrous porphyrins like deoxyMb (Figure 5a).^{24,25} In contrast to these three mutant complexes, rHSA(HF/L185Q)-heme and rHSA(HF/L185H)-heme show weaker intensity in the Soret band region and greater intensity in the visible region (Figure 5b).

On the basis of all the UV-vis absorption and MCD spectral results, we concluded that the reduced ferrous heme is axially coordinated by His-142 at the core of the heme pocket in rHSA-(mutant) and forms a five-N-coordinate high-spin ferrous complex under an Ar atmosphere in the case of HF/L185N, HL/L185N, and HL/L185Q mutants (Figure 6a,b,d). In addition to the His-142 ligation, Gln-185 and His-185 partially interact with the sixth coordinate position of the central Fe²⁺ ion of the

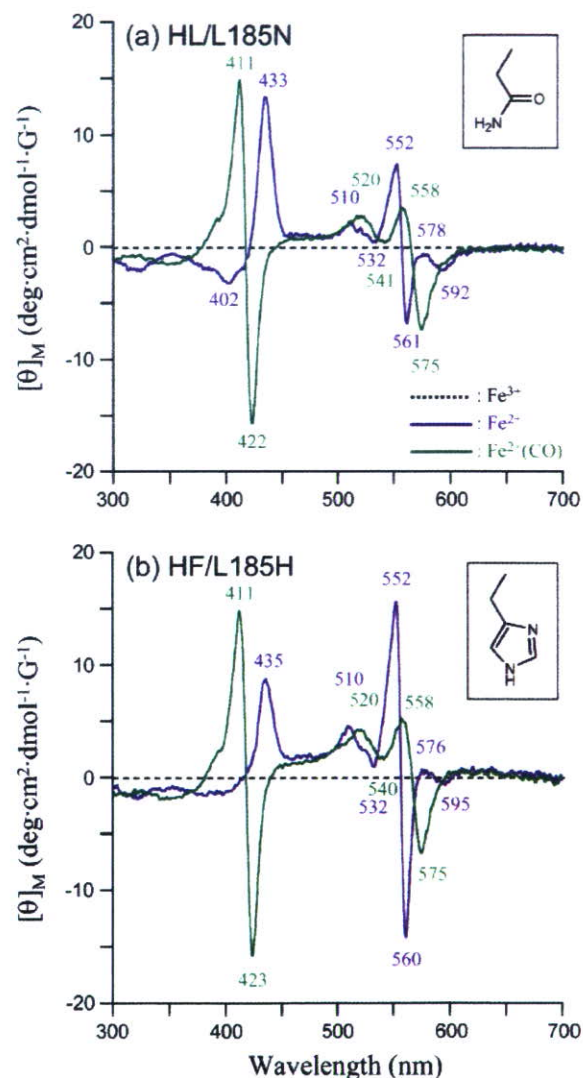


Figure 5. MCD spectral changes of (a) rHSA(HL/L185N)-heme and (b) rHSA(HF/L185H)-heme complexes in 50 mM potassium phosphate buffered solution (pH 7.0, 22 °C).

heme in the HF/L185Q and HF/L185H mutants in spite of the bulky aromatic ring of Phe-161 (Figure 6c,e). We postulated that rHSA(HL/L185Q)-heme would also form a six-coordinate

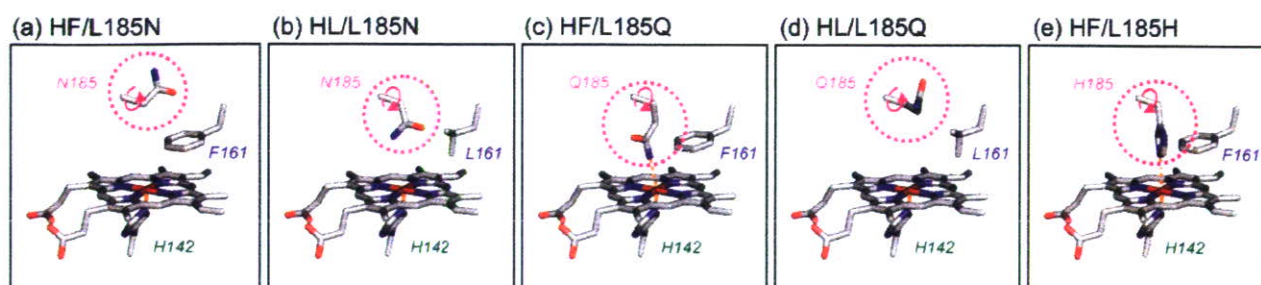


Figure 6. Structural models of the heme pocket in rHSA(triple mutant)-heme complexes: distal-side effects of engineered amino acids at position 185.

low-spin complex, because the small Leu-161 should allow additional room for rotation of Gln-185. However, it yielded a five-coordinate high-spin ferrous complex (Figure S1). This suggests that the flexible Gln-185 may interact with neighboring amino acids (Figure 6d) and underscores the difficulty in accurately predicting the impact of amino acid substitutions.

Upon exposure of the rHSA(HF/L185N)-heme and rHSA(HL/L185N)-heme solutions to O_2 , the UV-vis absorptions immediately changed to that of the O_2 adduct complex at 22 °C (Figure 3a, Table 1).^{11,26,27} However, the rHSA(HL/L185Q)-heme complex bound O_2 only at 5 °C (Figure S1a) and was observed to autoxidize rapidly at 22 °C. This rapid oxidation may suggest that the distal side of the heme has an open structure, which allows easy access of water to the heme, thereby facilitating autoxidation.²⁸ The rHSA(HF/L185Q)-heme and rHSA(HF/L185H)-heme complexes, both of which exhibit side-chain interactions with the sixth coordinate position of the heme, were immediately oxidized by O_2 even at low temperature (5 °C).

After introduction of CO gas, all the hemoproteins produced stable carbonyl complexes with identical absorption spectral patterns (Figures 3 and S1a, Table 1).^{11,26,27} In every case the carbonyl rHSA(triple mutant)-heme complexes exhibited the same S-shaped MCDs, which correspond to the *A*-term bands for the diamagnetic low-spin protoheme with CO and axial His coordinations (Figures 5 and S1b).^{24,25} This result implies that in the carbonyl complexes the Asn-185 and Gln-185 residues do not act as a proximal base instead of His-142.

O_2 and CO Binding Parameters of L185N Mutants. By use of laser flash photolysis, analysis of the kinetics of ligand binding to the double mutants rHSA(HF)-heme and rHSA(HL)-heme revealed that the asymmetric iron protoporphyrin IX molecule is accommodated in subdomain IB in two different orientations (180° rotational isomers).¹¹ As a result, there exist two geometries of the axial His-142 coordination to the central Fe^{2+} ion of the heme (species I and II). In species I, the proximal His coordinates to the heme without strain, while in species II, the ligation involves some distortion, resulting in weaker O_2 binding. The bending strain in the proximal His- Fe^{2+} bond in species II increases the dissociation rate constant and decreases the association rate for CO, whereas it increases the O_2 dissociation rate without changing the kinetics of the O_2 association.^{16c,19} Consequently, the entire absorption decay accompanying the CO recombination with rHSA(HF)-heme or rHSA(HL)-heme was composed of two single exponentials, but the rebinding process of O_2 followed a simple monophasic

decay. In rHSA(triple mutant)-hemes, this alternative geometry of the heme plane would also arise in the same manner.

We again used laser flash photolysis to characterize the O_2 and CO binding properties of the rHSA(triple mutant)-heme complexes. As expected, the binding behavior of O_2 for rHSA(HF/L185N)-heme and rHSA(HL/L185N)-heme was broadly similar to that of the double mutants. However, detailed analysis reveals that the absorption decay accompanied by O_2 rebinding to the heme was composed of two very similar phases (Figure S2, Supporting Information). Numerous investigations of the synthetic iron porphyrins have demonstrated that the “distal-side steric effect” is the only factor that influences the association rate constant for O_2 .^{16c,19} The double-exponential profiles for O_2 association are therefore likely to indicate that there are two distinct conformations of the distal Asn-185 above the heme. The amplitude ratio of the two phases was approximately 1:1 for rHSA(HL/L185N)-heme, suggesting that half of the Asn residue may turn toward the inside of the heme pocket and the other turns to the outside (Figure 6b). These two conformers of the distal Asn-185 residue also influence the association rate for CO. If we were to take this minimal effect into account, the CO rebinding process would have to be analyzed as four phases. However, (i) it would be too complicated to comprehend the fundamental aspects of the ligand binding properties of rHSA(triple mutant)-heme, and (ii) the observed distal-side effect is less significant compared to the major proximal-side steric effect in this system. Hence, the absorbance decays after laser pulse irradiation to rHSA(HF/L185N)-hemeCO and rHSA(HL/L185N)-hemeCO were fitted by biphasic kinetics. The ratio of the amplitude of the dominant fast phase (species I) and minor slow phase (species II) was approximately 7:3 for rHSA(HF/L185N)-heme and 3:2 for rHSA(HL/L185N)-heme. These values were within the same range observed in the rHSA(double mutant)-heme complexes.¹¹ Concomitantly, the O_2 association rate of rHSA(HF/L185N)-heme or rHSA(HL/L185N)-heme was determined as one value by weighted averaging of the $k_{on}^{O_2}$ values for the two phases (Table 2).

In general, k_{off}^{CO} is a simple indicator of the bending strain in the proximal His coordination to the central Fe^{2+} ion.^{16c,19} rHSA(HF)-heme, rHSA(HL)-heme, rHSA(HF/L185N)-heme, and rHSA(HL/L185N)-heme exhibited similar k_{off}^{CO} values in species I (0.008–0.013 s^{-1}) and they are identical to that of Hb(α) (R-state) (0.009 s^{-1}) (Table 3).²⁹ This result indicated that the axial His-142 ligation to the heme in these artificial hemoproteins has the same features as that of Hb.

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Table 2. O₂ Binding Parameters of rHSA(mutant)-Heme Complexes^a

hemoproteins	$k_{on}^{O_2}$ ($\mu\text{M}^{-1}\text{s}^{-1}$)	$k_{off}^{O_2}$ (ms^{-1})		$P_{1/2}^{O_2}$ (Torr)	
		I	II	I	II
rHSA(HF)-heme ^b	20	0.10	0.99	3	31
rHSA(HL)-heme ^b	7.5	0.22	1.70	18	134
rHSA(HF/L185N)-heme	26	0.10	1.03	2	24
rHSA(HL/L185N)-heme	14	0.02	0.29	1	14
rHSA(HL/R186L)-heme	25	0.41	8.59	10	209
rHSA(HL/R186F)-heme	21	0.29	7.01	9	203
Hb(α) (R-state) ^c	33 ^d	0.013 ^e		0.24	
Mb ^f	14	0.012		0.51	
RBC ^g				8	

^a In 50 mM potassium phosphate buffered solution (pH 7.0) at 22 °C. I or II indicates species I or II. ^b Reference 11. ^c Human hemoglobin α -subunit. ^d In 0.1 M phosphate buffer (pH 7.0, 21.5 °C); ref 30. ^e In 10 mM phosphate buffer (pH 7.0, 20 °C); ref 31. ^f Sperm whale myoglobin, in 0.1 M potassium phosphate buffer (pH 7.0, 20 °C); ref 13b. ^g Human red cell suspension, in isotonic buffer (pH 7.4, 20 °C); ref 32.

Table 3. CO Binding Parameters of rHSA(mutant)-Heme Complexes^a

hemoproteins	k_{on}^{CO} ($\mu\text{M}^{-1}\text{s}^{-1}$)		k_{off}^{CO} (s^{-1})		$P_{1/2}^{CO}$ (Torr)	
	I	II	I	II	I	II
rHSA(HF)-heme ^b	6.8	0.72	0.009	0.061	0.0011	0.068
rHSA(HL)-heme ^b	2.0	0.27	0.013	0.079	0.0053	0.240
rHSA(HF/L185N)-heme	7.7	1.09	0.008	0.043	0.0008	0.032
rHSA(HL/L185N)-heme	6.8	1.60	0.008	0.039	0.0010	0.020
rHSA(HL/R186L)-heme	5.0	0.57	0.011	0.165	0.0018	0.234
rHSA(HL/R186F)-heme	7.9	1.12	0.010	0.148	0.0010	0.107
Hb(α) (R-state) ^c	4.6 ^d		0.009 ^e		0.0016 ^f	
Mb ^g	0.51		0.019		0.030	

^a In 50 mM potassium phosphate buffered solution (pH 7.0) at 22 °C. I or II indicates species I or II. ^b Reference 11. ^c Human hemoglobin α -subunit. ^d In 50 mM potassium phosphate buffer (pH 7.0, 20 °C); ref 33. ^e In 0.1 M phosphate buffer (pH 7.0, 20 °C); ref 29. ^f Calculated from (k_{on}^{CO}/k_{off}^{CO})⁻¹. ^g Sperm whale myoglobin, in 0.1 M potassium phosphate buffer (pH 7.0, 20 °C); ref 13b.

Effect of Asn-185 Residue on O₂ Binding Affinity. The O₂ and CO binding parameters for the rHSA(HF)-heme and rHSA(HF/L185N)-heme complexes did not show any significant differences. The bulky benzyl side chain of Phe-161 may retard rotation of the polar amide group of Asn-185 and thereby maintain the polarity and size of the distal pocket (Figure 6a). In contrast, there are marked differences in the comparison of the O₂ and CO binding parameters for rHSA(HL)-heme and rHSA(HL/L185N)-heme. First, the presence of Asn rather than Leu at position 185 resulted in 2- and 3–6-fold increases in the association rate constants for O₂ and CO, respectively. As described above, the kinetics of O₂ binding to rHSA(HL/L185N)-heme actually consist of two phases. The Asn may partly rotate upward, which provides a somewhat greater space for the distal pocket. This presumably increases the association rate constants. Second, Asn-185 induced 18- and 10-fold increases in the O₂ binding affinity for species I and II, respectively (Table 2); these increases were predominantly due to the 6–11-fold diminution of the $k_{off}^{O_2}$ values. This corresponds to a free energy difference of $-1.8 \text{ kcal mol}^{-1}$ at 22 °C that may be attributable to a H-bond interaction with the bound O₂. This is consistent with the observation that, in HbO₂ and MbO₂, the distal His-64 stabilizes the coordinated O₂ by $-0.6\text{--}1.4 \text{ kcal mol}^{-1}$ due to the H-bonding.¹³ Unfortunately, attempts to measure the stretching frequency of the bound O₂ molecule in rHSA(HL/L185N)-heme by infrared spectroscopy failed because the O₂ adduct complex was not sufficiently stable.

Nevertheless, it is noteworthy that the high O₂ binding affinity ($P_{1/2}^{O_2}$ 1 Torr) for rHSA(HL/L185N)-heme is now close to that of natural Hb(α) (0.24 Torr)^{30,31} and Mb (0.5 Torr)¹³ (Table 2).

Replacement of L182 or R186 by His. Leu-182 and Arg-186 were also considered to be good candidates for introduction of the distal His, so we prepared the rHSA(HL/L182H) and rHSA(HF/R186H) triple mutants (Figure 1). Modeling trials demonstrated that neither of these introduced histidines is coplanar with the Fe–O–O moiety. Rather, they are positioned off to the side, so that there may be an oblique interaction with the coordinated O₂ and the heme center.

The rHSA(HL/L182H)-hemin complex and its reduced form showed spectra similar to those of rHSA(HF/L185H)-heme. In contrast, the color of the ferric rHSA(HF/R186H)-hemin solution was bright red, and the UV–vis absorption spectrum clearly showed the formation of a bis-histidine-coordinated low-spin ferric complex (Figure 7a).^{22,26} The MCD intensity of the S-shaped curve in the Soret band region (Figure 8) was higher than that observed with rHSA(HF/L185H)-hemin (Figure 4). The chemical reduction of the Fe³⁺ complex results in very sharp β , α bands in the visible absorption spectrum (529, 560 nm) (Figure 7a). In MCD, we observed the loss of the strong C-terms in the Soret band and the appearance of intense A-terms corresponding to the α band (Figure 7b). They all resembled those of the typical bis-histidyl hemochrome, for example, cytochrome *b*₅,³⁴ soluble guanylylase,^{35a} and bis-imidazole-bound protoheme,^{22,26,35b} as well as Hpx.^{1b} It can be concluded that rHSA(HF/R186H)-heme produced a strong six-coordinate low-spin ferrous complex under an Ar atmosphere. Unfortunately, the ferrous forms of both rHSA(HF/R186H)-heme and rHSA(HL/L182H)-heme were readily autoxidized upon the addition of O₂ gas. It is known that bis-histidyl hemochromes are rapidly oxidized by O₂ via an outer-sphere mechanism.²¹ We have demonstrated that this also applies to our artificial hemoprotein, the rHSA(mutant)-heme system.

O₂ and CO Binding Parameters for R186L and R186F Mutants. We have clearly shown that the O₂ binding equilibrium and kinetics of rHSA-heme complexes may be significantly enhanced by site-directed mutagenesis. In fact, the O₂ binding affinity of the rHSA(HL/L185N)-heme complex (1 Torr) was shown to be similar to those of Mb and the high-affinity R-state of Hb(α). However, for saline solutions of artificial rHSA-heme complexes to provide effective lung-to-tissue O₂ transport in vivo, the affinity should be reduced to render it more similar to the affinity of human RBC ($P_{1/2}^{O_2}$ 8 Torr).³² This requires an O₂ binding affinity that is intermediate between the values observed for rHSA(HL)-heme and rHSA(HL/L185N)-heme.

Both site-directed mutagenesis and synthetic porphyrin approaches have previously shown that an effective way to diminish the O₂ binding affinity of the heme is to introduce a

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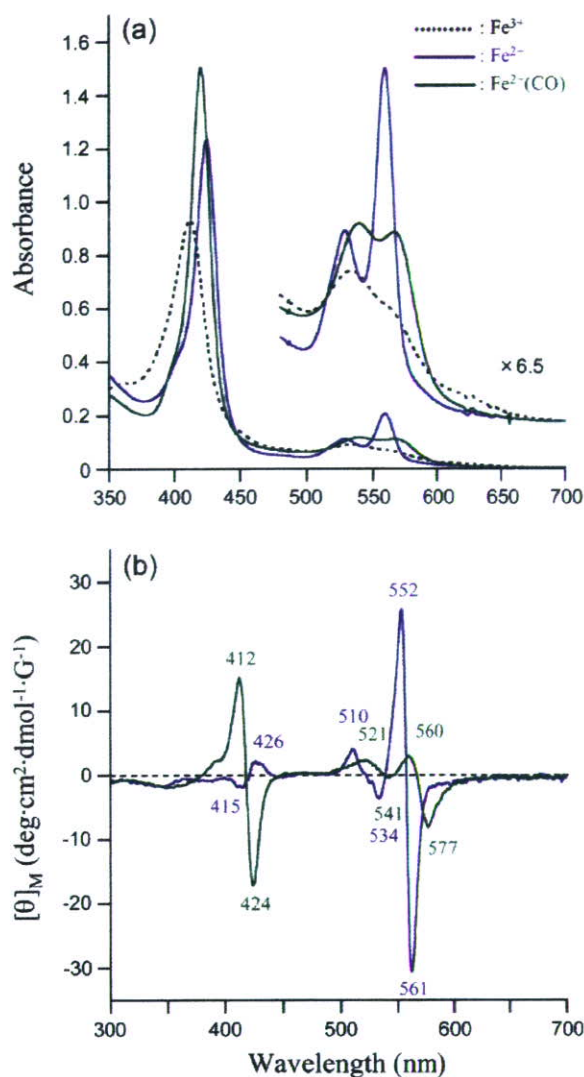


Figure 7. (a) UV-vis and (b) MCD spectral changes of rHSA(HF/R186H)-heme complex in 50 mM potassium phosphate buffered solution (pH 7.0, 22 °C).

hydrophobic amino acid (or substituent) around the O₂ binding site.^{13,14,17–19} We expected that increasing the hydrophobicity of the distal side of the heme pocket by insertion of a nonpolar residue would reduce the O₂ binding affinity of the rHSA-heme complex. The most suitable position for this introduction could be at Arg-186, which is the entrance of the heme pocket and is rather close to the central Fe²⁺ ion.

Thus, we designed new triple mutants rHSA(HL/R186L) and rHSA(HL/R186F) in an effort to prepare rHSA-based artificial hemoproteins having the same O₂ binding affinity as human RBC (Figure 9). An important structural factor in these mutants is Y161L, which is likely to allow rotation of the isopropyl group of Leu-185 above the O₂ coordination site.

MCD spectra in the Soret band region of ferric rHSA(HL/R186L)-hemin and rHSA(HL/R186F)-hemin both showed very low intensity, essentially the same as that observed for rHSA(HL)-hemin (Figure 8). The reduced ferrous form demonstrated the characteristic UV-vis absorption and MCD spectra of the five-N-coordinate high-spin complex under an Ar atmosphere (Table 1; Figure S3, Supporting Information).

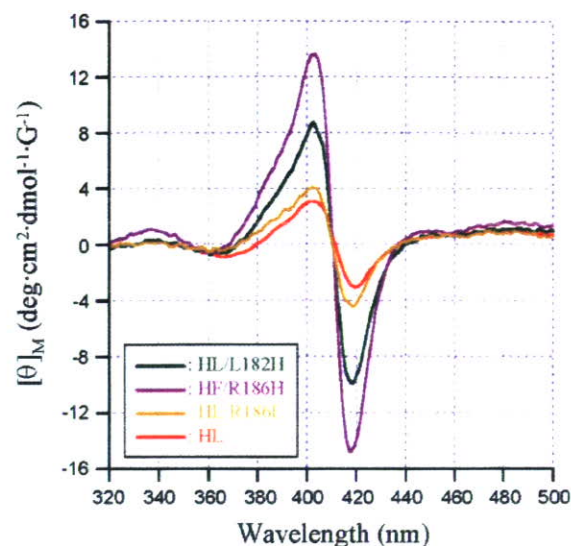


Figure 8. MCD spectra of rHSA(182-mutant)-hemin and rHSA(186-mutant)-hemin complexes in 50 mM potassium phosphate buffered solution (pH 7.0, 22 °C).

Upon bubbling of O₂ gas through the solutions, the spectral patterns were shifted to that of the O₂ adduct complex. The distinct features of all the spectra were quite similar to those of rHSA(HL)-heme.

Following laser flash photolysis, the absorption decays associated with O₂ recombination with rHSA(HL/R186L)-heme and rHSA(HL/R186F)-heme were monophasic, which suggests that the distal space in the pocket is uniform, in contrast to the L185N mutants. The kinetics for CO rebinding were still composed of double exponentials, consistent with the existence of two different geometries of the axial His-142 coordination to the central Fe²⁺ ion of the heme.

We previously showed that rHSA(HF) binds O₂ with significantly higher affinity than rHSA(HL) and reasoned that the presence of Leu rather than Phe at position 161 allowed a downward rotation of the adjacent L185 side chain that restricted access to the O₂ binding site on the heme group and reduced the affinity by a factor of 6 (Table 2).^{11b} Strikingly, however, insertion of Leu or Phe at position 186 in the presence of Leu-161 [as in rHSA(HL/R186L)-heme and rHSA(HL/R186F)-heme complexes] yielded $k_{on}^{O_2}$ and k_{on}^{CO} values that were 3–4-fold higher than those of rHSA(HL)-heme. The presence of a hydrophobic residue at position 186 may restrict the mobility of Leu-185 and thereby enhance access to the O₂ binding site (Figure 9).

Overall, the O₂ and CO binding parameters of rHSA(HL/R186L)-heme and rHSA(HL/R186F)-heme were more similar to those of rHSA(HF)-heme. In species I, for example, the k_{off}^{CO} values were almost identical, which again implies unhindered axial coordination structures of His-142 to the heme; as a result, the CO binding affinities of these triple mutants ($P_{1/2}^{CO}$ 0.0010–0.0018 Torr) were close to that of the rHSA(HF)-heme complex. In contrast, the O₂ dissociation rate constants of rHSA(HL/R186L)-heme and rHSA(HL/R186F)-heme were 3–4-fold higher than found for rHSA(HF)-heme, which modestly reduced the O₂ binding affinities (higher $P_{1/2}^{O_2}$). This could be due to the increase in the hydrophobicity in the distal pocket. Crucially, the O₂ binding affinities of rHSA(HL/

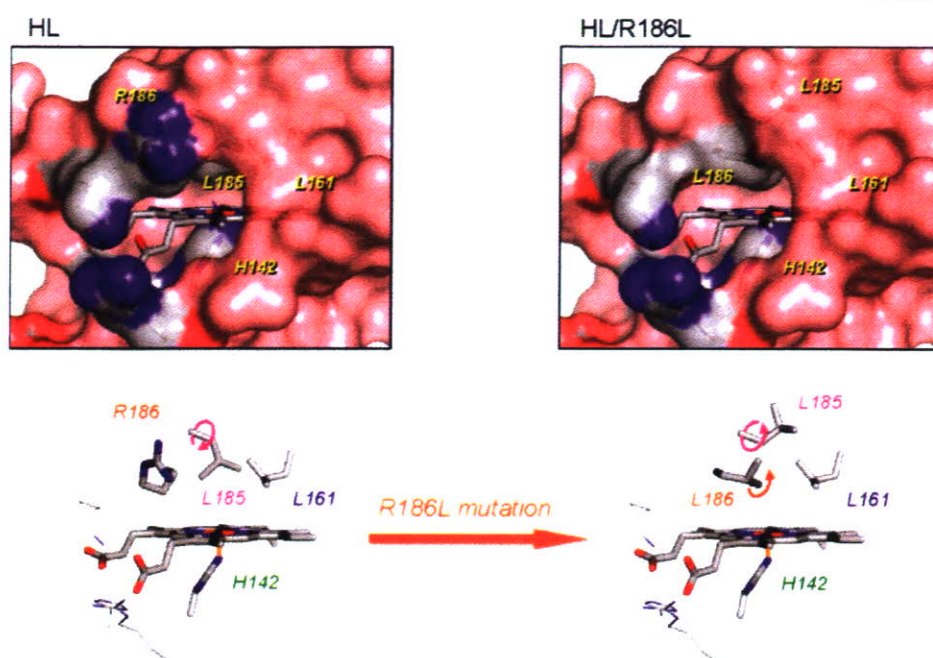


Figure 9. Structural models of rHSA(HL)–heme and rHSA(HL/R186L)–heme complexes. Introduction of R186L mutation may induce upward rotation of the L185 residue.

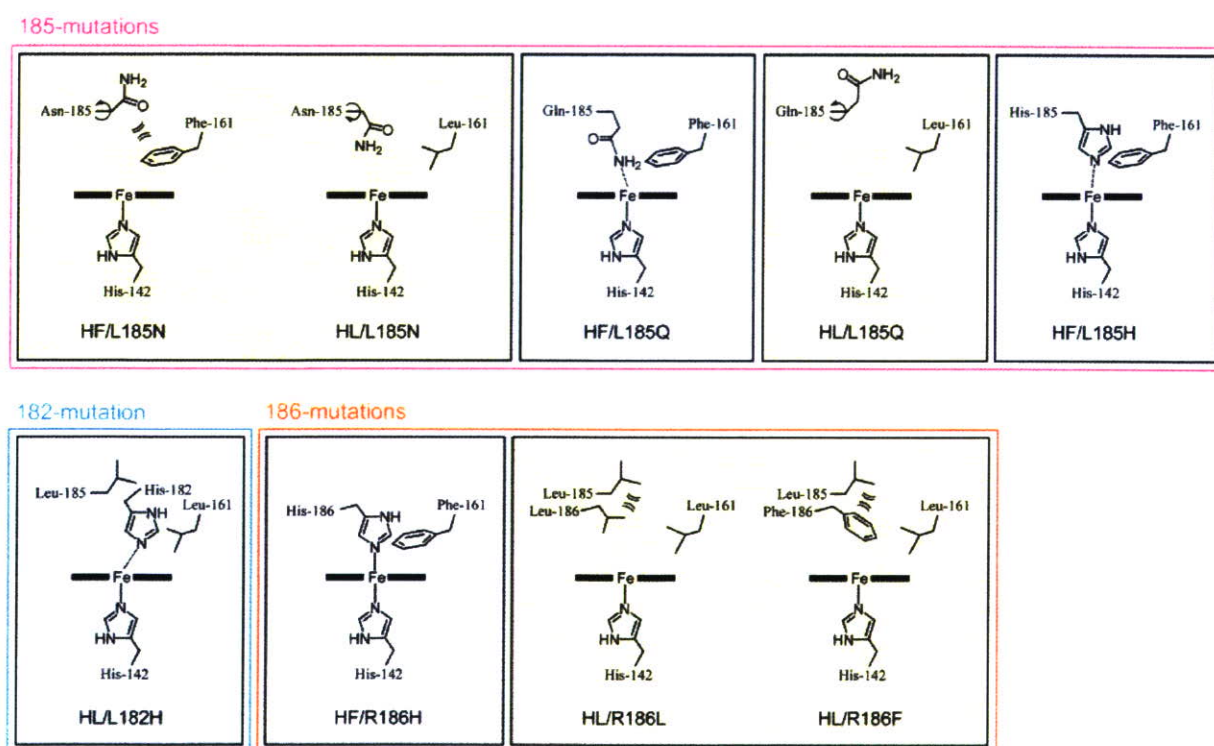


Figure 10. Schematic illustrations of the engineered distal amino acids in the heme pocket of rHSA(triple mutant)–heme: (yellow) five-coordinate high-spin ferrous complex; (blue) five-coordinate high-spin complex including six-coordinate low-spin ferrous complex; (pink) six-coordinate low-spin ferrous complex.

R186L)–heme ($P_{1/2}^{O_2}$ 10 Torr) and rHSA(HL/R186F)–heme ($P_{1/2}^{O_2}$ 9 Torr) are essentially indistinguishable from that of human RBC ($P_{1/2}^{O_2}$ 8 Torr). These results show that there are several different combinations of mutations that can confer the RBC-like O_2 binding affinity to the prosthetic heme group.

Another possibility, yet to be explored, is that insertion of a proximal His into the 186 position would construct a distal pocket on the opposite side of the porphyrin plane (the Ile-142 side), that would provide somewhat different O_2 binding properties of the heme.

Conclusions

Transport of O₂ by rHSA–heme complexes could be of great clinical importance, not only as a blood alternative but also as an O₂-providing therapeutic fluid. Such a synthetic compound has the potential advantage of not having to be matched to the recipient's blood type; moreover, it could be prepared in controlled facilities without viral contamination.

We have previously demonstrated that rHSA–heme complexes can be engineered to bind O₂ reversibly;¹¹ however, these complexes did not display optimal O₂ binding affinities. By use of structure-based mutagenesis of HSA combined with chemical modification of the synthetic iron–porphyrin, we have attempted to modify the heme pocket architecture so as to refine the O₂ binding properties of rHSA–heme complexes. By focusing on modifications on the distal side of the heme binding pocket in rHSA, we have successfully engineered distinct rHSA(triple mutant)–heme complexes with a broad range of O₂ binding affinities. Schematic illustrations of the engineered distal amino acids in the heme pocket of the different rHSA mutants are shown in Figure 10. These include mutants such as rHSA(HL/L185N) with affinities that mimic the high affinity of Hb(α) ($P_{1/2}^{O_2}$ 0.24 Torr) and others [e.g., rHSA(HL/R186L)] with affinities similar to that of human RBC ($P_{1/2}^{O_2}$ 8 Torr).

The highest affinity mutants rHSA(HL/L185N) and rHSA(HF/L185N) both contain Asn-185, which has a short amide side chain that significantly enhances the O₂ binding affinity, particularly when the neighboring amino acid is Leu-161. The N–H bond of the Asn-185 may face the terminal oxygen atom of the Fe–O₂ moiety, providing an amide dipole that stabilizes the O₂ binding to the heme. This interpretation is consistent with the findings of Chang et al.,³⁶ who first demonstrated that the dipole–dipole interaction between the Fe–O₂ and amide group can produce kinetic and thermodynamic control of the dioxygenation of the model hemes. In contrast, introduction of the larger Gln and His side chains at position 185 partly provided a six-coordinate heme character and therefore did not stabilize O₂ binding.

In a different approach, substitution of the polar Arg-186 at the entrance of the heme pocket with Leu or Phe caused a useful reduction in the O₂ binding affinity, yielding $P_{1/2}^{O_2}$ values that are very close to that of the human RBC and therefore well adapted for O₂ transport in vivo (Table 2). The impact of these substitutions may be due to their interaction with the adjacent residue, L185, which results in enhanced access to the O₂ binding site.

Other mutations were deleterious to O₂ binding but nevertheless produced complexes that might have other uses. For example, rHSA(HF/R186H)–heme formed a typical bis-histidyl Fe³⁺ or Fe²⁺ complex. In the circulation, free hemin is known to participate in the Fenton reaction to produce the highly toxic hydroxyl radical. However, it is sequestered by Hpx, in which the bis-histidyl coordination tightly fixes the hemin with the highest binding affinity of any known protein.¹² In the same manner, rHSA(HF/R186H) has a bis-histidine clamp for hemin and might conceivably be exploited as an antioxidant reagent to protect the body from oxidative damage after blood heme overload.

On the other hand, it would be of great importance to study the NO binding property of rHSA(mutant)–heme for practical medical applications. Some of the Hb-based blood substitutes leak through the vascular endothelium and capture the endothelial-derived relaxing factor, NO, that elicits an acute increase in blood pressure by vasoconstriction.³⁷ Our rHSA(mutant)–heme would bind NO in the same way as Hb, but it would not induce such hypertension, because the albumin carrier has low permeability through the muscle capillary pore.³⁸

Ultimately, to fully understand the structural basis of the effects of the various mutations on O₂ binding, it will be necessary to examine the structural details of the heme binding pocket. Crystal structural analysis of the rHSA(mutant)–heme complexes is now underway. Structural information should enhance our ability to design mutations that will further optimize the O₂ binding properties of these complexes.

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Supporting Information Available: UV–Visible absorption and MCD spectra of rHSA(HL/L185Q)–heme, absorption decay of O₂ rebinding to rHSA(HL/L185N)–heme after laser flash photolysis, and MCD spectra of rHSA(HL/R186F)–heme complex. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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Heme Pocket Architecture in Human Serum Albumin: Regulation of O₂ Binding Affinity of a Prosthetic Heme Group by Site-Directed Mutagenesis

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Summary: We present the O₂ binding properties of recombinant human serum albumin (rHSA) mutants complexed with an iron(II) protoporphyrin IX as a prosthetic heme group. Iron(III) protoporphyrin IX (hemin) is bound within subdomain IB of HSA with weak axial coordination by Tyr-161. In order to confer O₂ binding capability to this naturally occurring hemoprotein: (i) a proximal histidine was introduced into position Ile-142; and (ii) the coordinated Tyr-161 was replaced with hydrophobic Leu using site-directed mutagenesis. It provided a recombinant HSA double-mutant [rHSA(I142H/Y161L) = rHSA(HL)]. The rHSA(HL)-heme formed a ferrous five-coordinate high-spin complex with axial ligation of His-142 under an Ar atmosphere. This artificial hemoprotein binds O₂ at room temperature. Laser flash photolysis experiments demonstrated that O₂ rebinidng to rHSA(HL)-heme displays monophasic kinetics, whereas the CO recombination process obeyed a double-exponential pattern. This might be attributable to the two different geometries of the axial imidazole coordination arising from the two orientations of the porphyrin plane in the heme pocket. The O₂ binding affinity of rHSA(HL)-heme was considerably lower than those of R-state hemoglobin (Hb) and myoglobin (Mb), principally because of the high O₂ dissociation rate constant. The third mutations have been introduced into the distal side of the heme (at position Leu-185 or Arg-186) to increase the O₂ binding affinity. The rHSA(HL/L185N)-heme showed high O₂ binding affinity ($P_{1/2}^{O_2}$: 1 Torr), which is 18-fold greater than that of the original double mutant rHSA(HL)-heme and which is rather close to those of Hb (R-state) and Mb. Furthermore, replacement of polar Arg-186 with Leu or Phe adjusted the O₂ binding affinity ($P_{1/2}^{O_2}$) to 10 Torr, which is almost equivalent to value for human red blood cells.

Keywords: biomimetics; heme; human seruma albumin; O₂ binidng; proteins

Introduction

Human serum albumin (HSA), the most abundant plasma protein (4–5 g/dl) in our circulatory system, is characterized by its remarkable ability to bind widely various endogenous and exogenous compounds^[1] such as fatty acids, bilirubin, bile acids, thyroxine,^[2,3] and a wide range of drugs.^[4] Hemin [iron(III) protoporphyrin IX] released from methemoglobin is also captured by HSA with a high binding constant ($K \approx 10^8 \text{ M}^{-1}$).^[5] This strong affinity of HSA for hemin has stimulated efforts to develop albumin as an artificial hemoprotein which can mimic the O₂ binding ability of hemoglobin (Hb) and myoglobin (Mb).^[6,7] HSA consists of a helical monomer of 66.5 kDa containing three homologous domains (I–III), each of which comprises of A and B subdomains.^[8] Crystallographic studies have revealed that hemin is bound within a narrow D-shaped hydrophobic cavity in subdomain IB with axial coordination of Tyr-161 to the central ferric ion and electrostatic interactions between the porphyrin propionates and a triad of basic amino acid residues (Arg-114, His-146 and Lys-190) (Figure 1).^[9,10] In terms of the general hydrophobicity of this α -helical heme pocket, the subdomain IB of HSA potentially has similar features to the heme binding site of Hb or Mb. However, if one reduces the HSA–hemin to obtain the ferrous complex, it is rapidly oxidized by O₂, even at low temperature, because HSA lacks the proximal histidine that enables the prosthetic heme group to bind O₂ and serves to regulate the O₂ binding affinity (Figure 1). In order to confer the O₂ binding capability to this

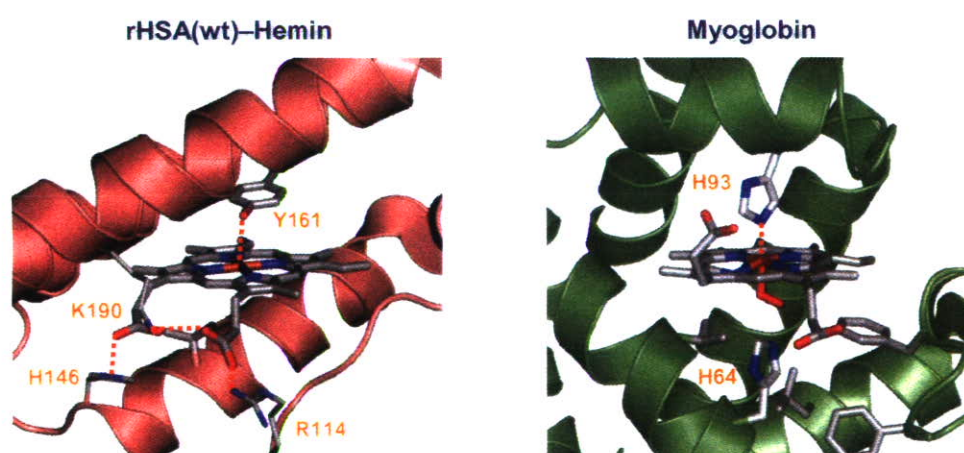


Figure 1. Heme pocket structure in subdomain IB of HSA (left; 1O9X from ref. 9) and heme pocket structure of Mb (1MBO).

naturally occurring hemoprotein, we have introduced a proximal histidine into the heme binding site of HSA by site-directed mutagenesis; it would provide axial coordination to the central ferrous ion of the heme and thereby promote O₂ binding.^[11] Moreover, to modulate its O₂ binding affinity, we have added further modification to the distal side of the heme. The O₂ binding properties of several rHSA(mutant)–heme complexes have been characterized kinetically and compared to those of the natural Hb, Mb, and red blood cells (RBC). We have shown that our mutagenesis approach can create a new class of albumin-based artificial hemoprotein which would serve as an O₂ carrier.

Results and Discussion

Double-mutations to confer the O₂ binding capability

The detailed structure of the heme binding site in HSA revealed by crystallographic studies allows the design of mutagenesis experiments to construct a tailor-made heme pocket for O₂ binding.^[9,10] In fact, Tyr-161 was the first candidate to introduce a proximal histidine (Figure 1). However, the Y161H mutation was not done because our simulation indicated that the distance from His-161 to the central Fe would be too great (4.0 Å). Instead, modeling experiments suggested that the favorable positions for the axial imidazole insertion would be Ile-142 (Figure 2a). The N(histidine)–Fe distance was estimated as 2.31 Å for H142 (compared to 2.18 Å for Mb). We therefore designed a recombinant HSA (rHSA) double-mutant I142H/Y161L [= rHSA(HL)].

The specific mutations were introduced into the HSA coding region in the plasmid vector (pHIL-D2 HSA) using the QuikChange mutagenesis kit (Stratagene), and clones were expressed in the yeast *Pichia pastoris* (Invitrogen Corp.). The rHSA–hemin complexes were prepared fundamentally according to our previously reported procedures.^[9,11]

The MCD spectra of the rHSA(HL)–hemin showed S-shaped patterns in the Soret band region, which resembled that of ferric Mb.^[12] One water molecule is known to coordinate axially to the sixth position of the central ferric ion of the hemin in metMb. Our MCD result suggests that the rHSA(HL)–hemin is also in a ferric high-spin complex, having a water molecule as the sixth ligand.