Direct observation of ligand migration within human hemoglobin at work

Naoya Shibayama\(^1\), Ayana Sato-Tomita\(^9\), Mio Ohkii\(^8\), Kouhei Ichiyanagia\(^9\), and Sam-Yong Parkc

\(^*\)Division of Biophysics, Department of Physiology, Jichi Medical University, Tochigi, 329-0048 Shimotsuke, Japan;\(^\#\)Research Complex at Harwell, Rutherford Appleton Laboratory, Harwell, Didcot, Oxon OX11 0FA, United Kingdom; and \(^\circ\)Drug Design Laboratory, Graduate School of Medical Life Science, Yokohama City University, Tsurumi, 230-0045 Yokohama, Japan

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Hemoglobin is one of the best-characterized proteins with respect to structure and function, but the internal ligand diffusion pathways remain obscure and controversial. Here we captured the CO migration processes in the tense (T), relaxed (R), and second relaxed (R2) quaternary structures of human hemoglobin by crystallography using a high-repetition pulsed laser technique at cryogenic temperatures. We found that in each quaternary structure, the photodissociated CO molecules migrate along distinct pathways in the \(\alpha\) and \(\beta\) subunits by hopping between the internal cavities with correlated side chain motions of large nonpolar residues, such as \(\alpha\text{Trp}(A12), \alpha\text{105Leu}(G12), \beta\text{J15Trp}(A12),\) and \(\beta\text{I1Phe}(E15)\). We also observe electron density evidence for the distal histidine \(\alpha\text{58/63His(E7)}\) swing-out motion regardless of the quaternary structure, although less evident in \(\alpha\) subunits than in \(\beta\) subunits, suggesting that some CO molecules have escaped directly through the E7 gate. Remarkably, in T-state Fe(II)-Ni(II) hybrid hemoglobins in which either the \(\alpha\) or \(\beta\) subunits contain Ni(II) heme that cannot bind CO, the photodissociated CO molecules not only dock at the cavities in the original Fe(II) subunit, but also escape from the protein matrix and enter the cavities in the adjacent Ni(II) subunit even at 95 K, demonstrating the high gas permeability and porosity of the hemoglobin molecule. Our results provide a comprehensive picture of ligand movements in hemoglobin and highlight the relevance of cavities, nonpolar residues, and distal histidines in facilitating the ligand migration.

Hemoglobin | ligand migration | X-ray crystallography | allostery

Human hemoglobin (Hb), an \(\alpha_2\beta_2\) tetrameric oxygen transport protein that binds gaseous ligands such as O\(_2\) and CO cooperatively at the four heme iron, serves as a model for studying protein–ligand interactions (Fig. 1A). After the pioneering work by Perutz (1), the mechanisms of Hb and its reaction with ligands have been explained based on the crystal structures of the deoxy tense (T) and fully liganded relaxed (R) quaternary states (2–5). However, a fundamental question still under debate concerns the pathways for ligands from the solvent to the heme iron, since crystal structures display no obvious gas channels leading to the heme sites (Fig. 1B).

Previous crystallographic analysis of T-state deoxy Hb under pressurized Xe identified the positions of Xe binding cavities in the \(\alpha\) and \(\beta\) subunits (Fig. 1C and D) (6). Four are located in \(\alpha\), labeled \(\alpha\text{Xe1}, \alpha\text{Xe2}, \alpha\text{Xe3},\) and \(\alpha\text{Xe6},\) and two are in \(\beta, \beta\text{Xe1}\) and \(\beta\text{Xe2}\), using the same nomenclature as in the previous study (6). It was suggested that these internal cavities may play a role as transient areas for ligands migrating through the protein interior in both subunits of human Hb. In support of this idea, recent molecular dynamics (MD) simulations have demonstrated that kinetically accessible ligand diffusion tunnels in Hb encompass the reported Xe cavities regardless of the quaternary structure (7, 8). On the other hand, however, experimental data on CO rebinding kinetics after photolysis of distal pocket mutants of Hb bound to CO (COHb) do not support this idea, but instead suggest that most ligands enter and escape directly through the distal His(E7) gate pathway (9, 10). Therefore, controversy remains as to whether the Xe-binding cavities play a functional role in Hb.

For directly observing the ligand migration in Hb, X-ray crystallographic analysis of COHb after photolysis is obviously preferred but is quite challenging and complex, due to the relatively low apparent quantum yield for the photolysis of COHb (11), as well as the need to address the quaternary structure dependence of gas diffusion as predicted by MD simulations (7). In addition, excitation light transfer to the sample, together with the rapid kinetics of CO recombination, place limits on crystal thickness and measurement temperature. To date, only two crystallographic approaches have been reported on this topic for Hb. One is a cryogenic approach to increase the level of CO photolysis by lowering the temperature to 25 to 35 K, where CO rebinding is very slow and virtually negligible during the measurement (12). It has been shown that continuous illumination of crystals of the T and R states of human Hb allows a high level of CO photolysis, but most of the photodissociated CO molecules reside at the primary docking site (located ~3.5 Å from the iron) within the distal heme pocket (DP), with the exception of a weak CO electron density present in \(\beta\text{Xe2}\) (located ~8.5 Å from the iron) in the T state (12). While near-liquid helium temperatures

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1To whom correspondence may be addressed. Email: shibayam@jichi.ac.jp.

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Significance

Human hemoglobin is the textbook example of the stereochemistry of an allosteric protein and of the exquisite control that a protein can exert over ligand binding. However, the fundamental basis by which the protein facilitates the ligand movement remains unknown. In this study, we used cryogenic X-ray crystallography and a high-repetition pulsed laser irradiation technique to elucidate the atomic details of ligand migration processes in hemoglobin after photolysis of the bound CO. Our data clarify the distinct CO migration pathways in the individual subunits of hemoglobin and unravel the functional roles of the internal cavities and neighboring amino acid residues in ligand exit and entry. Our results also demonstrate the high gas permeability and porosity of hemoglobin, facilitating O\(_2\) delivery.
are beneficial for impeding CO rebinding by decreasing the energy of CO, they also impede migration of the CO from the DP, as previously reported for myoglobin (Mb) (13, 14).

The second approach is the use of room temperature (288 K) time-resolved Laue crystallography to track the structural evolution of R-state COHb at a set of time delays between the pump laser pulse and the probe X-ray pulse (i.e., 100 ps, 1 ns, 10 ns, 100 ns, 1 μs, and 10 μs) (15). Although ligand migration and protein relaxation are expected to occur at room temperature, the experiment incurred the problem of a rather small amount of dissociated CO even at 100 ps after the pulse laser irradiation. To deal with this problem, a dataset corresponding to a hypothetical fully photolyzed crystal was calculated by linear extrapolation of structure-factor amplitudes using the scalar approximation. Specifically, the measured structure-factor amplitudes were extrapolated to 100% photolysis by assuming a photolysis level of 15%, where bound CO vanished from the extrapolated maps for both α and β subunits by visual assessment. The generated 100-ps maps showed that the dissociated CO molecules moved 1.5 to 2.0 Å away from the iron toward the primary docking site. However, no electron density of the CO was found in any other locations within the protein throughout the measurement period, which is inconsistent with the MD simulations. Clearly, there remains a considerable need for additional investigation on Hb.

In this study, we used a high-repetition (10 kHz) 1.2-ns-pulsed 532-nm Nd:YAG laser at a temperature range of 95 to 140 K, where the photodissociated CO overcomes the initial energy barrier and gradually migrates from the DP to more remote cavities in the protein during repeating dissociation and geminate recombination (16, 17). We also use crystals of the T, R, and R2 quaternary states of human Hb to explore the effect of quaternary structure on ligand migration. Note that the structural differences between R and R2 are as large as those between R and T, and that recent MD simulations suggest that ligand escape pathways vary somewhat among T, R, and R2 (7).

Results and Discussion

T-State Crystals of CO-Bound Fe(II)-Ni(II) Hybrid Hbs. We first investigated the CO migration process in the T quaternary structure of human Hb using crystals of two symmetric CO-bound Fe(II)-Ni(II) hybrid Hbs, XL[α(Fe-CO)β(Ni)]2 and XL[α(Ni)β(Fe-CO)]2, in which either the α or β subunits contain Ni(II) heme and the two αβ dimers are cross-linked by a fumaryl group between the two β2Lys(EF6) residues. It has been shown that these doubly CO-liganded hybrids yield well-diffractionally, optically thin iso-
morphous T-state crystals (12). It has also already been established that Ni(II)-heme binds neither O2 nor CO and mimics the deoxy Fe(II)-heme with respect to the effects on the functional properties of the adjacent Fe(II) subunits in the same protein (18, 19), and that the ββ fumaryl cross-link between the two β2Lys(EF6) residues has little affect on the structure and function of Hb (20).

The X-ray diffraction datasets for the T-state crystals of XL [α(Fe-CO)β(Ni)]2 and XL[α(Ni)β(Fe-CO)]2 were collected in a 95 K nitrogen gas stream under continuous irradiation by 10-kHz pulsed laser light with an average power density of 47 mW mm−2, along with control datasets on the CO-bound states without laser irradiation (SI Appendix, Table S1). Both crystals diffracted to 1.40- to 1.45-Å resolution with (light) and without (dark) laser irradiation, although relatively small crystals of 15 to 30 μm thickness were used for light transfer. The 2Foobs − Fcalc electron density maps around the α and β hemes in the dark and light structures at 95 K without (dark) and with (light) laser irradiation of 60 min demonstrate the initially bound CO and the achievement of nearly full photolysis, respectively (Fig. 2). Interestingly, in the α subunits, a detectable amount of CO exists in the primary docking site within the DP (located ~2.9 Å from the iron) after photolysis, whereas in the β subunits, a smaller amount of CO remains in the DP. This is consistent with the time-resolved Laue crystallographic data showing much faster CO escape from βDP than from αDP on photolysis (15), and is in line with the MD simulations indicating a smaller relative population of photolyzed ligand in βDP than in αDP (7).
The CO migration and the protein structural changes associated with photolysis are more clearly seen in the $|F_{\text{obs}}|_{\text{light}} - |F_{\text{obs}}|_{\text{dark}}$ difference Fourier electron density maps obtained from the same set of crystals (Fig. 3). The maps contoured at ±3.5σ confirm the loss of bound CO, the out-of-plane motion of the iron toward the proximal side, and the movement of the distal histidine α58β63His(E7) toward the site previously occupied by CO in both Fe(II) subunits (Fig. 3 A and D). The concomitant movement of the F-helix is more significant in α(Fe) subunits than in β(Fe) subunits. Moreover, the β heme undergoes a deligation-linked sliding from pyrrole C to B. These observations highlight with sticks. (Image)

Unexpectedly, we find that in XL[α(Fe-CO)]β(Ni)2, the dissociated CO molecules not only dock at the cavities in the original α(Fe-CO) subunits (Fig. 3A), but also escape from the protein and enter the cavities in the adjacent β(Ni) subunits even at 95 K (Fig. 3B). As the same situation is seen with the counterpart hybrid XL[α(Ni)]β(Fe-CO)2 (Fig. 3 C and D), we discuss the invasion of CO into the Ni(II) subunits after the description of the CO migration pathway in the β(Fe-CO) subunits of XL[α(Ni)]β(Fe-CO)2.

CO Migration in α(Fe) Subunits in the T-State. In XL[α(Fe-CO)]β(Ni)2, in addition to the electron density of the photodissociated CO remaining in αDP as mentioned above (SI Appendix, Fig. S1A), a strong electron density feature definitely corresponding to CO is observed in αXe2 among helices A, B, E, and G (Fig. 3A and see Fig. 1C), surrounded by hydrophobic amino acid residues, such as α14Trp(A12), α17Val(A15), α21Ala(B2), α24Tyr(B5), and α109Leu(G16). This site is relatively close to the solvent and is not shielded by the other subunits of the tetramer. Concomitantly with the appearance of CO, the side chains of α14Trp(A12) and α105Leu(G12) move in the direction of αXe1 as a result of a collision of CO (Fig. 3A). In addition, a relatively weak electron density feature of CO appears in αXe1 (Fig. 3A). According to the displacement of α14Trp(A12), the side chain distance between α14Trp(A12) and α17Val(A15) expands to form a transient channel for escape of CO to the solvent. These observations strongly suggest that a likely pathway for CO migration within T-state subunits is DP $\rightarrow$ αXe2 $\rightarrow$ solvent. This pathway is consistent with a recent MD study on O2 migration within human Hb, showing that a high percentage of ligand molecules escape via αXe2 in both T and R (7, 8).

CO Migration in β(Fe) Subunits in the T-State. In XL[α(Ni)]β(Fe-CO)2, the photodissociated CO is clearly observed in βXe2 between helices B and E (Figs. 1D and 3D), circumscribed by the hydrophobic side chains of β28Leu(B10), β67Val(E11), β68Leu(E12), and β106Leu(G8). It is likely that the dissociated CO has overcome the β67Val(E11) barrier to migrate into βXe2 in the back of DP. In our previous crystallographic study on the same T-state hybrid Hbs at 25 K, although most of the dissociated CO remained in DP in each subunit, a weak electron density peak was detected in βXe2, in agreement with our present results.

In XL[α(Ni)]β(Fe-CO)2 at 95 K, another positive electron density feature of CO is present in βXe1 between helix G, helix H, and the β heme (Figs. 3D and 4B), surrounded by the hydrophobic amino acid side chains of β71Phe(E15), β103Phe(G5), β134Val(H12) and β137Val(H15) and the vinyl group of heme.
of the photodissociated CO (Fig. 4B). The replacement of water with CO results in a dumbbell-shaped positive feature of CO and a weak spherical negative feature of water, as shown in Fig. 4B. Similar electron density features are observed in the counterpart hybrid XLa[α(Fe-CO)]β(Ni)]2 (Fig. A4, and SI Appendix, Fig. S2). A previous crystallographic study on T-state deoxy Hb under pressurized Xe also reported the replacement of this water by Xe1 in β subunits (6).

Taken together, our results suggest that a likely CO migration pathway within T-state β subunits is DP → βXe2 → βXe1 → solvent (Fig. 3D). This pathway agrees well with the MD study, showing that a high percentage of ligand molecules escape via the β-internal tunnel encompassing βXe2 and βXe1 in both T and R (7). The MD simulations also indicate that the open conformation of β71Phe(E15), which is favored in T and disfavored in R or R2, permits ligands to migrate between βXe1 and the central water-filled cavity of Hb (7). Consistent with this calculation, the \( F_{\text{obs}} - F_{\text{cal}} \) dark map around β71Phe(E15) in T-state XLα(Ni)β(Fe-CO)]2 provides evidence for its sparsely-populated, closed conformation moving to a position of the open conformation on photolysis (Fig. 3D, Inset, lower right).

It is important to point out here that the difference Fourier map for T-state XLa[α(Ni)]β(Fe-CO)]2 suggests a transient occurrence of a swing-out motion of the distal β65His(E7) on photolysis (Fig. 3D, Inset, lower left), although no such motion can be detected for the distal α58His(E7) in the counterpart hybrid XLα(Fe-CO)]β(Ni)]2. Our structural analyses on the R and R2 states of Hb provide more evidence of the distal α58/β65His(E7) swing-out motion (as described below in more detail), suggesting that at least some CO molecules can escape directly through a transient E7 channel as well as through an internal cavity network.

**CO Invasion into Ni(II) Subunits Indicates the High Gas Permeability of the T State.** Regarding the CO invasion into Ni(II) subunits, the CO docking sites observed in β(Ni) subunits of XLα(Fe-CO)]β(Ni)]2 (Fig. 3B) are very similar to those in β(Fe-CO) subunits of the counterpart hybrid XLa[α(Ni)]β(Fe-CO)]2 (Fig. 3D), confirming the distribution of the photodissociated CO in β subunits of Hb. Similar confirmation can be made for α subunits by comparing the CO docking sites between α(Fe-CO) and α(Ni) subunits (Fig. 3A and C and SI Appendix, Fig. S1B). We note here that the dumbbell-shaped electron density in each Ni(II)-subunit cavity (Fig. 3 B and C) can be modeled by a CO molecule (SI Appendix, Figs. S1–S3) but not by a chain of two water molecules, because the oxygen atoms of hydrogen-bonded water molecules are too far apart.

To further demonstrate that the changes in electron density in Ni(II) subunits are due mainly to CO and not to a temperature rise caused by laser irradiation of Ni(II) heme, we performed another series of X-ray measurements using the isomorphous T-state crystal of XLα(Ni)β(Ni)]2, in which all four subunits contain Ni(II) heme and two β2LYs(EF6) residues are cross-linked by a fumaryl group. For these measurements, data collection and experimental conditions (including temperature, laser irradiation time, crystal size, and resolution) were similar to those for the crystals of XLa[α(Fe-CO)]β(Ni)]2 and XLα(Ni)]β(Fe-CO)]2 (SI Appendix, Table S2 and Fig. S4). The \( F_{\text{obs}} - F_{\text{cal}} \) dark Fourier electron density maps (contoured at similar density levels as in Fig. 3) show no indication of an increase in electron density in any cavities in XLa[α(Ni)]β(Ni)]2 with laser irradiation (SI Appendix, Fig. S5 A and B). There is only a negative electron density feature at βXe1, signifying displacement of the preexisting water molecule. These results, together with the data in Fig. 3, strongly indicate that the photodissociated CO molecules are able to exit from and reenter the protein matrix even at 95 K, demonstrating the high gas permeability and porosity of Hb. Our data also reveal that CO migration is not just a passive diffusion process but is coupled with side chain motions of key residues, such as α14Trp(A12), α105Leu(G12), β36His(E7), and β71Phe(E15) (Fig. 3 F).

The obvious question is why ligands can migrate between the subunits of Hb even in the frozen glassy state at temperatures <160 K. While the answer is not yet clear, a recent simulation study suggests that many ligand molecules exist from and enter the protein matrix via the central cavity of Hb without escaping to the bulk solvent (7). The larger central cavity in T is expected to facilitate ligand diffusion between the subunits within the tetramer compared with R/R2. The experiments described below provide additional evidence in support of this expectation.

**R- and R2-State Crystals of COHb.** We next investigate the CO migration processes in two representative relaxed quaternary structures, R and R2. Recent structural studies using X-ray crystallography (21, 22), NMR (23), and cryo-EM (24) have shown that a fully-ligated Hb exists in an ensemble of relaxed conformations, varying between the R and R2 boundaries. The R and R2 crystals used in this study are those of COHb C [β6Glu(A3) → Lys] and COHb A (wild type), respectively, both of which are of ~30 μm thickness. Mutant Hb C is used because it forms less soluble, more resistant crystals compared with Hb A (25–27) due to the β6Glu-to-Lys surface mutation, which does not alter the ligand-binding properties of Hb. We recently demonstrated that thin crystals of R-state COHb C are of high diffraction quality and suitable for photolysis experiments (28).

The X-ray diffraction datasets for the R and R2 crystals were collected at 95 K using the same procedures as for the T crystals. Note, however, that CO recombination is much faster in R/R2 than in T (29), so the photoproduct yield of R/R2 was only ~30% under the irradiation conditions used here. Therefore, we carried out additional data collection at 140 K to facilitate CO migration, based on the temperature-dependent CO migration in Mb crystals (30). At both temperatures with (light) and without (dark) laser irradiation, the R and R2 crystals diffracted to 1.40- to 1.60-Å and 1.60- to 1.70-Å resolution, respectively (SI Appendix, Table S1). As seen in the \( F_{\text{obs}} - F_{\text{cal}} \) dark difference Fourier maps of R and R2 (Figs. 5 and 6), although the CO photolysis features are weaker in R/R2 than in T, especially in β subunits with faster recombination kinetics than α (15), there are overall similarities among the difference Fourier maps of T, R, and R2, helping us confirm that the electron density features observed in R and R2 likely reflect important aspects of the CO migration processes rather than noise.
It is important to mention that when the time of laser irradiation was varied from 2 to 60 min at both temperatures, there were no significant changes in the ligand electron density in the cavities of R/R2 (SI Appendix, Table S3 and Figs. S6 to S9), providing no evidence of ligands moving from one site to another with increasing irradiation time. This is in marked contrast to the results obtained with crystals of CO-bound Mb (MbCO), which showed increasing CO density in the Xe-binding cavities with time under illumination with a 532-nm pulsed laser at 100 to 140 K (16). Thus, in the present study, we focus on the temperature-dependent evolution of the electron densities of the photodissociated CO molecules rather than on their time evolution.

**CO Migration in the R and R2 States.** As expected, in R and R2, the photodissociated CO molecules migrate to more remote sites with increasing temperature. A comparison of Fig. 5 A and C and Fig. 6 A and C clearly shows that an increase in temperature from 95 K to 140 K decreases the occupancy of CO in αXe3 close to αDP and lined by the distal α88His(E7) Cβ atom but, conversely, increases the CO occupancy in αXe2. This migration behavior of CO is in line with the MD simulations (7) showing that many of the ligand copies that enter αXe3 return to αDP (during an early stage of simulation) before migrating to the α internal tunnel encompassing αXe2 and αXe1. The simulations estimate that 18% and 33% of the ligand molecules escape to the solvent via αXe3 in R and R2, respectively, whereas 34% and 10% of escapes occur via αXe2/αXe1 in R and R2, respectively.

As may be expected, CO most likely cannot migrate between the internal cavities and the surrounding bulk solvent at 95 K, which is well below the solvent glass transition temperature (31), even though with laser irradiation the temperature is likely to be slightly higher than that nominally set using a nitrogen gas stream. Such a view may be correct for R/R2 at 95 K (Figs. 5 and 6), but not for T at 95 K, as mentioned above (Fig. 3), suggesting a higher gas permeability of T compared with R/R2. The CO distribution in T-state Hb appears to have already reached equilibrium among subunits at 95 K, with relatively unhindered diffusion of ligands via the larger central cavity. Consequently, the patterns of CO movement in T at 95 K more closely resembles those in R/R2 at 140 K rather than at 95 K.

A further detailed comparison of the difference Fourier maps of T, R, and R2 reveals subtle but important differences among them in terms of the side chain motions of key residues as well as the patterns of CO movement. For example, unlike T with β71Phe(E15) in its open conformation, in R and R2 this residue adopts a closed conformation, partially capping βXe1 before and during photolysis at 95 K (Figs. 5B and 6B). This closed conformation expands the volume of a non–Xe-binding cavity below β15Trp(A12) (blue dotted circles in Figs. 5B and 6B), allowing the entrance of CO at 95 K. Correlated side chain displacement of β15Trp(A12) is evident in R (Fig. 5B). When the temperature is raised to 140 K, the phenyl ring of β71Phe(E15) transiently switches to an open conformation (an alternative rotamer, as shown in Fig. 5G), which simultaneously ejects CO from the cavity.
below β15Trp(A12) and permits CO migration from βXe1 to βXe2 (Figs. 5D and 6D) and probably further to the central water-filled cavity of Hb, as suggested by the MD simulations (7). It is interesting to note that previous time-resolved resonance Raman spectroscopic studies on solution COHb found a transient change in the Raman lines of COHb (12) (15Trp(A12) in a very short time after photolysis (32, 33), which may relate to the observed displacements of these Trp(A12) residues as seen in Figs. 5 and 6.

There is also electron density evidence that the distal His(E7) transiently swings out of DP in both subunits in R2 at 140 K (Fig. 6 F and G) and only in β subunits in R at 140 K (Fig. 5F). No detectable displacements occur at 95 K in R and R2. We speculate that these deligation-induced transient motions of His(E7) are a sign of direct exit of CO from DP to the solvent, supporting the distal His(E7) gate hypothesis (34, 35), in which the ligand entry to and exit from DP is gated by swinging of the imidazole ring of His(E7) out of DP. It should be noted that the movement of the distal His(E7) toward the site occupied by CO is less visible in R/R2 at 140 K (Figs. 5F and 6 F and G) than in T at 95 K (Fig. 3A and D). This result is consistent with the photolysis-induced structural changes observed at 25 to 35 K (12) and also in line with the view that both liganded and unliganded heme can be accommodated equally well within the more plastic R state (36), as the strain energy retained by the distal His(E7) is much smaller in R/R2 than in T.

In addition, in R and R2, a positive electron density feature appears in a small cavity beside the α heme (blue dotted circles in Figs. 5A and C and 6C), whereas no electron density is observed in T (Fig. 3A). This site corresponds to the position of βXe1 in β subunits (Fig. 1D). However, the following facts are important to note. Previous studies of the crystal structure of R-state horse deoxy Hb (Protein Data Bank [PDB] ID code 1IBE) (36) showed that one water molecule is located at this cavity in the unliganded R state but not in the CO-liganded R state (PDB ID code 1G0B) (37), suggesting a ligation-linked water molecule displacement in R. Moreover, comparing βXe1 and the α cavity, amino acid differences at the G10 and H16 positions (i.e., β107Gly[G10] → α102Ser and β138Ala[H16] → α133Ser) decrease the size and hydrophobicity of the α cavity, making it less accessible to CO and Xe. Thus, it is likely that the observed electron density beside the α heme in R/R2 is at least partly due to a water molecule entering from the solvent on deligation of the α heme.

Comparison with Related Studies. Our present results suggest that some CO molecules can escape directly through a transient E7 channel in addition to an internal cavity network in all quaternary states T, R, and R2. Consistent with this idea, recent MD simulations by Shadrina et al. (7) showed that an average of 23%, 31%, and 46% of the ligand escaped to the solvent directly from the DP of T, R, and R2, respectively, with the remainder escaping through the interior tunnels. Moreover, previous simulations by Lucas and Guallar (38) have shown an average of 40% and 51% of the ligand escaping from the distal path of T and R2, respectively. Both simulation studies indicated a larger preference for the distal path in β subunits, especially in R2 (7, 38), in agreement with the
present findings. Recent simulations also have suggested an interesting mechanism in which the ligand can escape to the solvent directly from the DP even with the His(E7) gate closed (8). Unfortunately, our data neither support nor directly contradict this mechanism.

A recent room temperature time-resolved X-ray analysis of R-state COHb crystals after photolysis found no electron density of the photodissociated CO in any of the reported Xe cavities (35). This contradiction with the present data on the same R crystals (Fig. 5A–D) is unexplained but possibly could arise from the highly transient and dynamic nature of migrating ligand, making its detection difficult, especially at room temperature. Indeed, although the multiphasic CO geminate rebinding in the R-state COHb crystal at room temperature indicated the existence of at least one CO docking site other than DPs (15), no electron density of CO was found in any of possible sites. The existence of discrete docking sites for the photodissociated CO within Hb has also been demonstrated by a previous kinetics study that found biphasic geminate rebinding of CO to R-state Hb encapsulated in wet silica gels (39).

Finally, we should consider and discuss the lessons learned from studies of the monomeric oxygen storage protein Mb, whose ligand migration mechanism has been extensively investigated using various experimental and simulation techniques. In 1966, Perutz and Matthews (34) proposed that O2 enters mammalian Hb and Mb by a short, direct channel gated by the distal His(E7) near the solvent edge of the heme. Subsequently, the existence of internal Xe-binding cavities was discovered in the Mb crystal structure (40), leading some researchers to argue that instead of passing through the E7 channel, diatomic gaseous ligands, including O2 and CO, may enter and exit globins through apolar tunnels involving Xe sites (41). MD simulations have provided theoretical support for this idea (42, 43).

Direct evidence for movement of photodissociated CO molecules into the Xe-binding cavities in Mb comes from X-ray crystallographic studies at room and cryogenic temperatures (13, 14, 16, 30, 44–46). However, the migration of photodissociated CO between the Xe sites in MbCO crystals cannot by itself define the ligand exit pathway, because the ligand exit process involves overcoming activation barriers and the resultant high-energy transition states cannot be observed directly. Kinetic measurements on ligand entry and exit and the lifetimes of the photodissociated states are needed. Scott et al. (35) experimentally examined the effects of large-to-small amino acid substitutions on the rates of O2 entry into and exit from Mb in solution using 90 mutants at 22 different positions and found that mutations at the Xe sites and along the proposed apolar tunnels had little effect on measured rates for ligand entry and exit, although they often affected the geminate kinetics. Time-resolved room temperature X-ray measurements of MbCO crystals after photodissociation of CO showed that the lifetime of CO electron density in the Mb Xe1 cavity (located on the proximal side of the heme pocket) increased dramatically when the size of the B10 side chain (located at the DP wall) was increased (47). These findings indicate that although photodissociated ligands can migrate into the Xe-binding cavities, they must move back to the DP to exit the protein. Since the active site structures and the E7 channels in the subunits of human Hb are similar to those in Mb, we should keep in mind the possibility that, as in Mb, some CO molecules in the internal cavities may return to the DP in Hb. However, there is far less information about the functional role of apolar tunnels in human Hb, and the distal His(E7) channel is not the universal pathway for ligand entry into and exit from all globins (48). Further studies combining X-ray crystallography and mutagenesis as well as MD simulations could improve our understanding of the ligand migration mechanism of human Hb.

Conclusions

Our high-quality electron density maps provide a comprehensive picture of CO migration processes in the T, R, and R2 quaternary structures of human Hb and highlight the relevance of cavities, nonpolar residues, and the distal His(E7) in rapid ligand exit and entry in Hb. Despite a similar folding topology in both subunits of Hb, the photodissociated CO diffuses by hopping between the internal cavities along the pathway in a different direction in each subunit of all quaternary states. The distribution of CO in the cavities depends somewhat on the quaternary state. The side chain of His(E7) also acts as a transient gate for ligand in all quaternary states, although this is less evident in α subunits than in β subunits. In addition to the distal e863His(E7), α14Trp(A12), α105Leu(G12), β15Trp(A12), and β17Phe(E15) are revealed as the key residues controlling ligand migration in each quaternary state of Hb. The present results strongly suggest the existence of multiple ligand migration pathways in both subunits of Hb (7, 37, 39) and emphasize the functional relevance of the high gas permeability and porosity of the T-state Hb molecule (49, 50) in facilitating rapid O2 diffusion and delivery to the tissues.

Materials and Methods

Detailed information on the materials and methods used in this study, including sample preparation, crystallization, X-ray structural determination, and data collection and refinement statistics, are provided in SI Appendix. Data Availability. Structural data have been deposited in the Protein Data Bank (https://www.wwpdb.org/) with PDB ID codes 6K9A for T-state XL[Fe–CO][Ni], (dark, 95 K), 6KAE for T-stateXL[Fe–CO][Ni], (light 60 min, 95 K), 6KAH for T-stateXL[Fe–CO][Ni], (dark, 95 K), 6KAI for T-state XL[Fe–CO][Ni], (dark, 95 K), 6KAQ for R-state COHb C (dark, 95 K), 6KAP for R-state COHb C (light 60 min, 95 K), 6KAR for R-state COHb A (light 60 min, 140 K), 6KAS for R-state COHb A (dark, 95 K), 6KAT for R-state COHb A (light 60 min, 95 K), 6KAU for R-state COHb A (dark, 95 K), 6KAV for R-state COHb A (light 60 min, 140 K), 6LSV for R-state COHb A (light 2 min, 140 K), 6LSX for R-state COHb A (light 2 min, 95 K), 6LSY for R-state COHb A (light 2 min, 95 K), 6LVQ for T-stateXL[Fe–CO][Ni], (dark, 95 K), and 6LXQ for T-stateXL[Fe–CO][Ni], (light 60 min, 95 K) (details in SI Appendix, Tables S1 to S3).

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