Directed inter-domain motions enable the IsdH *Staphylococcus aureus* receptor to rapidly extract heme from human hemoglobin

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INTRODUCTION

*Staphylococcus aureus* is an opportunistic bacterial pathogen that causes a wide range of life-threatening illnesses such as pneumonia, meningitis, endocarditis, toxic shock syndrome, bacteremia, and septicemia [1–3]. During infections, *S. aureus* relies on iron as a nutrient, which it extracts from human hemoglobin (Hb) using the IsdH surface receptor. Heme extraction is mediated by a tridomain unit within the receptor that contains its second (N2) and third (N3) NEAT domains joined by a helical linker domain. Extraction occurs within a dynamic complex, in which receptors engage each globin chain; the N2 domain tightly binds to Hb, while substantial inter-domain motions within the receptor enable its N3 domain to transiently distort the globin’s heme pocket. Using molecular simulations, Markov modeling, and quantitative measurements of heme transfer kinetics, we show that directed inter-domain motions within the receptor play a critical role in the extraction process. The directionality of N3 domain motion and the rate of heme extraction is controlled by amino acids within a short, flexible inter-domain tether that connects the N2 and linker domains. In the wild-type receptor directed motions originating from the tether enable the N3 domain to populate configurations capable of distorting Hb’s pocket, whereas mutant receptors containing altered tethers are less able to adopt these conformers and capture heme slowly via indirect processes in which Hb first releases heme into the solvent. Thus, our results show inter-domain motions within the IsdH receptor play a critical role in its ability to extract heme from Hb and highlight the importance of directed motions by the short, unstructured, amino acid sequence connecting the domains in controlling the directionality and magnitude of these functionally important motions.
the LN3-Hb interface) (Fig. 1A) [26]. A subsequent 2.6 Å structure of the IsdB-Hb complex revealed deformation of Hb’s F-helix, which contains the hemin iron-coordinating axial His87 residue [12]. Subsequent structural studies of the IsdB-Hb complex revealed a similar mode of binding and F-helix distortion, suggesting that IsdB and IsdH extract Hb’s hemin using a related mechanism [27]. Interestingly, cellular studies suggest that IsdB is the primary Hb receptor in S. aureus, while IsdH may function to prevent Hp-mediated removal of Hb from the blood in addition to extracting its hemin [21, 28].

Biophysical, computational, and structural studies of IsdH have shed light onto the mechanism underpinning hemin extraction. Hb binding is primarily mediated by the N2 domain, as the N2-Hb interface forms with a dissociation constant (K_D) of 7.5 ± 0.1 μM that contributes ~95% of the total binding standard free energy required for complex formation. In contrast, the LN3-Hb interface in which Hb’s F-helix is partially unwound forms with much weaker affinity (K_D > mM) [9, 11, 24]. Quantitative measurements that selectively measured the rate of hemin transfer from tetrameric Hb revealed that IsdH complex accelerates hemin release from the alpha globin chain up to 13,400-fold as compared to the indirect process in which Hb first spontaneously releases hemin into the solvent, and identified two conserved receptor subsites within the Hb-LN3 interface that are important for transfer [9, 11]. Structural experiments and molecular dynamics (MD) simulations of the IsdB-Hb complex suggest that the receptor lowers the activation energy associated with hemin removal from the alpha subunit by inducing strain in the His8-Fe³⁺ axial bond and by increasing the concentration of nearby water molecules that compete with Hb’s His8’s Nε atom for the iron atom within hemin. After breakage of the His8-Fe³⁺ axial linkage the released hemin is then transferred ~12 Å to the binding pocket in IsdH’s N3 domain where its metal is coordinated via the phenol oxygen of Y642 [11, 12, 26].

NMR studies indicate that the IsdH receptor undergoes significant inter-domain motions when bound to Hb [10, 29]. The motions occur at the junction between its N2 and linker domains, suggesting that LN3 domains form a rigid unit that transiently engages and distorts Hb’s hemin pocket while the N2 domain remains affixed to each globin [29]. Here, we show using computational and experimental approaches that these inter-domain motions are functionally important and defined by a semi-disordered tether sequence that connects the N2 and linker domains (residues P466-Y475). MD simulations analyzed through Markov state models of wild-type and mutant receptors reveal that the tether plays a key role in controlling both the positioning and direction of motion of the LN3 unit relative to Hb. These directed domain motions raise the effective molar concentration of the LN3 unit near Hb to ~15 mM, enabling it to populate a low affinity Hb-LN3 interface in which Hb’s heme pocket is distorted to promote hemin release. Residues in the tether are mobile in the simulations and absent in crystal structures of Hb-IsdH complexes, yet nevertheless direct inter-domain motions by transiently adopting a closed conformation that projects the LN3 unit toward Hb. Altering the sequence of the tether disrupts its ability to direct functionally important domain motions dramatically slowing the rate at which hemin is captured.

**Figure 1.** Structure of the IsdH-Hb complex. (A) Crystal structure of the IsdH-Hb complex (PDB code: 4XS0). The N2 (red) and LN3 (orange) subunits in the receptor form contacts with alpha globin chain in Hb (gray). The tether sequence and the N2-Hb and LN3-Hb interfaces are labeled. The hemin molecule in Hb is shown in space-filling format (black). The bounds of the tether (blue) are shown as spheres. (B) The panel shows the domain organization of IsdH with the amino acid sequence of the tether shown below. The tether sequence is formed by residues P466-Y475 (blue). Also shown are primary sequences of the tether mutants explored in this study, as well as the homologous tether sequence in IsdB.

**METHODS**

**Molecular dynamics simulations.** Initial atom coordinates for the WT IsdH were taken from a 2.55 Å resolution IsdH:Hb dimer structure (PDB: 4XS0) [12]; the hemoglobin dimer was deleted, and missing residues were built. The protein was then neutralized with Na⁺ and solvated with a 150 mM NaCl buffer using the OPLS-AA force field [30, 31]. The system was parameterized using the AMBER ff14SB [32] while the heme and Fe-N bond parameters were taken from previous work [29]. The system was minimized and heated to 300 K over 20 ps while restraining the protein, then the restraints were relaxed over 100 ps. Three separate 500 ns simulations were carried out under constant temperature and pressure with a Langevin thermostat with a collision frequency of 1 ps⁻¹ and a Monte Carlo barostat set to 1 atm [33, 34]; electrostatic energies were calculated using particle-mesh Ewald while van der Waals energies were calculated with a cutoff of 10 Å. All simulations were done using the AMBER18 software suite with a timestep of 2 fs, unless otherwise specified [35]. Standard simulation analyses were performed with CPPTRAJ [36].

To further sample the interdomain motions, the N2-N3 distance and N2-H3-N3 dihedral was calculated for every 100 ps, then frames from the three simulations were clustered into 50 groups using these two projections. One structure from each group was then selected at random to seed a 500 ns simulation using the goal-oriented FAST protocol [37]; the trajectory set was used to create a temporary Markov model based on the distance and dihedral projections. Clusters in the model were selected based on a...
reward function which used their proximity to the goal state and on
the sampling of similar structures. Here, the goal state was any struc-
ture that was different than the initial crystal coordinates—that is,
states with similar coordinates as the WT IsdH:Hb dimer structures
were penalized. Selected structures were prepared and run for 100 ns,
then added to the trajectory set. 20 rounds of this sampling were pro-
duced, leading to a total of 25 µs of sampling. The resulting trajectorys
from the FAST protocol were then represented by a final Markov
model consisting of 200 states. All models were built using the
KMeans clustering algorithm and a lag time of 50 ns was selected
based on the implied timescales of the model. Distance and dihedral
measurements were done using CPPTRAJ [38], while Markov model
building and coarse-graining were done with MSMBuilder [38]. Visu-
alization was done using PyMol version 2.3.0 [39].

To sample the N3 binding process, we first formed a coarse Mar
k model by lumping the 200 microstates into 75 macrostates using
the BACE algorithm [40]. This algorithm was selected in order to
clump clusters that behaved similar in the model and to ensure that
these clumps were sufficiently different. A representative apo-IsdH structure was taken from each macrostate, then aligned the N2 domain to
the IsdH:Hb dimer structure (PDB:4X80); the IsdHN2N3 in the crys-
tal structure was then deleted, leaving the Hb structure and the aligned
IsdHN2N3. Of these 75 initial structures, 51 did not lead to structural
overlap between the N3 and Hb dimers. These structures were pre-
pared as described above and simulated for 100 ns. This set was then
used to initialized another FAST protocol of five rounds, where struc-
tures were rewarded for being similar to the WT IsdH:Hb dimer. The N3 RMSD relative to its positioning in the Hb bound crystal structure
(after aligning the a-subunits) was included as a third dimension dur-
ing the clustering. Outside the clustering dimensions selection, the
protocol followed was the same as mentioned above.

Neither the initial simulations nor the FAST protocol were able to
sample states similar to the N3:Hb interface seen in the crystal struc-
ture, thus the five aligned structures with the lowest N3 RMSD were
used to initialize steered MD simulations. The N3 RMSD was
steered towards zero over 50 ns with a force constant of 1.0
kcal/mol Å. Initial structures where the N3 RMSD was 5, 10, and 15
Å were pulled from these trajectories and used to seed three conven-
tional, unbiased 100 ns simulations—leading to 4.5 µs of sampling
states similar to IsdH:Hb dimer structure. The steered simulations
were done using NAMD2.12 [41] through the Colvars module [42],
while the seeded simulations were conducted as described above.
IsdH-Hb trajectories from the initial aligned structures, the FAST
rounds, and seeded from the steered simulations were then analyzed
by building a Markov model for each structural mutation. The con-
struction of the model was done in the same manner as the apo-IsdH
models, except with the N3 RMSD included as a third dimension in
the clustering and a lag time of 85 ns was used. Binding kinetics
were analyzed by coarse-graining the model to 25 macrostates using
BACE. Each structural mutation was built using Robetta [43] and
sampled in the same way described above. Two separate mutations
were introduced: a ten residue insertion consisting of an alternating
Gly-Ser sequence in between Pro466 and Asp467, and a ten residue
deletion (residues Pro466-Y475). Apo- and holo-IsdH mutant models
were produced following the protocol described above.

Preparation of mutant plasmids. All mutants are derived from
the parent gene encoding “wild-type” IsdH2503 as an insert at
BamHI(5')-XhoI(3') restriction sites the pET28a-derived plasmid
pSUMO, which encodes an N-terminal small-ubiquitin-like modifier
(SUMO) fusion and an N-terminal hexahistidine tag. Mutations to this
insert were introduced through QuikChange mutagenesis and single-
overlap extension (SOE) PCR using primers supplied in the supple-
mentary information. When SOE was used, the agarose-gel purified
assembled insert and plasmid backbone were digested with BamHI and
XhoI (NEB) and the insert ligated with T4 ligase. Plasmid DNA
was transformed into chemically competent BL21(DE3) E. coli cells
and plated on Luria broth (LB) agar supplemented with 50 µg/ml
kanamycin sulfate. Colonies were harvested, propagated for sequenc-
ing, and stored as stocks at -80°C in 1:1 LB:50% glycerol supple-
mented with 25 µg/ml kanamycin sulfate.

Preparation of IsdH and Hb. Mutants of apo-IsdH and apo-
Mb(H64Y/V68F) were prepared as previously reported [44–46].
Briefly, 10 mL LB supplemented with 50 µg/ml kanamycin sulfate
(LB/kan) was inoculated with E. coli BL21(DE3) cells bearing mutant
pSUMO-IsdH or pSUMO-apoMb(H64Y/V68F) plasmid and grown overnight at 37°C with 200 rpm shaking. 2 L of LB/kan were inocu-
lated with overnight culture and grown to OD600 = 0.6 with 200 rpm
shaking at 37°C. Cultures were induced with 2 mL 1 M IPTG and in-
cubated with 200 rpm shaking at 37°C for 20-24 hours. Cells were
harvested by centrifugation at 7000 rpm for 10 minutes in 500 mL
plastic collection vessels in a Beckman JA-10 centrifuge rotor. Cell
pellets from all mutants were slightly brown in color indicative E.
coli-sourced heme bound to IsdH. Cells were resuspended in 50-60
mL lysis buffer (50 mM NaHPO4, 300 mM NaCl, 10 mM imidazole,
PH 7.4) and frozen overnight at -20°C. Thawed cell pellets were
supplemented with Protease Inhibitor Cocktail, lysed by sonication,
and clarified at 15,000 rpm in a Beckman JA-20 centrifuge rotor. Clarified
lysates were applied to 5 mL HisPur Ni-NTA resin by co-incubation
at 4°C for 20 minutes. Flowthrough was eluted and bound protein
washed with 50 mL lysis buffer. Protein was denatured on-bead with
50 mL denaturing buffer (50 mM NaHPO4, 300 mM NaCl, 10 mM
imidazole, 6 M guanidine hydrochloride, PH 7.4) to release bound heme [46]. Heme was washed away with 4 x 50 mL washes of a 1:4
twice mixture of ethanol denaturing buffer. The denatured apo-protein
was refolded by an additional wash with 50 mL lysis buffer. Resin
was resuspended in ~30 mL lysis buffer and 1 mL Upl-1SUMO prote-
ase added as a 1 µg/mL solution in 50% glycerol/lysis buffer. Resin
mixture was incubated overnight with gentle rocking at 4°C to liberate
IsdH. The protein was then eluted and resin washed with 2 x 30 mL
lysis buffer, concentrated and buffer exchanged into heme transfer-su-
crose buffer (20 mM NaHPO4, 150 mM NaCl, 0.45 M sucrose, PH
7.5) using 15 mL Amicon 10 kDa cutoff centrifugal filters.

metHb0.1 was prepared as reported previously, with small modifica-
tions. Briefly, 50 mL LB supplemented with 10 µg/mL tetracycline
(LB/tet) was inoculated with E. coli BL21(DE3) cells bearing plasmid
pSGE1.1, which encodes the tetramer-stabilized Hb mutant Hb0.1.
Cultures were grown overnight at 37°C with 200 rpm shaking. 2 L of
M9 minimal medium supplemented with 10 µg/mL tetracycline was
prepared and inoculated with overnight culture and incubated at 37°C
with 200 rpm shaking to OD600 = 0.4-0.6. Hb0.1 expression was in-
duced by addition of 2 mL 8 mM hemin in 0.1 M NaOH and 2 mL 1
M IPTG. Expression was conducted at 25°C for 20-24 h with 200 rpm
shaking. Cells were harvested by centrifugation at 7000 rpm for 10
minutes in 500 mL plastic collection vessels using a Beckman JA-10
centrifuge rotor. Cells were washed twice with Hb0.1 lysis buffer (20
mM Tris, 17 mM NaCl, PH 8.5) to remove excess heme, resuspended
in 50-60 mL Hb0.1 lysis buffer and frozen overnight at -20°C. Thawed
cell mixtures were bubbled with carbon monoxide to prevent autox-
idation. Cells were treated with 1 M benzamidine HCl, and lysed
by sonication. The crude lysate was clarified at 15,000 rpm in a Beck-
man JA-20 centrifuge rotor. Clarified lysate was applied to 10 mL
HisPur Co-NTA resin and incubated with gentle shaking at 4°C for 20
minutes to maximize binding. For purification, all buffers were
sparged thoroughly with CO to ensure the Hb0.1 Fe(II)-CO state is
maintained. The binding supernatant was eluted and resin washed
with 50 mL Hb0.1 lysis buffer, 50 mL wash buffer 1 (20 mM Tris,
500 mM NaCl, PH 8.5), 50 mL wash buffer 2 (20 mM Tris PH 8.5),
and finally eluted with 3 x 25 mL elution buffer (20 mM Tris, 100
mM imidazole, PH 8.5). The eluted fraction was applied to 5 mL Q-
Sepharose resin equilibrated with 20 mM Tris PH 8.5. Bound Hb0.1
was washed with 50 mL 20 mM Tris PH 8.5 and eluted with 3 x 20
mL 30 mM NaHPO4 PH 6.9. Eluted fraction of Hb0.1-CO was cen-
trated to 500 µM concentration and oxidized with 5 molar equiva-
lents of potassium ferricyanide (K3Fe(CN)6) for 1 hour at 4°C. Excess
oxidant was removed by gel filtration over Sephadex G-25, and eluted
metHb0.1 was buffer exchanged into heme transfer-sucrose buffer as
described for IsdH using 15 mL Amicon 3 kDa cutoff centrifugal fil-
ters.

Hemin transfer measurements. Rates of hemin transfer were
measured with an Applied Photophysics SX-20 stopped-flow spectro-
photometer under the following conditions: 5 µM metHb0.1 (heme
basis) and 150 μM heme acceptor (apo-IsdH or apo-Mb(H64Y/V68F)) in 20 mM NaPO4, 150 mM NaCl, 0.45 M sucrose, pH 7.5. Temperatures were varied from 15-37°C. Each mutant tested was measured in technical triplicate with 1000-second acquisition time at 405 nm (IsdH trials) and 600 nm (apo-Mb(H64Y/V68F) trials). Rate constants $k_{fast}$ and $k_{slow}$ were obtained through fitting the $A_{405}$ curve. The absorbance of the reaction mixture at 405 nm at any given time is the sum of absorbance contributions from holo-Hb0.1, holo-IsdH, and background absorbance (given by constant $A_{405}^{back}$). $A_{405} = A_{405}^{Hb0.1} \cdot \frac{[\text{holoHb0.1}]}{[\text{heme}]} + A_{405}^{IsdH} \cdot \frac{[\text{holoisdH}]}{[\text{heme}]} + A_{405}^{back}$. The transfer of heme from Hb0.1 to IsdH results in an overall reduction in absorbance because $e_{405}^{isdH}$ is lower than $e_{405}^{Hb0.1}$. This process is fitted to a two-exponential decay function a fast phase and a slow phase, governed by rate constants $k_{fast}$ and $k_{slow}$. We constrain each process to 50% of the span, where span = $A_{405}^{t=0}$ – plateau. $A_{405}(t) = plateau + span \cdot (0.5e^{-k_{fast}t} + 0.5e^{-k_{slow}t})$

With limited experiment time, the plateau value for a given reaction is approximated as the minimum absorbance value obtained after reaction for 1000 seconds, the typical length of data acquisition. For fast mutants, like WT IsdH, this timeframe is sufficient to observe complete reaction with metHb0.1. Slower mutants do not react completely within 1000 seconds, so the span value is only a fraction of that observed with WT. In our analysis, the WT span is set as the maximum, describing complete transfer of 5 μM heme from Hb0.1 to IsdH. For each slow mutant, we convert fractional amplitudes observed with WT to IsdH. For each slow mutant, we convert fractional amplitudes obtained after 1000 seconds to the concentration of heme transferred as calibrated by the WT span. This allows us to report the rates in terms of the concentration of heme transferred per second, enabling more meaningful comparisons of the transfer kinetics of the receptor mutants.

RESULTS

Inter-domain motions enable the LN3 unit to transiently engage Hb’s heme pocket. NMR studies of IsdH$^{N2N3}$ in its Hb-free state (hereafter called apo-IsdH$^{N2N3}$) revealed the presence of significant inter-domain motions between its N2 and linker domains, and suggested that these motions persist even after the receptor binds to Hb [10, 29]. To probe these domain rearrangements, we performed molecular dynamics (MD) simulations of apo-IsdH$^{N2N3}$ and the IsdH$^{N2N3}$:Hb complex. Three 500 ns simulations of apo-IsdH$^{N2N3}$ were performed, revealing large scale inter-domain motions between the N2 domain and the LN3 unit that are enabled by coordinate rearrangements in the ten amino acid tether sequence (residues P466-Y475) (Fig. S1). Two metrics were used to quantify the inter-domain motions: the inter-domain separation between the centers of mass (COM) of the N2 and N3 domains, and the degree of twisting defined by the dihedral angle between the N2 COM, the intervening linear linker domain, and the N3 COM (Fig. 2A). Each 500 ns simulation showed distinct inter-domain trajectories despite starting from identical initial conformations in which the domains are separated by ~49 Å (Fig. 2B). To sample longer time scale receptor motions, 20 rounds of ten x 100 ns MD simulations were performed using the FAST protocol (see methods), and trajectories were analyzed by constructing a Markov model [37]. The model provides a population estimate for 200 microstates, each corresponding to a distinct inter-domain arrangement between the N2 and LN3 units. These parameters were then used to generate a free energy landscape of inter-domain motions. In the apo-receptor the landscape is relatively flat, with most conformers accessible at room temperature; their free energies are only 2-4 kcal/mol above the ground state (Fig. 2C). However, a closed state in which N2 and N3 domains are proximal to one another is slightly preferred. This agrees with our original three 500 ns trajectories, which revealed that the compact state can be stable for hundreds of nanoseconds. In the apo-state the receptor also transiently samples the conformation of the Hb-bound form visualized by crystallography (star), although it represents only a small fraction of the conformers sampled in
equilibrium. The MD-derived energy landscape is also compatible with an NMR model of the apo-receptor, which samples a narrower distribution of receptor conformers (indicated by black dots on the landscape) [10].

To explore the effects Hb binding on receptor-inter-domain motions we performed MD simulations of the IsdH^{N2N3}:Hb complex and constructed a Markov model. Initially, we obtained diverse apo-state configurations by coarse graining the apo-state Markov model into 75 macrostates using BACE [40], and then we selected a representative structure at random from each macrostate. A Hb dimer was then introduced into the system by aligning the N2 domain in each representative structure to the N2 domain located in the crystal structure of the IsdH^{N2N3}:Hb complex in which Hb is dimeric (PDB: 4XS0). This procedure was used because the affinity of N2 for Hb’s A-helix is much stronger than the LN3 unit that contacts the heme pocket [11]. After alignment, some of the structures exhibited atomic overlap between the coordinates of the Hb dimer and the LN3 unit and were eliminated. This left approximately 50 starting structures of the IsdH^{N2N3}:Hb complex that were each simulated for 100 ns, followed by further sampling using the FAST protocol and steered-MD, leading to 4.5 µs of total sampling (see methods).

An analysis of the Markov model of the MD trajectories reveals that the tether undergoes wide ranging coordinate rearrangements that enable the LN3 unit to move relative to the N2 domain and Hb. A comparison of the free energy landscapes of these inter-domain motions in the free- and Hb bound forms of receptor reveals substantial differences (Fig. 2C). Not only has the most stable conformation of the receptor changed, but the energy landscape is less flat and more well shaped when the receptor binds to Hb. Thus, IsdH remains flexible when in complex with Hb as predicted from NMR measurements, albeit less flexible than when it is in its apo-state. Interestingly, in the MD simulations the LN3 unit receptor only transiently samples conformations that resemble the crystal structure of the extraction complex (red star), which are approximately 4 kcal/mol higher in energy than the most stable conformer. This is in marked contrast to the crystal structure complex in which the LN3 is engaged with Hb (Fig. 1A). A comparison of the conformer ensembles of apo-IsdH^{N2N3} and IsdH^{N2N3}:Hb suggests that Hb binding occurs via a conformational selection mechanism as only a subset of apo-IsdH conformers are capable of engaging Hb via the their N2 domains without causing unfavorable atomic overlap with the remainder of the receptor. Collectively, the MD simulations reveal that the tether sequence connecting the N2 and LN3 units plays an instrumental role in orchestrating inter-domain motions in the receptor that persist even after it has bound to Hb.

Altering the receptor’s inter-domain tether slows the rate of hemin extraction from Hb. To gain insight into the functional consequences of inter-domain motions, we measured the hemin extraction kinetics of a series of receptor variants in which the tether sequence connecting the N2 and LN3 unit was altered (Fig. 1B). As described previously, kinetic measurements employed a tetramer stabilized form of Hb (Hb0.1) and an IsdH^{N2N3} variant that preferentially binds to the α-globin within Hb0.1 (*IsdH^{N2N3}, IsdH^{N2N3} containing F365Y and A369F substitutions within its N2 domain) [12]. Using these protein reagents enables measurement of the rate of hemin extraction from only the α-subunit of tetrameric Hb, avoiding complications associated with Hb dissociation into its dimeric and monomeric forms. Hemin extraction rates were determined by tracking the change in Soret absorbance at 405 nm using a stopped-flow UV/Visible spectrophotometer (Fig. 3A). Hemin transfer from Hb to *IsdH^{N2N3} is biphasic, with rapid and slow changes in A_{405} characterized by k_{fast} and k_{slow} rate constants, respectively (Fig. 3B). For fully functional receptors k_{fast} defines the rate constant describing active hemin extraction from Hb’s α-globin, whereas the value of k_{slow} characterizes transfer to the receptor that occurs via an indirect process in which Hb first releases hemin into the solvent followed by hemin binding to the apo-receptor [11]. To assess the effects of tether mutations on hemin transfer we compared the initial rate of hemin transfer (µM s⁻¹), since for several of the receptor variants the transfer reaction was not completed during the time frame of the stopped-flow experiment. The kinetics data is reported in Table S1 and was obtained as described in the Methods section.

Rapid mixing experiments using native *IsdH^{N2N3} reveal that it actively extracts hemin from Hb at a rate of 0.93 ± 0.05 µM s⁻¹. This is significantly faster than the rate at which Hb0.1 spontaneously releases hemin into the solvent, consistent with an active extraction process. Housing the N2 and LN3 units within the same polypeptide is critical, since a polypeptide containing only the isolated LN3 unit (IsdH^{LN3}) captures hemin from Hb0.1 435-fold slower than the native *IsdH^{N2N3} receptor (0.002 ± 0.004 µM s⁻¹). Adding a polypeptide encoding the N2 domain to this reaction (*IsdH^{N2}) only marginally affects the rate of hemin capture by IsdH^{N3}, indicating that the main function of N2 is to tether the LN3 unit to Hb0.1 and its N2 binding to the A-helix does not promote hemin release. Three receptor variants in which the tether was altered were studied: (i) *IsdH^{INS}, an insertion variant containing an additional ten residue (GS)₄ segment in the tether that is located after residue P466, (ii) *IsdH^{DEL}, a tether deletion variant that removes residues P466-Y475, and (iii) *IsdH^{REP}, a variant in which amino acid sequence P466-Y475 in the tether is replaced with (GS)₄ (Fig. 1B). Stopped-flow experiments reveal that replacing the tether’s amino acid sequence with (GS)₄ or lengthening it slows hemin capture dramatically, as the *IsdH^{REP} and *IsdH^{INS} variants are slowed 320- and 73-fold relative to *IsdH^{N2N3}, respectively. *IsdH^{DEL} in which the tether sequence is removed entirely is extremely slow at capturing hemin, with transfer rates 4140-fold slower than the native receptor. In fact, the rate of hemin transfer by *IsdH^{DEL} is substantially slower than the intrinsic rate of spontaneous hemin loss from Hb0.1. This result suggests that *IsdH^{DEL} acts to inhibit hemin loss from Hb0.1, perhaps by blocking hemin egress from Hb through steric occlusion by the misbound LN3 unit. This massive loss in activity is not caused by disruption of the tertiary structure of the receptor, as ¹⁵N-labeled *IsdH^{DEL} and *IsdH^{N2N3} exhibit similar NMR spectra. Collectively, our results show that the tether sequence serves as the nexus for inter-domain motions in the receptor and plays a critical role in coupling the destabilization of Hb’s hemin pockets with the direct extraction and capture of hemin by the N3 domain.
C. Traces are using the following receptor variants: nevertheless, tether is removed reveal that the LN3 unit remains dynamic. Nevertheless, tether sequence is lengthened, as evidenced by an expanded density which the tether sequence is lengthened, as evidenced by an expanded density. Simulations of the Hb:InsN3 complex indicated by a dashed line). Conformers in which the location of N3 in the crystal structure of the receptor more stably stays affixed to Hb’s pocket, sampling a range of conformations near Hb that resemble its positioning in the crystal structure. Fig. 4A, high atomic density colored red and the location of N3 in the crystal structure of the receptor complex indicated by a dashed line). Conformers in which the N3 domain is near Hb’s hemin, but translated as compared to the native receptor (Fig. 4C). Transfer measurements indicate that InsHDEL captures hemin very slowly, slower than the rate at which Hb spontaneously releases hemin into the solvent. This is compatible with the simulation data as the close proximity of the N3 domain relative to Hb may act to block hemin release, and because the domain is displaced, the LN3 unit may be unable to distort Hb’s F-helix to trigger hemin release.

A Markov model of the simulation data was constructed to further probe how the tether sequence impacts the ability of the LN3 extraction unit to engage Hb’s heme pocket. The wild-type and mutant receptor-Hb trajectories were coarse grained, grouped into ~150 microstates and a Markov model constructed that contained 25 macrostates. These macrostates were then placed into one of two categories: an associated group where the average minimum distance between Hb and the N3 domain was below 4 Å, and a dissociated group that was separated by greater than 4 Å. This definition allowed us to estimate the timescale of the N3-Hb interaction in a general manner, as the association group involves both direct transfer competent and incompetent Hb-receptor orientations. The average association transition times required for the N3 domain in each type of receptor to approach Hb’s heme pocket are shown in Fig 5A, providing insight into the frequency of engagement. Consistent with density plot analysis that shows that the N3 domains in the wild-type and tether deletion mutant tend to hover near Hb, in the InsHDEL and InsHINS proteins N3 associations occur within 7 μs.

The tether promotes directed inter-domain motions that orient the LN3 unit for hemin extraction. To gain insight into why altering the tether affects the rate of hemin extraction, we simulated the dynamics of the InsHINS and InsHDEL receptors when bound to Hb and produced a holostate Markov model as described above for native InsHINS. Altering the tether results in profound changes in the energetic landscapes describing inter-domain motions between the N2 and LN3 units (compare Figs. S2 and S3). These differences are highlighted in Fig. 4 which shows a probability-weighted atomic density map defining the positioning of the N3 domain during the course of each simulation. In the native InsHINS receptor the N3 domain in the LN3 unit primarily stays affixed to Hb’s pocket, sampling a range of conformations near Hb that resemble its positioning in the crystal structure (Fig. 4A, high atomic density colored red and the location of N3 in the crystal structure of the receptor complex indicated by a dashed line). Conformers in which the N3 domain is positioned distal to Hb are also sampled, but occur less frequently. In contrast, a much larger distribution of N3 positions is observed in simulations of InsHINS in which the tether sequence is lengthened, as evidenced by an expanded density cloud (Fig. 4B). Moreover, the most prevalent position sampled by N3 in the Hb:InsHINS complex no longer resembles the crystal structure, suggesting the tether sequence is required to properly position N3 near Hb, and explaining why its alteration slows the rate of hemin removal. Simulations of the Hb:InsHDEL complex in which the tether is removed reveal that the LN3 unit remains dynamic. Nevertheless, its positioning is biased toward a closed state in which the N3 domain is near Hb’s hemin, but translated as compared to the native receptor (Fig. 4C). Transfer measurements indicate that InsHDEL captures hemin very slowly, slower than the rate at which Hb spontaneously releases hemin into the solvent. This is compatible with the simulation data as the close proximity of the N3 domain relative to Hb may act to block hemin release, and because the domain is displaced, the LN3 unit may be unable to distort Hb’s F-helix to trigger hemin release.

Figure 3. Stopped-flow measurements of hemin transfer. (A) Representative stopped-flow UV/Vis absorbance changes during the course of reaction. In the experiment apo-InsH2N3 (150 μM) is rapidly mixed with holo-Hb0.1 (20 μM) in its ferric state at 25°C. Traces are colored on a scale from t = 0 s (red) to t = 1000 s (blue). (B) Graph showing the change in absorbance at 405 nm after mixing holo-Hb0.1 with different tether variants of apo-InsH2N3. Transfer measurements were performed at 25°C using the following receptor variants: apo-InsH2N3, InsHDEL, InsHINS, InsHREP and N2 + LN3 added in trans. The data shows that altering the tether significantly slows transfer, which remains incomplete even after 1000 s. (C) Eyring plots for InsH tether variants that were used to determine the thermodynamic activation parameters (ΔH‡ and ΔS‡) of hemin transfer. The variants show distinct thermodynamic signatures indicating that they scavenge hemin via both direct and indirect mechanisms.
Figure 4. Atomic density plots describing the positioning of the N3 domain in wild-type and mutant receptors. The positioning of the N3 domain in simulations of complexes containing Hb bound to αIsdH<sup>WT</sup> (left), αIsdH<sup>INS</sup> (middle) and αIsdH<sup>DEL</sup> (right) are shown. A space filling model of dimeric α/β Hb is shown (gray) with its hemin molecules drawn in stick format. In each image the structure of the most populated form of the receptor is shown as a cartoon. The positioning of the N3 domain sampled in each simulation trajectory is represented as an atomic density map (blue = low density, red = high density). The positioning of the N3 domain observed in the crystal structure of the complex (PDB: 4XS0) is indicated by dashed outline (black) and presumably represents the active form of the receptor. The N3 domain in simulations containing αIsdH<sup>WT</sup> have the greatest degree of density overlap with the crystal structure, whereas in αIsdH<sup>INS</sup> the N3 density is more disperse. In αIsdH<sup>DEL</sup> simulations the N3 domain is translated relative to its positioning observed in the crystal structure of the receptor–Hb complex.

In the case of αIsdH<sup>DEL</sup> these associated states presumably represent inactive configurations in which the N3 domain is translated relative the Hb’s hemin (Fig. 5B). Moreover, compatible with density plots that show that the N3 domain in αIsdH<sup>INS</sup> samples a wider range of configurations that are removed from Hb, lengthening the tether slows the rate of N3 association with Hb, which requires on average more than 15 μs. Less substantial differences between the receptor types are observed when the average N3 dissociation times are compared, however dissociation events for both mutations occur on a longer timescale.
If the tether acts to bias inter-domain motions toward Hb, does it also impart a directional prejudice that increases the likelihood that the LN3 unit encounters Hb in an extraction competent orientation? Previous studies identified two sub-sites in the LN3 unit that promote hemin release from Hb by contacting its F-helix, the “L” and “N3” sub-sites [29] (Fig. 6A). We reasoned that LN3 conformers in the MD trajectory that were oriented similarly to the crystal structure of the IsdH<sup>N2N3</sup>-Hb complex would be poised to extract hemin and perhaps enriched in simulations of the wild-type receptor. To investigate this issue, we specified the orientation of receptors in the MD trajectories using three vectors whose relative positioning provides insight into the separation and orientation of the LN3 unit relative to Hb (Fig. 6B). These included a vector connecting points within N2 domain (N2 vector), a vector through the linker domain connecting N2 to N3 (L vector) and a vector connecting points within N3 (the N3 vector). N2 and Hb are tightly bound and move with respect to one another during the simulation. Therefore, the angle between the L and N2 vectors (N2-L angle) is sensitive to LN3 motions that alter its displacement from Hb, while the dihedral angle generated by rotating about the L vector (L dihedral angle) reports on the orientation of the sub-sites within LN3 unit. Fig. 6B shows a plot of these angular parameters generated using the 25 macrostates that define the ensemble of receptor conformations present in the MD simulations. The crystal structure presumably represents a hemin extraction-competent configuration and has an N2-L angle and L dihedral angle of 79.2° and -60.2°, respectively (colored red in Fig. 6B). Each type of receptor samples a range of conformations that differ from the crystal structure based on their angle and dihedral metrics. However, inspection of the "IsdH<sup>WT</sup>" macrostates reveals that they are enriched for conformers that resemble the crystal structure (left). The active configuration orientation for the native receptor is also evident when the vectors for each protein are visually displayed after aligning their N2 vectors (Fig. 6B, top). Consistent with the atomic density plots, "IsdH<sup>WT</sup>" samples a range of conformations, but has a preference for closed configurations that resemble the crystal structure in both its distance separation and orientation of the sub-sites relative to Hb. In contrast, in the "IsdH<sup>NS</sup>" simulations the extraction unit samples a much wider range of orientations that
differ in their separation (N2-L angle) and orientation (L-dihedral angle) relative to Hb. Moreover, even though "IsdH^DEL" tends to position its LN3 unit near Hb, a large scatter in the dihedral angle indicates that its LN3 unit is misaligned with Hb. The conclusion that the tether plays a critical role in enabling the extraction unit to transiently engage Hb’s F-helix in functionally active orientations is further supported by measurements of the N3 pocket angle, a metric that measures the alignment of the heme binding pockets in Hb and the receptor (Fig. 5C). This analysis reveals that in simulations of the wild-type receptor ~20% of the conformers in the trajectory have pocket angles that are within 10° of the crystal structure, whereas the insertion and deletion tether mutations rarely sample this extraction competent configuration. Thus, we conclude that the tether sequence acts to directly motions in the LN3 unit such that the unit samples conformers that are properly oriented and positioned for hemin extraction.

**Perturbing the receptor’s inter-domain dynamics causes it to passively acquire hemin from the solvent.** If altering inter-domain dynamics slows hemin capture, does it also affect the mechanism through which heme is transferred from Hb to the receptor? To investigate this issue the temperature dependence of the rate of heme capture for wild-type and mutant forms of the receptor were measured and interpreted using an Eyring analysis (Table S2). A similar analysis was performed for the hemin transfer reaction from Hb to apo-Mb, which is known to occur through an indirect process in which Hb first releases heme into the solvent before it is bound by apo-Mb. Based on their measured extraction velocities, the tether variants can be ranked from fastest to slowest as: "IsdH^N2N3" > "IsdH^NS" > "IsdH^REP" > "IsdH^DEL" (Fig. 3B and Table S1). Interestingly, as the receptor heme capture efficiencies decrease ΔH° becomes progressively less favorable, trending from a favorable 14.9 cal/mol*K for "IsdH^N2N3" to an unfavorable -59.5 cal/mol*K for "IsdH^DEL" (Fig. 7). An opposite trend in ΔS° occurs, which becomes more favorable as the receptor become less able to rapidly capture heme; 22.6 ± 0.7 kcal/mol ("IsdH^N2N3") to 4.3 ± 1.7 kcal/mol ("IsdH^REP"). The energetic profile of "IsdH^DEL" is like that observed for apo-Mb, suggesting that it is unable to actively distort Hb’s heme pocket, instead scavenging heme that is released from Hb into the solvent.

**DISCUSSION**

Inter-domain motions play a key role in modulating protein function by altering their affinity and specificity for ligands, as well as governing the efficiencies with which they catalyze chemical transformations [47, 48]. Domains are frequently connected by short polypeptide tethers, whose length and amino acid composition constrains their range of motion [49, 50]. Here we show using MD simulations and quantitative measurements of heme transfer that a flexible inter-domain tether within the IsdH receptor plays a critical role in controlling domain motions that are needed to extract heme from Hb. Simulations of the native complex reveal that it is dynamic, with coordinate rearrangements in the tether enabling the LN3 unit to adopt a range of conformations relative to Hb, while the N2 domain remains affixed to the globin A-helix. Interestingly, these inter-domain motions are biased, such that within the cloud of receptor conformers there is an enrichment for LN3 units that are positioned near Hb’s heme pocket (Figs. 2, 4) that are also
properly oriented to distort its F-helix (Figs. 5, 6). The specific amino acid sequence of the tether is important for function, as a "IsdHREP" receptor variant in which the tether is replaced with glycine and serine residues acquires heme 320-fold slower than the wild-type receptor. The finding that tether residues enable the low-affinity LN3 unit to move even after the receptor binds to Hb is consistent with NMR studies [10]. It is also compatible with the crystal structure of IsdH[P2N3]-Hb, as electron density for 7 of the 10 residues in the tether (P466-E472) were missing suggesting that they may be flexible [26, 12]. However, in the crystal structure of the complex, the LN3 adopts a single conformation in which it is affixed to Hb, presumably because crystal packing interactions restrict its mobility.

Our results suggest that within the dynamic complex the LN3 unit repeatedly engages Hb, but it only infrequently forms extraction competent configurations in which heme is removed. Comparison of the extraction kinetics of the native receptor and an isolated LN3 unit reveals that the motional bias imparted by the native tether increases the effective concentration of the LN3 unit near Hb’s heme to ~15 mM. This presumably drives the formation of the weaker LN3-Hb interface in which Hb’s heme pocket is distorted. A Markov model of the simulation data reveals that the LN3 unit samples a range of configurations that generally resemble the crystal structure. Using the LN3 position in the crystal structure in which sub-sites are aligned with Hb to define "transfer competence", the model predicts that only 5% of complexes in equilibrium are competent for direct transfer, which is in agreement with previous NMR results [29]. Based on the mean first passage times of LN3, Hb association and dissociation events occur within ~10 μs (Fig. 5).

Using the rate constant of direct heme transfer (ksuming) for "IsdHP2N3", the mean lifetime (τ) of the functional "IsdHP2N3"-Hb complex under pseudo-first order conditions is ~1 s. In comparing these values, we postulate that LN3 can potentially undergo ~107 association and dissociation events prior to successfully extracting heme. This implies that only a small subset of LN3-Hb association events within the dynamic complex satisfy conditions needed to successfully transfer heme from Hb to the receptor.

The tether connecting the N2 and LN3 units dictates inter-domain motions that are important for heme extraction. This is evident from studies of "IsdHP2N3", a receptor variant that contains a (GS)5 insertion after residue P466 in the tether. Relative to the wild-type protein it exhibits a large 73-fold reduction in the transfer rate, which coincides with the results of MD simulations that reveal that the dynamic LN3 unit samples a much larger, more isomerogenic conformational landscape in which the orientational constraints imposed by the native tether sequence are lost (Figs. 4-6). The increased motional freedom caused by the insertion also extends the mean first passage time needed to associate with Hb’s heme pocket from 7 to >14 μs, resulting in greater exploration of LN3 dissociated states that are unable to extract heme (Fig. 5). In "IsdHDEL" the tether is deleted entirely. This rather aggressive mutation still yields a well-folded protein as determined by 15N-HSQC experiments, but causes a very substantial ~4100-fold reduction in the rate of heme capture. In the "IsdHDEL" the receptor’s ability to orient its LN3 unit for productive Hb engagement is lost, as both the positioning of the LN3 unit (Fig. 3) and orientation relative to Hb (Fig. 5) are altered as compared to the native receptor.

An analysis of the activation energies of heme transfer reveals that altering the tether causes the receptor to capture heme from Hb via an indirect process in which heme is first released into the solvent before being captured by the receptor (Fig. 7A). For a series of receptor variants, a plot of their activation energies reveals that slower transfer rates relate to ΔS‡ becoming more negative and ΔH‡ becoming smaller in magnitude. Both parameters are correlated with the degree of receptor impairment and the correlation holds true for mutant receptors with either altered tethers ("IsdHREP", "IsdHDEL", and "IsdHP2N3") or amino acid substitutions in receptor sub-sites that directly contact Hb’s F-helix in the crystal structure ("IsdH[N3-]Helix", "IsdH[N3-]Lys6") (Fig. 7B).

Transfer of heme from Hb to apo-Mb occurs through indirect mechanism and exhibits similar activation energies as the most defective IsdH receptors. Therefore, we postulate that that in both the tether and sub-site receptor mutants, an increasing proportion of heme is transferred via an indirect mechanism wherein heme is first released from Hb into the solvent. Presumably in the indirect process heme solvation causes the large unfavorable change in ΔS‡, whereas the direct heme transfer reaction is enthaplycally limited because the receptor must distort Hb’s F-helix. Thus, combined with the MD results, the transfer data suggest that directed inter-domain motions orchestrated by the tether sequence are needed to populate conformers that provide a route for direct heme transfer between Hb and N3.

A comparison of the tether conformations sampled during the simulations provides insight into how it directs restricted, and functionally important LN3 motions. The primary sequence of the tether (66PDDYDEETY475) is notably populated with carboxylate sidechains that are documented to induce disorder [51]. In simulations of the native IsdH-Hb complex these sidechains frequently participate in intratether hydrogen bonding interactions with the polypeptide backbone that bias the tether’s conformation toward a compact state. Interestingly, in many of these condensed configurations the side chain of Y475 in the tether forms sustained interactions with a hydrophobic patch on the N2 domain thereby constraining the LN3 unit so that it is positioned near Hb’s heme pocket. This closed state is captured in the crystal structure of the complex where the Y475 aromatic ring is docked against a relatively hydrophobic surface on the N2 domain formed by P351 and P374, and poised to donate a hydrogen bond to E375 (Fig. 8). This constrained configuration is only transiently sampled during the simulations explaining why the LN3 unit exhibits directed motions (Fig. 4) and why the tether mutants remove heme slowly as they disrupt this interaction. The tether sequence in IsdB also contains conserved tyrosine residue (Y273 in IsdB numbering) and is enriched with anionic residues (Fig. 1A), suggesting that similar tether-N2 contacts will bias its LN3 unit toward conformations that are poised to extract heme.

Domains connected by flexible tethers are often thought to exhibit disordered motions in which they freely re-orient with respect to one another. In this study we have shown that despite exhibiting a high degree of conformational flexibility, a short inter-domain tether in IsdH nevertheless restricts and directs motions of its heme extraction unit when the receptor is bound to Hb. These motions enable the LN3 extraction unit to populate a low affinity interface in which Hb’s heme pocket is transiently distorted. Altering the tether misa-
ligns these motions, reducing the rate of direct heme scavenging and causing the receptor to slowly acquire hemin through an indirect process. The functionally important motions in IsdH are akin to those observed in protein chaperones, polyketide synthases, and prolyl isomerases [49, 50, 52, 53], which also employ inter-domain tethers that determine the frequency and geometry of molecular interactions. In IsdH these motions are likely biologically important, as they enable S. aureus to overcome Hb’s strong affinity for its hemin molecules without the need to form a high affinity, long-lived receptor-Hb complex. In the well-studied HasA-HasR transfer complex from Serratia marcescens, hemin is transferred by forming a high affinity complex in which the protein-protein binding free energy is used to weaken HasA’s contacts with hemin [54–56]. In contrast, both the IsdB and IsdH receptors bind to Hb with modest affinity and harness inter-domain dynamics to facilitate heme removal. Extraction within this weak and dynamic complex presumably maximizes the rate of heme flow into the cell, enabling repeated rounds of Hb capture and heme removal to occur on the microbial surface.

**Figure 8.** Model of on-target LN3 engagement A) Two macrostates derived from simulations of “IsdH{N2,N3}.Hb demonstrate the LN3 unit in disengaged (light orange) and engaged (orange) states, which reveal that Y475 (blue spheres) defines the primary contact between the tether and a conserved solvent-exposed patch on N2 (green). This interaction enforces an inter-domain geometry that positions the LN3 subsites (purple) in proximity with the heme-binding site of Hb (gray). B) A molecular model of the docked Tyr475 (blue sticks) with the cognate conserved patch residues on N2.

**REFERENCES**


