Structure–function analyses reveal key features in *Staphylococcus aureus* IsdB-associated unfolding of the heme-binding pocket of human hemoglobin

Catherine F. M. Bowden, Anson C. K. Chan, Emily J. W. Li, Angelé L. Arrieta, Lindsay D. Eltis, and Michael E. P. Murphy

From the Department of Microbiology and Immunology, Life Sciences Institute, University of British Columbia, Vancouver, British Columbia V6T 1Z3, Canada

Received for publication, July 12, 2017, and in revised form, October 19, 2017

Edited by Norma Alliewell

IsdB is a receptor on the surface of the bacterial pathogen *Staphylococcus aureus* that extracts heme from hemoglobin (Hb) to enable growth on Hb as a sole iron source. IsdB is critically important both for *in vitro* growth on Hb and in infection models and is also highly up-regulated in blood, serum, and tissue infection models, indicating a key role of this receptor in bacterial virulence. However, structural information for IsdB is limited. We present here a crystal structure of a complex between human Hb and IsdB. In this complex, the α subunits of Hb are refolded with the heme displaced to the interface with IsdB. We also observe that atypical residues of Hb, His58 and His89 of αHb, coordinate to the heme iron, which is poised for transfer into the heme-binding pocket of IsdB. Moreover, the porphyrin ring interacts with IsdB residues Tyr440 and Tyr444. Previously, Tyr440 was observed to coordinate heme iron in an IsdB-heme complex structure. A Y440F/Y444F IsdB variant we produced was defective in heme transfer yet formed a stable complex with Hb ($K_d = 6 \pm 2 \mu M$) in solution with spectroscopic features of the bis-His species observed in the crystal structure. Haptoglobin binds to a distinct site on Hb to inhibit heme transfer to IsdB and growth of *S. aureus*, and a ternary complex of IsdB-Hb-Hp was observed. We propose a model for IsdB heme transfer from Hb that involves unfolding of Hb and heme iron ligand exchange.

*Staphylococcus aureus* is most commonly found as a member of the normal human flora, colonizing primarily on the hands and in the nostrils (1). However, *S. aureus* is also one of the main agents of nosocomial infections (2) and can cause a range of diseases in humans, from mild skin infections such as boils and folliculitis, to severe, life-threatening bloodstream infections (3). Establishing infection requires effective iron scavenging systems because iron trafficking is tightly controlled within the human body, both to mitigate the toxicity of free iron and to limit microbial growth. The latter has been referred to as a form of innate immunity called “nutritional immunity” (4). *S. aureus* overcomes this nutritional immunity by employing two mechanisms to access iron sources: the secretion and uptake of iron-chelating siderophores to acquire iron from host proteins such as transferrin and the expression of surface receptors for heme and hemoglobin. However, when presented with both heme and iron-bound transferrin as the only potential iron sources, *S. aureus* can acquire both forms of iron but preferentially utilizes heme (5).

*S. aureus* acquires heme using the Isd (iron-regulated surface determinant) system. This system consists of nine components: four surface proteins covalently anchored to the peptidoglycan that reversibly bind heme (IsdA, IsdB, IsdC, and IsdH); an ABC transporter (IsdF) with an associated lipoprotein (IsdE); and two intracellular heme-degrading enzymes (IsdG and Isdl) (6–8). The function of a predicted membrane protein, IsdD, remains unknown. Lastly, sortase B (SrtB) is encoded in a gene cluster with *isdCDEF* and functions to anchor IsdC to the peptidoglycan, whereas the remaining Isd surface proteins (Isd-ABH) are anchored by sortase A, the housekeeping sortase of the cell (7). The expression of all Isd components appears to be classically regulated by the Fur regulator and is repressed in the presence of iron (8). IsdB and IsdH stand apart in the Isd system in that they are the only components capable of binding hemo-proteins. IsdH can bind hemoglobin (Hb),2 haptoglobin (Hp), and the Hp-Hb complex. Hp is a high-abundance serum protein that binds free Hb with high affinity but does not, in itself, constitute an iron source. IsdB can bind Hb and the Hp-Hb complex but not Hp alone (9, 10). Thus, heme is stripped from Hb at the bacterial cell surface by IsdB or IsdH and is transferred in a unidirectional relay to IsdE via IsdA and IsdC (11–13). The heme is then imported into the cell by the permease IsdF for degradation by the homologous enzymes IsdG and Isdl to liberate iron for use by the cell (14, 15).

1 To whom correspondence should be addressed: Dept. of Microbiology and Immunology, 2350 Health Sciences Mall, Life Sciences Institute, University of British Columbia, Vancouver, BC V6T 1Z3, Canada. Tel.: 604-822-8022; E-mail: Michael.Murphy@ubc.ca.

2 The abbreviations used are: Hb, hemoglobin; Hp, haptoglobin; metHb, methemoglobin; r.m.s.d., root mean square deviation; PDB, Protein Data Bank; ITC, isothermal titration calorimetry; NRPMI, metal-depleted RPMI; EDDHA, ethylenediamine-\(N,N'\)-bis(2-hydroxyphenylacetic acid).

This work was supported by Canadian Institutes of Health Research Grant MOP-49597 (to M. E. P. M.) and a Natural Sciences and Engineering Research Council of Canada Postgraduate Scholarship - Doctoral scholar-

1 The atomic coordinates and structure factors (code SVMM) have been deposited in the Protein Data Bank (http://wwwpdb.org/).
The four cell wall–anchored Isd proteins, IsdABCH, share three features: an N-terminal secretion signal, a C-terminal sortase signal for cell wall anchoring, and one to three copies of a NEAT (for near transporter) domain (16). IsdA and IsdC contain a single NEAT domain, which binds heme. IsdB and IsdH contain two and three NEAT domains, respectively, but only their C-terminal NEAT domains, IsdBN2 and IsdHN2, respectively, bind heme. The N-terminal and central NEAT domains of IsdH, recombinantly expressed individually as IsdHN1 and IsdHN2, are each able to bind Hb, Hp, and Hp-Hb (9, 17); however, the N-terminal IsdB domain alone, IsdBN1, does not bind Hb (18), despite >40% sequence identity between IsdBN1, IsdHN1, and IsdHN2 and the shared ability of the full-length proteins to bind Hb. Both IsdBN1 and IsdBN2 must be present and contiguous with the intervening “linker” region (represented as IsdBN1N2) for high affinity Hb binding. Moreover, IsdBN1N2 removes heme from oxidized Hb, known as methemoglobin (metHb), the form of Hb produced upon red blood cell lysis in the bloodstream. By contrast, IsdBN1N2 does not remove heme from oxyHb, the reduced, oxygen-bound form of Hb that is present in intact red blood cells, demonstrating specificity toward the probable biologically relevant form of Hb encountered by S. aureus during infection (18).

Structures of the Isd proteins have been elucidated by X-ray crystallography or NMR, revealing the mode of heme coordination within NEAT domains (17, 19–25). Recently, crystal structures of various portions of IsdH in complex with Hb have also been reported: IsdHN1-metHb (17), IsdHN2-metHb (19), and a heme-transfer–deficient variant of IsdHN2N3 (Y642A) complexed with metHb (19, 26). These complex structures provide insight into the extensive interactions between IsdH and Hb. In these structures, the heme remains largely encapsulated by metHb with minor or no distortions of the αHb heme binding pocket. Therefore, these structures only provide insight into the initial step of the heme transfer process. These insights likely extend to IsdB-Hb, given the similarities between the two surface receptors. Nevertheless, IsdB is more important than IsdH in both in vitro growth on Hb and a mouse abscess model of infection (10, 27). IsdB is also a dominant antigen that is highly up-regulated in blood, serum, and a cage model of tissue infection (28–31), corroborating the key role of this receptor.

Herein, we report the crystal structure of an IsdBN1N2-Hb complex in which the heme is positioned between the two proteins, consistent with an intermediate state of heme transfer. Two point mutations were then introduced into IsdBN1N2 to trap a similar species in solution, which was characterized spectroscopically. In concordance, kinetic analyses of heme transfer revealed a multistep transfer process. Furthermore, we demonstrate that although IsdBN1N2 can bind to the Hp-metHb, as was previously demonstrated for IsdH, it is unable to remove heme from the complex. This finding correlates with our observation that S. aureus did not utilize Hp-Hb as a sole iron source.

Results

Crystal structure of an IsdBN1N2-Hb complex

Because IsdBN1N2 does not extract heme from oxyHb (18), this form of Hb was used to obtain an IsdBN1N2-Hb complex for crystallization. Crystals of an IsdBN1N2-Hb complex were grown from solutions of citric acid or malonic acid, pH 5.0–5.5, and 2.1–2.4 M ammonium sulfate. Small crystals (~0.05–0.1 μm) appeared after 2–4 weeks and contained both proteins (Fig. 1, left panel). A 3.6 Å resolution data set from a single crystal was collected, and the structure was solved by molecular replacement of the individual components. The structure revealed a central Hb a2b2 tetramer surrounded by four IsdB molecules in the asymmetric unit (Fig. 1, right panel). An intact copy of the recombinant IsdBN1N2 construct is bound to each αHb subunit; however, only the first NEAT domain of IsdB (IsdBN1) is observed bound to each βHb subunit. The 19-kDa band observed in the left panel of Fig. 1 is consistent with proteolysis of IsdBN1N2 during crystallization and is presumed to be IsdBN1. Inspection of crystal packing within the unit cell revealed that modeling the missing components of IsdB would result in a large scale steric clash with αHb-bound IsdBN1N2, suggesting that the partial proteolysis allowed for crystallization of the complex.
The structure of each IsdBN^N1N2^ molecule resembles a dumbbell with the two NEAT domains joined by an α-helical linker (Fig. 2A, left panel), as was observed for IsdH^N2N3 (19, 26). The loop between IsdBN^N1^ and the linker region is flexible, as evidenced by poor electron density; conversely, well ordered density is observed between the linker and the second NEAT domain (IsdB^N2), which includes a short 3_10 helix. The IsdBN^N1^ and IsdBN^N2^ domains of intact IsdB^N1N2^ each make numerous interactions with αHb (average interface area of 780 and 736 Å², respectively), and similar interactions are observed for IsdB^N2^ bound to βHb (634 Å²). No interactions are observed between the linker region and Hb.

The Hb chains of the complex described here reveal major structural rearrangements as compared with the structure of isolated oxyHb (PDB code 2DN1). These changes are situated primarily in the two αHb chains of the IsdB^N1N2^-Hb structure, which superimpose poorly with the equivalent α chains in the free oxyHb structure (r.m.s.d. of 3.3 and 3.4 Å over all Ca) as compared with alignments between the βHb chains (r.m.s.d. of 1.4 and 1.3 Å over all Ca). In the α subunit of free human oxyHb, heme is bound in a pocket between the E and F helices, with His^87^ (F helix) coordinating directly to the heme iron and the distal His^58^ (E helix) forming a hydrogen bond to the heme-bound dioxygen molecule. To our knowledge, the heme iron axial residue is His^87^ in all previously reported Hb crystal structures regardless of the gaseous heme ligand. Upon binding to IsdB^N1N2^, a major deformation of the αHb heme pocket occurs, with the F helix and part of the E helix of each αHb chain (encompassing residues αAsp^74^ to αArg^82^) unwinding entirely (Fig. 2B). Clear electron density in this region indicated that heme (modeled at full occupancy) remains bound to βHb (Fig. 3A) but has been displaced 5 Å toward the heme-binding pocket of IsdB^N2^. This heme displacement was accompanied by the unexpected direct coordination of the heme iron by αHis^28^ and αHis^89^ (Fig. 3B). Through reorganization of the polypeptide chain, αHis^87^ has pivoted out of the heme pocket and...
points away from the heme iron (Ne2 translation of 13.4 Å). In its place, αHis99 (originally on the F helix) has moved from a solvent-exposed position to coordinate to the heme iron (13.2 Å translation in Ne2). Binding to IsdBN1N2 also resulted in the extension of the C-terminal H helix by another half turn, such that Tyr140 occupies the void left by the unwound F helix, with an accompanying Cα shift of 7.5 Å and the placement of the side chain phenol group within 3.2 Å from Nα of His87.

Eleven residues between helices C and D (αTyr42 to αSer52) have also rearranged to accommodate the interaction with IsdBN1N2 (Fig. 2A, right panel). For the β chains of Hb, insufficient density was present to model most of the F helix (βLeu188 to βVal198 in chain B and βPhe85 to βVal198 in chain D), and no bound heme was observed (Fig. 2C). Otherwise, minimal differences were observed between the β chains of free and IsdB-complexed Hb.

The IsdBN2 domain interacts with the heme-binding site of αHb and superposed well with the structure of free IsdBN2 (20) (r.m.s.d. of 1.4 Å over 109 Cα). However, the β7-β8 loop (Val185 to Tyr440), which coordinates heme iron via Tyr440, is curved inward in the complex structure, making contacts with αHb, and is poised to receive the heme molecule (Fig. 3C). Tyr440 closely abuts αHis89 of Hb, whereas Tyr444 is positioned adjacent to the heme pyrrole ring, forming a π-stacking network of interactions. Interestingly, Tyr140 is positioned further back than Tyr444, suggesting that Tyr444 plays a role beyond simply stabilizing the position of Tyr440 in the heme pocket, as was previously proposed (20), and appears to be required for the heme transfer process. The second heme iron–coordinating residue of holo IsdBN2, Met362, has poor electron density in the complex structure and is likely conformationally flexible. Conversely, the IsdBN2 propionate-binding residue Ser361 remains engaged in hydrogen-bonding with the propionate, with Glu354 also participating in this interaction. The second heme propionate is not observed to form H-bonds to either Hb or IsdBN1N2.

Proteolyzed IsdBN1 (chains J and H) interact with βHb and are structurally similar to the IsdBN1 domains in intact IsdBN1N2 (chains E and F), which interact with αHb in the crystal structure (r.m.s.d. of 0.9–1.2 Å over 133 Cα). Minor differences are observed mainly in the β2-β3 and β7-β8 loops (numbered

Figure 3. Binding of IsdBN1N2 to Hb induces major changes in heme environment. A, stereo view of an Fo – Fc omit positive difference map (green) of the heme in the IsdBN1N2–αHb interface contoured at 3σ (refined with torsion-based simulated annealing). An anomalous density map (pink) contoured at 3σ is overlaid unambiguously support correct placement of the heme. B, the complex structure is overlaid with the structure of the αHb chain from the oxyHb structure (PDB code 2DN1). The 2DN1 αHb structure is shown in green; αHb and IsdBN1N2 from the IsdBN1N2–Hb structure are shown in beige and blue, respectively. Relevant amino acid side chains are shown as sticks. Only the heme moiety from the structure presented here is shown (dark green). C, the IsdB heme pocket is positioned to accept αHb-heme. IsdB and αHb residues situated at the binding interface near the heme are shown as blue and beige sticks, respectively. The remainder of the structure is shown as cartoons. The αHb-heme is shown in green, with the heme iron (coordinated by αHis89 and αHis99) shown as an orange sphere.
S. aureus IsdB-associated unfolding of human hemoglobin

Figure 4. Electronic spectra of IsdBYYFYF combined with heme and Hb. A, addition of excess IsdBYYFYF to metHb results in a distinctly altered spectrum (blue); the heme spectrum alone (red), metHb spectrum alone (purple), the mixture of IsdBYYFYF with heme (green), and reaction of wild-type IsdBNN2 with metHb (orange) are shown for comparison. AU, absorbance units. B, a closer look at the visible region of the spectra presented in A.

An IsdB heme pocket variant traps a bis-His metHb

Although the IsdBNN2-Hb structure clearly showed the Hb-heme in a bis-His coordination state, a bis-His state involving the native proximal and distal heme ligands was previously observed in crystals of horse metHb grown at pH 5.4 as compared with pH 7.1 (33). Although the heme ligands differ, the low pH of our IsdBNN2-Hb crystallization solution could have similarly induced bis-His coordination. However, incubation of metHb or oxyHb at pH 5.5 overnight, with or without IsdBN1N2, did not produce spectral changes associated with formation of a bis-His state (data not shown). Nonetheless, the observation of conformational changes in horse metHb to give a bis-His coordination state is a precedent for the conformational rearrangements observed in the IsdBNN2-Hb structure.

A double mutant of IsdBNN2 (IsdBYYFYF) was created to prevent heme transfer by replacing Tyr440 (heme iron coordinating) and Tyr444 (H-bonds to Tyr440) with Phe residues in the heme pocket. In the complex structure, these two tyrosine residues are also juxtaposed to αHis89 of Hb, the heme-coordinating residue that is part of the unwind F helix. Upon addition of free heme to apo-IsdBYYFYF, the spectra displayed a Soret peak at 403 nm, slightly blue-shifted relative to holo-IsdBN1N2, which peaks at 405 nm (Fig. 4A). The α/β region of the IsdBYYFYF + heme spectrum was nearly featureless with a minor peak at ~600 nm, similar to the spectrum of free heme in that region (Fig. 4B).

As previously reported (18), the addition of IsdBNN2 to metHb resulted in a large-scale shift in the intensity and shape of the Soret peak as heme was transferred, with minor changes in peak intensities in the visible region (Fig. 4A). Visible region peak wavelengths remained at ~500 and 535 nm, with a charge-transfer band at 630 nm, indicating that the heme was a high-spin ferric species in both holo-IsdBNN2 and metHb (34). Addition of IsdBYYFYF to metHb also caused a large-scale shift in the Soret peak; however, the spectra was distinct from that formed upon heme addition to IsdBYYFYF (Fig. 4A). The Soret peak became red-shifted, to 410 nm, and a distinct shoulder developed at ~360 nm. Moreover, dramatic changes occurred in the α/β region (Fig. 4B); the charge-transfer band at 630 nm disappeared, and the highest absorption peak shifted to 533 nm with a shoulder peak at 564 nm. This spectrum is nearly identical to that of a ferric, low-spin, bis-His hemechromophore form of human Hb observed when incubating purified α- or β-globin with heme (but not for heme addition to whole globin) (34). The change to a characteristic bis-His hemechromophore spectrum upon addition of IsdBYYFYF to 4 µM metHb was titratable and saturable (Fig. 5, A and B). A plot of the absorption change at a single wavelength (410 nm) as a function of added IsdBYYFYF plateaued at ~5 µM, indicating the stoichiometry of the reaction was ~1:1 (Fig. 5C), as seen for interaction of IsdBNN2 and Hb in the crystal structure.

The binding of IsdBYYFYF to metHb was confirmed by ITC. Titration of metHb into IsdBYYFYF resulted in an exothermic reaction as observed by the negative change in enthalpy (Fig. S1). Analysis of the data with a one-site model gave a $K_d$ of 6 ± 2 µM and a stoichiometry (N) of ~0.5 (average of three runs), implying two IsdBYYFYF molecules bound to one metHb monomer. Because Hb has two unequal subunits that may interact with IsdBNN2 differently, the $K_d$ and stoichiometry measurements are assumed to be the average of binding to αHb and βHb. A previous ITC study between IsdBNN2 and carb oxyhemoglobin reported a $K_d$ of 0.42 ± 0.05 nM (18). The weaker interaction of IsdBYYFYF for metHb may be due to a conformational change in the structure of metHb analogous to that observed in the IsdBNN2-Hb crystal structure.

Rapid heme transfer from metHb to IsdBNN2

To identify potential intermediates in the heme transfer pathway, the kinetics of heme transfer from metHb to IsdBNN2 was investigated by stopped-flow spectroscopy. Mixing of 2 µM metHb with 20 µM IsdBNN2 resulted in spectral changes between 180 and 730 nm that were complete within 10 s, consistent with previous reports using a full-length IsdB construct (11). Coincident with heme transfer, the Soret peak underwent a large shift in intensity, particularly between ~350 and 420 nm, along with spectral changes in the α/β region (Fig. 6A).
Although the intensity of the Soret absorption peak increased during the first ~60 ms (Fig. 6B), the overall effect was a reduction in intensity with broadening of the Soret band.

Two wavelengths (406 and 428 nm) were chosen for single-wavelength stopped-flow spectroscopy under pseudo-first order conditions, with metHb held constant at 1 μM and IsdB\textsuperscript{N1N2} increasing from 5 to 40 μM. The kinetics at 428 nm were simpler (Fig. 6C), because this wavelength was outside the range that increased in the first 60 ms (~350 – 420 nm). A single exponential fit yielded observed rate constants (κ\textsubscript{obs}) that varied hyperbolically with IsdB\textsuperscript{N1N2} concentration (Fig. 6D), consistent with a two-step heme transfer mechanism (20) with a rate constant for heme transfer from metHb to IsdB\textsuperscript{N1N2} of 0.35 ± 0.02 s\textsuperscript{-1}. At 406 nm, the kinetics were more complex. Curve fitting yielded four phases with differing amplitudes, rates, and concentration dependences (Fig. S2). In the first phase only, the rate was linearly dependent on IsdB\textsuperscript{N1N2} concentration (Table 1; see also Fig. S2C) and thus reflected concentration dependent collision events. Phases 2–4 did not display strong concentration dependence in their rates (Fig. S2, D–F), suggesting that they are associated with steps in the heme transfer process after IsdB\textsuperscript{N1N2}-metHb complex formation.

**Hp prevents heme transfer from metHb to IsdB\textsuperscript{N1N2}**

Hp is a serum α2-sialoglycoprotein that binds metHb released from lysed erythrocytes to prevent oxidative damage (35, 36). The fundamental unit of human haptoglobin is a polymorphic αβ dimer composed of light chains (α) and heavy chains (β). The β chain is encoded by a single allele, whereas the α chains come in two forms: α\textsubscript{1} and α\textsubscript{2} (36). These alleles generate three possible phenotypes. Phenotype 1-1 (where both copies of the α gene are α\textsubscript{1}) is the simplest, with Hp forming a homodimer of two αβ heterodimers. Phenotypes 2-1 and 2-2 can form heterogeneous cyclical or linear multimers of increasing size (37). Within the human host, the normal plasma Hb concentration is less than 5 μN, whereas Hp is generally present in the range of 0.3–2 mg/ml (35) or roughly 6–50 μM depending on the phenotype.

Binding of metHb by Hp has been proposed to make heme inaccessible to microbial pathogens (38). Therefore, to investigate the effect of Hp on heme uptake by IsdB from Hb, excess IsdB\textsuperscript{N1N2} was added to metHb preincubated with increasing concentrations of human Hp. A pooled mixed phenotype Hp was used to model normal human serum; however, the exact molar concentration was unknown. Therefore, Hp was used at the high end of the normal range for Hp serum concentration (2 mg/ml). Preincubation of 2 μM metHb with 2 mg/ml Hp resulted in electronic spectra that remained unchanged for 5 min after the addition of IsdB\textsuperscript{N1N2} (Fig. 7A). Only by decreasing the Hp concentration to 0.02 mg/ml or lower was a spectral change in the Soret region observed, implying heme transfer similar to that observed in the absence of Hp (Fig. 7B). Thus, Hp effectively blocked heme transfer from metHb to IsdB\textsuperscript{N1N2}. Inhibition was not due to the inability of IsdB\textsuperscript{N1N2} to bind metHb when the latter is complexed to Hp because IsdB\textsuperscript{N1N2} was able to pull down metHb in both the presence and absence of Hp but did not interact with Hp alone (Fig. 7C). To elucidate the stoichiometry of the Hp inhibition, the heme transfer assay was repeated using Hp phenotype 1-1, which revealed a stoichiometric ratio of approximately one Hp 1-1 molecule (consisting of two αβ subunits) to one α\textsubscript{2}β\textsubscript{2} metHb tetramer (Fig. S3).

**S. aureus growth on Hb-Hp as sole iron source**

To examine the effects of Hp on Hb-heme utilization in vivo, the growth of *S. aureus* strain Newman was evaluated in iron-depleted RPMI media. As expected, *S. aureus* was able to grow in medium supplemented with 2 μM heme iron or 0.2 μM oxyHb as a sole iron source (Fig. 8; see also Fig. S4). A higher concentration of heme was
required because 0.2 μM heme did not support growth of *S. aureus* (data not shown). OxyHb is expected to rapidly oxidize to metHb in culture. Assuming one Hp αβ dimer binds to one dimer of Hb, growth on Hb with the addition of 0.44–10 μg/ml Hp reduced the growth of *S. aureus* in a concentration-dependent manner. Conversely, addition of Hp to medium supplemented with heme did not reduce the growth of *S. aureus*, implying that Hp had a Hb-specific inhibitory effect.

### Table 1

| IsdB<sup>N1N2</sup> (μM) | 5 μM | 10 μM | 17 μM | 25 μM | 40 μM |
|--------------------------|------|-------|-------|-------|-------|
| **Amplitude**            | 0.15 ± 0.02 | 0.13 ± 0.01 | 0.14 ± 0.01 | 0.12 ± 0.01 | 0.124 ± 0.003 |
| **Amplitude**            | 1.9 ± 0.2 | 2.1 ± 0.2 | 5 ± 1 | 4 ± 1 | 4 ± 1 |
| **Amplitude**            | 0.16 ± 0.02 | 0.19 ± 0.02 | 0.16 ± 0.02 | 0.17 ± 0.01 | 0.16 ± 0.02 |
| **Amplitude**            | 0.41 ± 0.03 | 0.42 ± 0.05 | 0.5 ± 0.1 | 0.5 ± 0.1 | 0.5 ± 0.1 |
| **Amplitude**            | 0.48 ± 0.06 | 0.48 ± 0.06 | 0.47 ± 0.07 | 0.45 ± 0.07 | 0.50 ± 0.06 |
| **Amplitude**            | 0.12 ± 0.02 | 0.13 ± 0.03 | 0.14 ± 0.04 | 0.15 ± 0.03 | 0.13 ± 0.04 |
| **Amplitude**            | 0.20 ± 0.03 | 0.21 ± 0.06 | 0.22 ± 0.08 | 0.26 ± 0.08 | 0.22 ± 0.08 |

---

**Figure 6.** Heme transfer kinetics from metHb to IsdB<sup>N1N2</sup>. A and B, electronic spectra collected with a stopped-flow spectrophotometer equipped with a photodiode array. A, spectra recorded over 15 s after mixing of 2 μM metHb with 20 μM IsdB<sup>N1N2</sup>. B, an increase in the Soret peak was observed over the first 60 ms, from 350–420 nm. **AU**, absorbance units.

C, a representative single-wavelength (428 nm) stopped-flow spectroscopy experiment where 1 μM metHb was mixed with 17 μM IsdB<sup>N1N2</sup>. The **gray bars** represent the standard error of four replicates, and the **black curve** is a fit to a single exponential equation. The residuals for the experiment are in the inset. D, the observed transfer rate (k<sub>obs</sub>) from 1 μM metHb is plotted as a function of IsdB<sup>N1N2</sup> concentration. The **line** is a hyperbolic fit assuming a two-step reaction model. Each point represents the mean, and the **bars** are the standard errors of four replicates. The residuals of the data to the model are in the inset.
**Discussion**

IsdB is the major Hb receptor functioning at the interface between the bacterial cell surface and the extracellular environment. The heme extraction function of IsdB is supported by the observed growth deficiency of *S. aureus* on nanomolar concentrations of Hb as a sole iron source upon deletion of IsdB (27) and more directly by heme transfer assays between metHb and IsdB (18). In contrast, IsdB^{N1N2} was unable to extract heme...
from oxyHb within hours of incubation, and this form of Hb was used in crystallization trials to obtain the structure of a complex between the two proteins.

Crystals of IsdB\(^{\text{NIN2}}\)-Hb formed slowly over 2 weeks, sufficient time for oxyHb to be oxidized to metHb. Indeed, the heme groups from the βHb subunits are no longer observed in the complex structure and presumably were extracted by IsdB\(^{\text{NIN2}}\) that was subsequently degraded, leaving only the IsdB-N1 domain. The conformations of the αHb subunits in the complex with IsdB\(^{\text{NIN2}}\) are a large departure from all previously published structures of Hb. The large distortion of the polypeptide chain and the bis-His heme coordination state are accompanied by a shift of the heme from Hb to the interface with IsdB. The structure suggests that heme is removed from βHb before αHb, which may be a consequence of the kinetics or thermodynamics of crystal formation. The observed bis-His form of Hb observed in the crystal structure may be an artifact of crystallization and represent an off-path species. Attempts to identify this putative intermediate in solution by stopped-flow kinetics have not yet met with success, possibly because the putative intermediate is short-lived. However, spectroscopic analysis of the IsdB\(^{\text{NIN2}}\) variant mixed with metHb supports the trapping of a bis-His heme complex in solution, thereby supporting the possible existence of a bis-His heme intermediate in the heme transfer pathway by the wild-type protein.

An attractive model of heme transfer from Hb to IsdB is suggested when the IsdB\(^{\text{NIN2}}\)-Hb complex structure is overlaid with the structures of oxyHb (PDB code 2DN1; α chain) and holo-IsdB\(^{\text{N2}}\) (PDB code 3RTL) (Fig. 9A). In the IsdB\(^{\text{NIN2}}\)-Hb structure presented here, the heme moiety is midway between the Hb and IsdB heme pockets (heme iron is ~5 Å from either position). The motion of the heme is not solely translational, as one of the propionate groups in the complex is in nearly the same position as observed in the holo-IsdB\(^{\text{N2}}\) structure (Fig. 9). Instead, H-bond interactions between the propionate and both Ser\(^{361}\) and Glu\(^{355}\) appear to anchor the heme as it rotates ~90° from the Hb heme pocket to the IsdB heme pocket, providing the most parsimonious route of heme transfer between the start and end states (as represented by the uncomplexed structures). Interestingly, the structures of oxyHb and our complex may also provide insight into the ambiguity observed in the electron density for the heme methyl/ethyl groups in the previously solved structure of holo-IsdB\(^{\text{N2}}\), which resulted in two possible heme conformations while maintaining the position of the propionate groups (20). Because it is unlikely that the heme face can flip over during transfer, one can argue that heme conformation A, as shown in Fig. 9B, is the biologically relevant heme binding mode when transferred from Hb.

The interaction between IsdB\(^{\text{N1}}\) and βHb may represent an interaction after a successful heme transfer event, because the βHb heme and the IsdB NEAT1 and linker domains are not present. Additionally, helix F of βHb, which contains βHis\(^{92}\) (equivalent to axial heme-coordinating His\(^{97}\) of αHb) and βHis\(^{97}\), are disordered in the absence of bound heme. βHis\(^{97}\) is a single turn helix away from the equivalent position of αHis\(^{99}\) and may participate in the formation of a similar bis-His conformation in βHb. For the transfer to occur, βHb heme must have oxidized because IsdB\(^{\text{NIN1}}\) cannot remove ferrous heme from Hb (18). This slow oxidation is expected because the crystals of the complex took weeks to form. Subsequent to the oxidation, the heme is rapidly transferred to IsdB\(^{\text{N2}}\), and the flexible loop between IsdB\(^{\text{N1}}\) and the linker is cleaved to allow for growth of the crystal. Interestingly, under physiological conditions, αHb oxidizes seven to ten times more quickly than βHb, especially at acidic pH (our crystals were produced at pH 5.5) (39, 40). The observation of heme removal from βHb in the crystal structure may be explained by the differential positioning of αHis\(^{99}\) and βHis\(^{97}\) in αHb versus βHb, respectively, which may be a rate-determining factor in the heme uptake mechanism. The asynchronous heme extraction from αHb and βHb observed in the crystal structure is mirrored by the kinetic analysis of heme transfer monitored at 408 nm. The four distinct kinetic phases observed may correspond to steps in heme extraction from the each Hb domain; however, alternative models are possible because of the presence of different subunits and the complex allosteric nature of Hb. The complexity of the heme transfer kinetics is expected for a substrate (Hb) that is a tetramer with two non-equivalent sites where the rate of heme extraction and the spectral change would vary with the heme site and the number of heme molecules bound to the...
tetramer. This complexity may also be hindering our attempts to observe spectral features of the putative bis-His intermediate. Nonetheless, the rates observed are similar or faster than the overall rate of heme transfer observed under steady-state conditions with IsdA as the ultimate heme recipient (18).

Superposition of the αβ tetramer of Hb in complex with IsdBN1N2 upon the tetramers of oxyHb (PDB code 2DN1), deoxyHb (PDB code 2DN2), or metHb (PDB code 3PSQ) revealed an unusual quaternary structure. The IsdBN1N2-bound tetramer exhibited neither a T-like (deoxy) nor an R-like (oxy or met) state conformation. This quaternary structure was also observed in a crystal structure of a heme-transfer deficient mutant of IsdHN2N3 in complex with metHb (PDB code 4XS0) (26). However, the overall IsdHN2N3-metHb structure is more similar to that of oxyHb (αHb, r.m.s.d. of 1.2 Å; βHb, r.m.s.d. of 0.5 Å) than our IsdBN1N2-metHb and structure oxyHb (αHb, r.m.s.d. of 3.3–3.4 Å; βHb, r.m.s.d. of 1.3–1.4 Å), Dickson et al. (26) noted that the Hb dimer in IsdBN1N2-metHb structure was most similar to structures of unusual liganded hemoglobins in a T-like state, such as a human sickle-cell variant of embryonic Hb. Our structure of IsdBN1N2-Hb supports the hypothesis that distortion of the Hb quaternary structure into a liganded T-like state is a characteristic of the Hb-binding and heme-extraction process by Isd proteins.

Hb is present at 10 mg/ml in normal serum (37) and has a high affinity for dimeric Hb ($K_d \approx 10^{-15}$ M (41)). Therefore, metHb present in serum is likely to exist as a complex with Hb. We have shown that heme transfer from metHb to IsdBN1N2 is inhibited by Hp, despite the ability of IsdB to bind Hp-Hb (9). In fact, under the conditions tested, heme transfer only occurred when the concentration of mixed-serotype Hp was decreased to 20 μg/ml, or ~500-fold lower than the concentration of Hp in human serum. The data presented here are consistent with a recent study demonstrating Hp inhibition of heme transfer from metHb to a similar IsdBN1N2 construct but not to an analogous IsdH construct (42). Moreover, in contrast to IsdH, IsdB did not inhibit binding of Hb-Hp to CD163 (42), the macrophage receptor for recycling HpH complexes (43).

Binding of Hp to Hb has also been shown to stabilize the heme–Hb interaction (44). A structural alignment of our IsdB-Hb crystal structure with the porcine (PDB code 4F4O) and human (PDB code 4XOL) Hb structures revealed no steric clashes between IsdBN1N2 and Hb (Fig. 10). Thus, inhibition of heme uptake from the Hp-Hb complex may reflect modulated stability of heme in the Hb pocket rather than direct blocking of the IsdB binding site by Hp. More specifically, examination of the Hp-Hb structures reveals a Hp loop (Pro327–Gly329 in human Hp; Pro268–Gly270 in porcine Hp) that interacts with F helix at Pro78 of αHb. This interaction needs to be disrupted to unwind the F helix for subsequent heme transfer to IsdB.

The IsdBN1N2 construct could possibly lack regions required to remove heme from Hp-Hb complexes in vitro. However, S. aureus was not able to use Hb-Hp as a sole iron source, whereas nanomolar concentrations of Hb alone supported S. aureus growth (Fig. 8). Previous studies have indicated that S. aureus can grow on Hb-Hp as sole iron source (45, 46); however, the quality of Hb was lower, and the concentration used in the growth medium was much higher than the more physiologically relevant concentration used here. In the work of Francis et al. (46), cells were grown in iron-rich tryptone and yeast extract, conditions under which the expression of the Isd system is repressed, and iron uptake was measured based on radioactivity of the cell pellet, a method that cannot distinguish between surface-bound and internalized iron. Recent studies have highlighted the importance of using human Hb from donors over commercial lyophilized Hb because of contamination with free heme (27, 47). During infection by S. aureus, hemolysin production and local hemolysis may be required to sufficiently increase the Hb concentration, allowing for excess Hb not bound to Hp.

**Experimental procedures**

**Cloning**

Plasmids were generated encoding residues 126 – 459 of IsdB with an N-terminal His tag and a thrombin cleavage site in pET28a(+)as previously described (18). Briefly, the IsdBN1N2 construct was subcloned from a GST-tagged construct cloned from S. aureus N315 chromosomal DNA (20) using a modified whole plasmid polymerase chain reaction method (48). A dou-
ble variant of IsdB\textsuperscript{N1N2} was created by subcloning from the IsdB\textsuperscript{N1N2} clone using the QuikChange site-directed mutagenesis kit (Agilent Technologies, Santa Clara, CA). All clones were verified by sequencing (Agencourt, Beverly, MA).

**Protein expression and purification**

Recombinant protein was overexpressed in *Escherichia coli* BL21 (DE3) cells. A 2-liter bacterial culture was grown from 2 ml of overnight culture in LB broth supplemented with 25 \(\mu\)g/ml kanamycin at 30 °C to an \(A_{600}\text{nm}\) of 0.7–0.9 and then induced with 0.5 mM of isopropyl \(\beta\)-D-thiogalactopyranoside and grown for another 18 h at 25 °C. The cells were pelleted by centrifugation; resuspended in 20 ml of 50 mM Tris, pH 8.0, 100 mM NaCl; and then lysed at 4 °C using an EmulsiFlex-C5 homogenizer (Avestin, Ottawa, Canada). Insoluble material was removed by centrifugation; the soluble lysate contained a mixture of apo and holo His\textsubscript{6} protein, and apo protein could be separated at 4 °C using a HisTrap nickel affinity column (GE Healthcare) by elution with an imidazole gradient. The apo protein was dialyzed against 50 mM Tris, pH 8.0, 100 mM NaCl and grown for another 18 h at 25 °C. The cells were pelleted by centrifugation; resuspended in 20 ml of 50 mM Tris, pH 8.0, 100 mM NaCl and then cleaved with thrombin at a 1:500 ratio by weight of His\textsubscript{6} protein to remove the His\textsubscript{6} tag, leaving behind a two amino acid (Gly-Ser) N-terminal artifact. Recombinant protein was then dialyzed against 20 mM HEPES, pH 7.4, for cation exchange chromatography using a Source 15S column (GE Healthcare), and purified protein was obtained by elution with a NaCl gradient. The resulting pure (>95% by SDS-PAGE) apo protein was dialyzed against 20 mM HEPES, pH 7.4, 80 mM NaCl for all studies. Hemoglobin was prepared from fresh human blood as described previously.

**Stopped-flow spectroscopic analysis of heme transfer from metHb to IsdB**

An SX.18MV stopped-flow reaction analyzer (Applied Photophysics Ltd., Leatherhead, UK) was used to investigate the possibility of heme-connection intermediates in the transfer process between metHb and IsdB\textsuperscript{N1N2}. The temperature of the optical cell and drive syringe chamber was maintained at 25 °C using a circulating water bath. Multiple wavelength data from 180 to 730 nm were collected using the photodiode array detector of the system directly coupled to the xenon light source (the practical range was from 250 to 730 nm). 20 \(\mu\)M apo-IsdB\textsuperscript{N1N2} and 2 \(\mu\)M metHb samples were in a buffer composed of 20 mM NaH\textsubscript{2}PO\textsubscript{4}, pH 7.4, 50 mM NaCl. Each acquisition was 15 s long with spectra collected at logarithmic intervals; four acquisitions were averaged.

A monochromator was used to collect single-wavelength data at 406 and 428 nm for the reaction of metHb and IsdB\textsuperscript{N1N2} in 20 mM HEPES, pH 7.4, 80 mM NaCl. The reactions were carried out with 2 \(\mu\)M metHb in one syringe (final concentration, 1 \(\mu\)M) and concentrations of apo-IsdB\textsuperscript{N1N2} ranging from 10 to 80 \(\mu\)M (final concentrations, 5–40 \(\mu\)M) in the second syringe; a minimum 5-fold excess of the IsdB acceptor was used to attain pseudo-first order conditions. The temperature was again maintained at 25 °C, and five 30-s acquisitions were performed at each wavelength for each concentration pair. The first 3 ms were in the dead time of the instrument and thus were excluded from analysis. At least four acquisitions were averaged, and curve fitting was performed using the ProDataSX software.

**Crystallization and data collection**

Solutions of apo-IsdB\textsuperscript{N1N2} and oxyHb in an equimolar (by monomer) ratio were mixed together at 10 mg/ml and immediately used to set up crystal trays. The IsdB\textsuperscript{N1N2}-Hb complex was crystallized in space group \(P2_12_12_2\) using reservoir solution containing 0.1 M malonic acid, pH 5.5, 2.2 M ammonium sulfate in 96-well sitting drop plates. Crystals appeared within a few days at room temperature but were allowed to grow to a sufficient size for ~4 weeks. Crystals were cryoprotected with 30% sodium malonate, pH 7.0, and flash frozen in liquid nitrogen. A 3.60 Å resolution data set was collected at the Stanford Synchrotron Radiation Lightsource on Beamline 7-1 (1.000 Å wavelength) at 100 K. The data were processed using HKL2000 (49) (Table 2). The data set was initially phased by molecular replacement using the oxyhemoglobin (PDB code 2DN1) and IsdB\textsuperscript{N2} (PDB code 3RTL) crystal structures and a SWISS-MODEL (50) generated IsdB\textsuperscript{N1} model (based on IsdB\textsuperscript{N2} PDB code 4IJ2) as search models with Phaser-MR (51) in the PHENIX suite of programs (52), which found the hemoglobin tetramer, three copies of IsdB\textsuperscript{N1} and three copies of IsdB\textsuperscript{N2}. Manual inspection of the IsdB\textsuperscript{N2} and IsdB\textsuperscript{N1} molecules, particularly at non-conserved bulky residues such as Tyr and Phe, revealed that electron density for one Phaser-placed copy of IsdB\textsuperscript{N1} was better fit by IsdB\textsuperscript{N2}, and the model was thus corrected to include four copies of IsdB\textsuperscript{N2} and two copies of IsdB\textsuperscript{N1}. SWISS-MODEL was also used to generate a model for the IsdB linker region based on the Hb-IsdH complex crystal structure (PDB code 4IJ2), which was added in a subsequent round of molecular replacement. Manual rebuilding with WinCoot (53) was used in all stages to complete the structure, and refinement was carried out using phenix.refine (54). Heme molecules were placed into omit difference maps, and the position of iron was confirmed by an anomalous difference map.

### Table 2

**X-ray data collection and refinement statistics for IsdB\textsuperscript{N1N2}-Hb structure**

| Data collection | Space group | \(P2_12_12_2\) |
|-----------------|-------------|----------------|
| Unit cell dimensions (Å) | \(a\) | 140.81 |
| | \(b\) | 209.11 |
| | \(c\) | 70.43 |
| Resolution range (Å) | 49.79–3.60 (3.66–3.60) |
| \(R_{	ext{merge}}\) | 0.12 (0.89) |
| \(I/\sigma(I)\) | 16.9 (2.1) |
| Completeness (%) | 99.8 (100) |
| Redundancy | 6.0 (6.0) |
| Unique reflections | 24903 |
| Wilson B (Å\(^2\)) | 116.5 |
| Refinement | No. atoms | 11827 |
| | Protein/heme | 11827 |
| | Water | 0 |
| Average \(B\)-factors (Å\(^2\)) | Hemoglobin | 102.2 |
| | IsdB | 116.3 |
| | Heme | 112.4 |
| | r.m.s.d. | 0.003 |
| | Bond lengths (Å) | 0.80 |
| | Bond angles (°) | |
| | PDB entry code | 5VMM |
The heme-iron ligand geometry was not restrained. The poly-peptide chain has excellent stereochemistry, with 95.2% of residues in the favored region of the Ramachandran plots and 4.5% in allowed regions, and 0.3% were outliers.

**Hb+IsdB**\(^{N1N2}\) crystal composition

Crystals from a well of oxyHb+IsdB\(^{N1N2}\) were looped and soaked in well solution to remove adventitiously bound protein and then dissolved in 5 \(\mu l\) of 20 mM HEPES, pH 7.4, 80 mM NaCl. Five small crystals in total were dissolved together and separated on a 15% SDS-PAGE gel to visualize the components of the crystal.

**Haptoglobin inhibition assay**

Human Hp of mixed serotype purified from plasma was purchased as a lyophilized solid (Athens Research and Technology, Athens, GA). 1 mg of Hp was dissolved in 100 \(\mu l\) of 20 mM HEPES, pH 7.4, 80 mM NaCl to give a concentration of 10 mg/ml. Spectra of 25 °C. Syringe and cell samples were co-dialyzed overnight at 200 V for 1 h and 5 min and stained using Coomassie Blue.

**IsdB**\(^{VYFY}\) titration of metHb

Spectra (250–750 nm) of metHb, IsdB constructs, or mixtures thereof were taken in a conventional Cary50 spectrophotometer with an optical path length of 1 cm in a quartz cell at room temperature (22 °C). 5 \(\mu l\) of apo-IsdB\(^{N1N2}\) or apo-IsdB\(^{VYFY}\), and spectra were immediately recorded; spectra did not change within 5 min of first recording. Additionally, 4 \(\mu l\) of His6-metHb in a total volume of 1000 \(\mu l\) was titrated with aliquots (3.7 \(\mu l\)) of apo-IsdB\(^{VYFY}\) in 1 \(\mu l\) increments from 1 to 9 \(\mu l\). Spectra (250–750 nm) were taken in a Cary50 with an optical path length of 1 cm in a quartz cell at 22 °C, with three replicates carried out. The initial absorbance for each replicate at 410 nm (i.e. 0 \(\mu l\) IsdB\(^{VYFY}\)) was subtracted from each subsequent titration absorbance value to yield the \(\Delta A\)110 nm, which was plotted against IsdB\(^{VYFY}\) concentration; a straight line could be fit through the first five titration points to indicate stoichiometry.

**S. aureus growth experiments**

RPMI media 1640 with l-glutamine and Na\(_2\)HCO\(_3\) (Sigma–Aldrich) was supplemented with 1% (w/v) casamino acids (BD, Sparks, MD). Metal-depleted RPMI (NRPMI) medium was prepared by first adding 7% (w/v) Chelex-100 (Sigma–Aldrich) to RPMI medium and stirring overnight. Chelex-100 was then removed from the medium, and 25 mM ZnCl\(_2\), 25 mM MnCl\(_2\), 100 mM CaCl\(_2\), and 1 mM MgCl\(_2\) were added. NRPMI medium was adjusted to pH 7.4, filter-sterilized, and stored at 4 °C.

Single colonies of *S. aureus* str. Newman on tryptic soy agar were inoculated into RPMI medium with 0.5 mM ethylenediamine-N\(_2\),N’-bis(2-hydroxyphenylacetic acid) (EDDHA, LG Standards GmbH, Teddington, UK). The cultures were incubated at 37 °C for 16–20 h on a shaker set to 200 rpm. Overnight cultures were centrifuged for 2 min at 11,000 \(\times\) g, and the pellet was washed three times in NRPMI with 0.5 mM EDDHA. The cells were then normalized to OD \(_{600}\) and subcultured 1:100 into 200 \(\mu l\) of NRPMI with 0.5 mM EDDHA. In addition to a no-iron control, cultures were supplemented with 200 \(nM\) Hb, 200 \(nM\) human Hb, and 4–400 \(\mu g/ml\) Hp, mixed serotype (Athens Research and Technology), or 2 \(\mu M\) of heme with and without 400 \(\mu g/ml\) haptoglobin. The cultures were grown in triplicate with constant shaking at fast-speed and high-amplitude settings in a Bioscreen C instrument (Growth Curve USA, Piscataway, NJ) at 37 °C. Growth experiments were repeated three times, and the average growth with standard deviations are plotted using GraphPad Prism 6 (GraphPad Software, La Jolla, CA).

**Author contributions**—C. F. M. B. and M. E. P. M. conceptualized the study and designed the experiments; C. F. M. B., A. C. K. C., E. J. W. L., and A. L. A. conducted the experiments; C. F. M. B., A. C. K. C., and E. J. W. L. analyzed the data; C. F. M. B. drafted the article, with critical revisions by A. C. K. C., L. D. E., and M. E. P. M. All authors reviewed the results and approved the final version of the manuscript.
S. aureus IsdB-associated unfolding of human hemoglobin

Acknowledgments—Use of the Stanford Synchrotron Radiation Lightsource, Stanford Linear Accelerator Center National Accelerator Laboratory is supported by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences under Contract DE-AC02-76SF00515. The Stanford Synchrotron Radiation Lightsource Structural Molecular Biology Program is supported by the Department of Energy Office of Biological and Environmental Research, and by the NIGMS, National Institutes of Health Grant P41GM103393.

References

1. Wertheim, H. F., Melles, D. C., Vos, M. C., van Leeuwen, W., van Belkum, A., Verbrugh, H. A., and Nouwen, J. L. (2005) The role of nasal carriage in Staphylococcus aureus infections. Lancet Infect. Dis. 5, 751–762 CrossRef Medline

2. Wispelwijn, H., Bischot, T., Tallent, S. M., Seifert, H., Wenzel, R. P., and Edmond, M. B. (2004) Nosocomial bloodstream infections in US hospitals: analysis of 24,179 cases from a prospective nationwide surveillance study. Clin. Infect. Dis. 39, 309–317 CrossRef Medline

3. Waness, A. (2010) Revisiting methicillin-resistant Staphylococcus aureus infections. J. Glob Infect. Dis. 2, 49–56 CrossRef Medline

4. Weinberg, E. D. (1975) Nutritional immunity: host’s attempt to withhold iron from microbial invaders. JAMA 231, 39–41 CrossRef Medline

5. Skaar, E. P., Humayun, M., Bae, T., DeBord, K. L., and Schneewind, O. (2004) Iron-source preference of Staphylococcus aureus infections. Science 305, 1626–1628 CrossRef Medline

6. Skaar, E. P., and Schneewind, O. (2004) Iron-regulated surface determinants (Ist) of Staphylococcus aureus: stealing iron from heme. Microbes Infect. 6, 390–397 CrossRef Medline

7. Mazmanian, S. K., Ton-That, H., Su, K., and Schneewind, O. (2002) An iron-regulated sortase anchors a class of surface protein during Staphylococcus aureus pathogenesis. Proc. Natl. Acad. Sci. U.S.A. 99, 2293–2298 CrossRef Medline

8. Mazmanian, S. K., Skaar, E. P., Gaspar, A. H., Humayun, M., Gornicki, P., Jelenska, J., Joachmiak, A., Missiakas, D. M., and Schneewind, O. (2003) Passage of heme-iron across the envelope of Staphylococcus aureus. Science 299, 906–909 CrossRef Medline

9. Dryla, A., Hoffmann, B., Gelbmann, D., Gieffing, C., Hanner, M., Meinke, A., Anderson, A. S., Koppensteiner, W., Konrat, R., von Gabain, A., and Nagy, E. (2007) High-affinity binding of the staphylococcal HarA protein to haptoglobin and hemoglobin involves a domain with an antiparallel eight-stranded β-barrel fold. J. Bacteriol. 189, 254–264 CrossRef Medline

10. Torres, V. J., Pischchany, G., Humayun, M., Schneewind, O., and Skaar, E. P. (2006) Staphylococcus aureus IsdB is a hemoglobin receptor required for heme iron utilization. J. Bacteriol. 188, 8421–8429 CrossRef Medline

11. Zhu, H., Xie, G., Liu, M., Olson, J. S., Fabian, M., Dooley, D. M., and Lei, B. (2008) Pathway for heme uptake from human methemoglobin by the iron-regulated surface determinants system of Staphylococcus aureus. J. Biol. Chem. 283, 18450–18460 CrossRef Medline

12. Muryo, N., Tiedemann, M. T., Pluym, M., Cheung, J., Heinrichs, D. E., and Stillman, M. J. (2008) Demonstration of the iron-regulated surface determinant (Isd) heme transfer pathway in Staphylococcus aureus. J. Biol. Chem. 283, 28125–28136 CrossRef Medline

13. Grigg, J. C., Vermeiren, C. L., Heinrichs, D. E., and Murphy, M. E. (2007) Heme coordination by Staphylococcus aureus IsdE. J. Biol. Chem. 282, 28815–28822 CrossRef Medline

14. Lee, W. C., Reniere, M. L., Skaar, E. P., and Murphy, M. E. (2008) Ruffling of metalloporphyrins bound to IsdG and IsdI, two heme-degrading enzymes in Staphylococcus aureus. J. Biol. Chem. 283, 30957–30963 CrossRef Medline

15. Reniere, M. L., Ukpb, G. N., Harry, S. R., Stee, D. F., Krull, R., Wright, D. W., Bachmann, B. O., Murphy, M. E., and Skaar, E. P. (2010) The IsdG-family of heme oxygenases degrades heme to a novel chromophore. Mol. Microbiol. 75, 1529–1538 CrossRef Medline

16. Andrade, M. A., Ciccarelli, F. D., Perez-Iratxeta, C., and Bork, P. (2002) NEAT: a domain duplicated in genes near the components of a putative Fe3+ siderophore transporter from Gram-positive pathogenic bacteria. Genome Biol. 3, RESEARCH0047 CrossRef Medline

17. Krishna Kumar, K., Jacques, D. A., Pischchany, G., Caradoc-Davies, T., Spriog, T., Malmircherghini, G. R., Langley, D. B., Dickson, C. F., Mackay, J. P., Clubb, R. T., Skaar, E. P., Guss, J. M., and Gell, D. A. (2011) Structural basis for hemoglobin capture by Staphylococcus aureus cell-surface protein, IsdH. J. Biol. Chem. 286, 38439–38447 CrossRef Medline

18. Bowden, C. F., Verstraete, M. M., Eltis, L. D., and Murphy, M. E. (2014) Hemoglobin binding and catalytic heme extraction by IsdB near iron transporters. Biochemistry 53, 2286–2294 CrossRef Medline

19. Dickson, C. F., Kumar, K. K., Jacques, D. A., Malmircherghini, G. R., Spriog, T., Mackay, J. P., Clubb, R. T., Guss J. M., and Gell, D. A. (2011) Structure of the hemoglobin-IsdH complex reveals the molecular basis of iron capture by Staphylococcus aureus. J. Biol. Chem. 289, 6728–6738 CrossRef Medline

20. Gaudin, C. F., Grigg, J. C., Arrieta, A. L., and Murphy, M. E. (2011) Unique heme-iron coordination by the hemoglobin receptor IsdB of Staphylococcus aureus. Biochemistry 50, 5443–5452 CrossRef Medline

21. Grigg, J. C., Vermeiren, C. L., Heinrichs, D. E., and Murphy, M. E. (2007) HAem recognition by a Staphylococcus aureus NEAT domain. Mol. Microbiol. 63, 139–149 CrossRef Medline

22. Pilpa, R. M., Fadeev, A. E., Villareal, V. A., Wong, M. L., Phillips, M., and Clubb, R. T. (2006) Solution structure of the NEAT (NEAR Transporter) domain from IsdH/HarA: the human hemoglobin receptor in Staphylococcus aureus. J. Mol. Biol. 360, 435–447 CrossRef Medline

23. Sharp, K. H., Schneider, S., Cockayne, A., and Paoli, M. (2007) Crystal structure of the heme-IsdC complex, the central conduit of the Isd iron/ heme uptake system in Staphylococcus aureus. J. Biol. Chem. 282, 10625–10631 CrossRef Medline

24. Watanabe, M., Tanaka, Y., Suenaga, A., Kuroda, M., Yao, M., Watanabe, N., Arisaka, F., Ohta, T., Tanaka, I., and Tsumoto, K. (2008) Structural basis for multimeric heme complexation through a specific protein-heme interaction: the case of the third neut domain of IsdH from Staphylococcus aureus. J. Biol. Chem. 283, 28649–28659 CrossRef Medline

25. Honner, B. A., Tripet, B. P., Elers, B. J., Stanisch, J., Sullivan-Sprinquette, R. K., Moore, R., Liu, M., Lei, B., and Copie, V. (2014) Solution structure and molecular determinants of hemoglobin binding of the first NEAT domain of IsdB in Staphylococcus aureus. Biochemistry 53, 3922–3933 CrossRef Medline

26. Dickson, C. F., Jacques, D. A., Clubb, R. T., Guss, J. M., and Gell, D. A. (2015) The structure of haemoglobin bound to the haemoglobin receptor IsdH from Staphylococcus aureus shows disruption of the native α-globin haem pocket. Acta Crystallogr. D Biol. Crystallogr. 71, 1295–1306 CrossRef Medline

27. Pischchany, G., Sheldon, J. R., Dickson, C. F., Alam, M. T., Read, T. D., Gell, D. A., Heinrichs, D. E., and Skaar, E. P. (2014) IsdB-dependent hemoglobin binding is required for acquisition of heme by Staphylococcus aureus. Infect. Immun. 82, 2215–2223 CrossRef Medline

28. Allard, M., Moisan, H., Brouillette, E., Gervais, A. L., Jacques, M., Lacasse, P., Diarra, M. S., and Malouin, F. (2006) Transcriptional modulation of some Staphylococcus aureus iron-regulated genes during growth in vitro and in a tissue case model in vivo. Microbes Infect. 8, 1679–1690 CrossRef Medline
S. aureus IsdB-associated unfolding of human hemoglobin

32. Pilpa, R. M., Robson, S. A., Villareal, V. A., Wong, M. L., Phillips, M., and Clubb, R. T. (2009) Functionally distinct NEAT (NEAr Transporter) domains within the Staphylococcus aureus IsdB/HarA protein extract heme from methemoglobin. J. Biol. Chem. 284, 1166–1176 CrossRef Medline

33. Robinson, V. L., Smith, B. B., and Arnone, A. (2003) A pH-dependent aquomet-to-hemichrome transition in crystalline horse methemoglobin. Biochemistry 42, 10113–10125 CrossRef Medline

34. Rachmilewitz, E. A., Peisach, J., and Blumberg, W. E. (1971) Studies on the interaction between human hemoglobin and its constituent chains and their derivatives. J. Biol. Chem. 246, 3356–3366 Medline

35. Langlois, M. R., and Delanghe, J. R. (1996) Biological and clinical significance of haptoglobin polymorphism in humans. Clin. Chem. 42, 1589–1600 Medline

36. Wobeto, V. P. d. A., Zaccariotto, T. R., and Sonati, M. d. F. (2008) Polymeric forms of haptoglobin and its clinical importance. Genet. Mol. Biol. 31, 602–620 CrossRef

37. Cheng, T. M., Pan, J. P., Lai, S. T., Kao, L. P., Lin, H. H., and Mao, S. I. (2007) Immunochemo property of human haptoglobin phenotypes: determination of plasma haptoglobin using type-matched standards. Clin. Biochem. 40, 1045–1056 CrossRef Medline

38. Eaton, J. W., Brandt, P., Mahoney, J. R., and Lee, J. T., Jr (1982) Haptoglobin: a natural bacteriostat. Science 215, 691–693 CrossRef Medline

39. Langlois, M. R., and Delanghe, J. R. (1996) Biological and clinical significance of haptoglobin polymorphism in humans. Clin. Chem. 42, 1589–1600 Medline

40. Tsuruga, M., Matsuoka, A., Hachimori, A., Sugawara, Y., and Shikama, K. (1998) The molecular mechanism of autoxidation for human oxyhemoglobin: a natural bacteriostat. Biochem. J. 329, 667–673 CrossRef Medline

41. Hwang, P. K., and Greer, J. (1980) Interaction between hemoglobin subunits in the hemoglobin-haptoglobin complex. J. Biol. Chem. 255, 3038–3041 Medline

42. Sæderup, K. L., Stokilde, K., Graversen, J. H., Dickson, C. F., Etzerodt, A., Hansen, S. W., Fago, A., Gell, D., Andersen, C. B., and Moestrup, S. K. (2016) The Staphylococcus aureus protein IsdH inhibits host hemoglobin scavenging to promote heme acquisition by the pathogen. J. Biol. Chem. 291, 23989–23998 CrossRef Medline

43. Nielsen, M. J., Andersen, C. B., and Moestrup, S. K. (2013) CD163 binding to haptoglobin-hemoglobin complexes involves a dual-point electrostatic receptor-ligand pairing. J. Biol. Chem. 288, 18834–18841 CrossRef Medline

44. Bunn, H. F., and Landl, J. H. (1968) Exchange of heme among hemoglobins and between hemoglobin and albumin. J. Biol. Chem. 243, 465–475 Medline

45. Dryla, A., Gelmann, D., von Gabain, A., and Nagy, E. (2003) Identification of a novel iron regulated staphylococcal surface protein with haptoglobin-hemoglobin binding activity. Mol. Microbiol. 49, 37–53 CrossRef Medline

46. Francis, R. T., Jr., Booth, J. W., and Becker, R. R. (1985) Uptake of iron from hemoglobin and the haptoglobin-hemoglobin complex by hemolytic bacteria. Int. J. Biochem. 17, 767–773 CrossRef Medline

47. Pischhany, G., Haley, K. P., and Skaar, E. P. (2013) Staphylococcus aureus growth using human hemoglobin as an iron source. J. Vis. Exp. 72, e50072

48. MacPherson, I. S., Rosell, F. I., Scofield, M., Mauk, A. G., and Murphy, M. E. (2010) Directed evolution of copper nitrite reductase to a chromogenic reductant. Protein Eng., Des. Sel. 23, 137–145 CrossRef Medline

49. Otwinowski, Z., and Minor, W. (1997) Processing of X-ray diffraction data collected in oscillation mode. Methods Enzymol. 276, 307–326 CrossRef

50. Biasini, M., Bienert, S., Waterhouse, A., Arnold, K., Studer, G., Schmidt, T., Kieber, F., Cassarino, T. G., Bertoni, M., Bordoli, L., and Schwede, T. (2014) SWISS-MODEL: modelling protein tertiary and quaternary structure using evolutionary information. Nucleic Acids Res. 42, W252–W258 CrossRef Medline

51. McCoy, A. J., Grosse-Kunstleve, R. W., Adams, P. D., Winn, M. D., Storoni, L. C., and Read, R. J. (2007) Phaser crystallographic software. J. Appl. Crystallogr. 40, 658–674 CrossRef Medline

52. Adams, P. D., Afonine, P. V., Bunkóczi, G., Chen, V. B., Davis, I. W., Echols, N., Headd, J. J., Hung, L. W., Kapral, G. J., Grosse-Kunstleve, R. W., McCoy, A. J., Moriarty, N. W., Oeffner, R., Read, R. J., Richardson, D. C., et al. (2010) PHENIX: a comprehensive Python-based system for macromolecular structure solution. Acta Crystallogr. D Biol. Crystallogr. 66, 213–221 CrossRef Medline

53. Emsley, P., Lohkamp, B., Scott, W. G., and Cowtan, K. (2010) Features and development of Coot. Acta Crystallogr. D Biol. Crystallogr. 66, 486–501 CrossRef Medline

54. Afonine, P. V., Grosse-Kunstleve, R. W., Echols, N., Headd, J. J., Moriarty, N. W., Mustyakimov, M., Terwilliger, T. C., Urzhumtsev, A., Zwart, P. H., and Adams, P. D. (2012) Towards automated crystallographic structure refinement with phenix.refine. Acta Crystallogr. D Biol. Crystallogr. 68, 352–367 CrossRef Medline