Functionally distinct NEAT (NEAr Transporter) domains within the 
*Staphylococcus aureus* IsdH/HarA protein extract heme from Methemoglobin

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The pathogen *Staphylococcus aureus* uses Iron-regulated surface determinant (Isd) proteins to scavenge the essential nutrient iron from host hemoproteins. The IsdH protein (also known as HarA) is a receptor for hemoglobin (Hb), haptoglobin (Hp) and the Hb-Hp complex. It contains three NEAT (NEAr Transporter) domains: IsdH\textsuperscript{N1}, IsdH\textsuperscript{N2} and IsdH\textsuperscript{N3}. Here we show that they have different functions; IsdH\textsuperscript{N1} and IsdH\textsuperscript{N2} bind Hb and Hp, while IsdH\textsuperscript{N3} captures heme that is released from Hb. The staphylococcal IsdB protein also functions as an Hb receptor. Primary sequence homology to IsdH indicates that it will also employ functionally distinct NEAT domains to bind heme and Hb. We have used site-directed mutagenesis and surface plasmon resonance methods to localize the Hp and Hb binding surface on IsdH\textsuperscript{N1}. High affinity binding to these structurally unrelated proteins requires residues located within a conserved aromatic motif that is positioned at the end of the beta barrel structure. Interestingly, this site is quite malleable as other NEAT domains use it to bind heme. We also demonstrate that the IsdC NEAT domain can capture heme directly from Hb, suggesting that there are multiple pathways for heme transfer across the cell wall.

*Staphylococcus aureus* causes a wide range of life threatening diseases such as pneumonia, septicemia, osteomyelitis, toxic shock syndrome, and cardiomycelitis (1). The rise of antibiotic resistance has created a pressing need for new drugs to treat infections caused by this microbe as methicillin-resistant *S. aureus* (MRSA) now accounts for up to ~65% of all clinical isolates, and new multi-drug resistant strains have emerged that are resistant to vancomycin, the preferred antibiotic of "last resort" to treat MRSA infections (2). Bacterial growth is limited by the availability of iron, which is an essential cofactor for enzymes involved in microbial metabolism. Although the human body contains large quantities of iron, little is directly available to *S. aureus* because it is sequestered intracellularly or bound to the carrier glycoproteins transferrin and lactoferrin. To overcome this bacteriostatic limitation, *S. aureus* has evolved a variety of mechanisms to assimilate iron (3,4). Of special interest are recently discovered iron-regulated surface determinant (Isd) proteins, which extract heme bound iron from host hemoproteins (5-8). The Isd system likely plays a significant role in virulence as heme contains ~75% of the body's total iron and it is *S. aureus*’ preferred source of iron (9). As sequence homologs of Isd proteins are also found in a number of other important pathogens (e.g. *Listeria monocytogenes*, *Bacillus anthracis*, and *Streptococcus pyogenes*), a greater understanding of the Isd-system could lead to the development of new therapeutically useful anti-infective agents (10-12).

Hemoglobin (Hb) in red blood cells contains large quantities of heme-iron, to which *S. aureus* gains access by secreting cytolytic toxins (leukocidin, and α- and δ-hemolysin) (13). Release of Hb from lysed erythrocytes significantly dilutes the protein causing it to dissociate and oxidize into methemoglobin (MetHb), a dimeric form of Hb that contains
single α and β globin chains as a heterodimer, each bound to a ferric heme (14,15). Because heme readily dissociates from MetHb and it can cause severe oxidative damage to host tissues or serve as a microbial nutrient, several host mechanisms exist to remove it from circulation (16). First, heme-laden MetHb present in the blood is tightly bound by the serum glycoprotein haptoglobin (Hp), and the Hp-MetHb complex is cleared from circulation by receptor-mediated endocytosis in the liver parenchymal cells (17,18). Second, free heme in the blood is eliminated when it binds to hemopexin and serum albumin proteins (15). Recent studies have begun to elucidate how \textit{S. aureus} circumvents these clearance systems in order to gain access to heme-iron. Free heme present in the blood can be scavenged by the heme transport system (Hts), a membrane protein complex that pumps heme into the cytoplasm (9). Heme is also aggressively acquired by capturing it from circulating MetHb and the Hp-MetHb complex using the Isd system (6,19). Although both the Hts- and Isd-systems import heme and are important for virulence, \textit{Staphylococcal} strains with disrupted Isd components cause less persistent infections in mice than wildtype strains and are unable to utilize Hb as an iron source in cell cultures (20,21).

The Isd pathway consists of nine proteins that work in concert to acquire heme (5-8). Three receptor proteins are covalently anchored to the cell wall by the SrtA sortase and are displayed on or near the cell surface (IsdA, IsdB, and IsdH (also known as HarA (Haptoglobin receptor A))) (6,19,22). In addition, a fourth receptor, IsdC, is attached to the cell wall by the SrtB sortase enzyme (21). MetHb is captured by IsdH and IsdB (6,19,20). Although these proteins share a high degree of sequence homology with one another, only IsdH is capable of binding Hp (19,23). In addition, full-length IsdH binds to heme (6), whereas it is unknown if IsdH can also bind to this molecule. The IsdA protein binds heme and may play a general role in bacterial adhesion, since the isolated protein interacts with an array of extracellular matrix proteins (22). After capture by the surface receptors, heme is thought to be transferred to IsdC, which is presumably embedded within the cell wall because it resists proteolytic degradation in whole cell digestion studies (6). The IsdC protein then passes the heme molecule to the membrane transporter IsdDEF complex where it is imported into the cytoplasm. Finally, iron is released when the tetrapyrrrole ring is cleaved by the monooxygenase IsdG or its paralog IsdI (24). Recent studies also suggest that heme imported by the Isd system can be directly incorporated in microbial proteins (8).

The Isd receptors IsdA, IsdB, IsdC, and IsdH bind to proteins and heme using NEAT (NEAR iron Transporter) domains. These conserved binding modules are ~125 residues in length and are named for their prevalence in bacterial genes whose genomic positioning is proximal to putative Fe^{3+} siderophore transporter genes (10). NEAT domains are also found in putative surface proteins in a number of other Gram-positive pathogens where they presumably function in iron import. Studies of isolated NEAT domains from \textit{S. aureus} have revealed that different domains bind to distinct ligands. For example, some NEAT domains only bind heme (e.g. IsdC) or only to other proteins (e.g. the first and second domains of IsdH), while other NEAT domains can bind both of these ligands (e.g. IsdA) (19,22,23,25). NMR and crystallography studies have revealed that NEAT domains adopt a β-sandwich fold that shares structural homology with immunoglobin-like proteins (23,25-28). In addition, the structures of the IsdC- and IsdA-heme complexes have revealed the mechanism of heme binding (26-28). However, it remains unknown how NEAT domains interact with other proteins.

The IsdH protein captures MetHb and Hp on the cell surface (19). Sequence analysis indicates that it contains three NEAT domains: IsdH^{N1}, IsdH^{N2}, and IsdH^{N3} (Fig. 1A). While the IsdH^{N1} and IsdH^{N2} domains bind to MetHb and Hp, the function of the C-terminal IsdH^{N3} domain is unknown (19). In this study we systematically investigated the functions of NEAT domains within IsdH. We show using alanine scanning mutagenesis and surface plasmon resonance (SPR) that Hp and MetHb bind at the same site on IsdH (19). Interestingly, this site is also used by the IsdA and IsdC NEAT domains to bind heme, suggesting that this surface is particularly well suited for recognizing a range of distinct ligands (26-28). We also show that the previously uncharacterized IsdH^{N3} domain binds heme and that it can capture this molecule from MetHb. A model of heme transfer is proposed in which...
domains IsdH\textsuperscript{N1} and IsdH\textsuperscript{N2} first bind MetHb or the Hp-MetHb complex on the cell surface. Heme is then captured by the adjacent IsdH\textsuperscript{N3} domain before subsequent transfer through the cell wall.

**EXPERIMENTAL PROCEDURES**

**Protein expression and purification.** The first NEAT domain from IsdH consists of residues Ala\textsuperscript{86} to Leu\textsuperscript{229} and is referred to as IsdH\textsuperscript{N1} (Fig. 1A). Two versions of IsdH\textsuperscript{N1} were studied. For the heme transfer assay untagged IsdH\textsuperscript{N1} was produced from a pet11a vector (Novagen) and purified as previously described (25,29). For the binding studies using surface plasmon resonance (SPR), a histidine-tagged fusion of IsdH\textsuperscript{N1} was used (his-IsdH\textsuperscript{N1}, residues Ala\textsuperscript{86} to Leu\textsuperscript{229} with the sequence Met-Gly-Ser-Ser-His-His-His-His-His-Ser-Ser-Gly-Leu-Val-Pro-Arg-Gly-Ser-His-Met at its N-terminus). Briefly, the DNA for IsdH\textsuperscript{N1} was PCR amplified from \textit{S. aureus} chromosomal DNA (strain RN4220) using primers that incorporated NdeI and BamHI restriction sites at the 5’ and 3’ ends of the DNA, respectively. This PCR product was then digested with NdeI and BamHI and ligated into a similarly cut pET15b vector. Plasmids encoding single amino acid mutants of his-IsdH\textsuperscript{N1} were constructed from the pET15b vector bearing the IsdH\textsuperscript{N1} sequence using the QuickChange Site-Directed Mutagenesis Kit (Stratagene) according to the manufacturer’s specifications. \textit{E. coli} BL21(DE3) cells containing the plasmids of either his-tagged wild-type or mutant IsdH\textsuperscript{N1} were harvested four hours after the addition of IPTG (isopropyl β-D-1-thiogalactopyranoside, Sigma) by centrifugation for 15 min at 5000 rpm at 4 °C in a JA-10 rotor, resuspended in lysis buffer (50 mM Tris-HCl (pH 7); 2.5 mM benzamidine (Acros); 1 mM PMSF (phenylmethylsulphonyl fluoride, Sigma-Aldrich); protease inhibitor cocktail II (Roche)), and lysed by three cycles of sonication. The lysed cells were centrifuged at 4 °C for one hour at 12,000 rpm in a SS-34 rotor. The supernatant was then applied to Talon nickel affinity beads (Clontech) and the his-tagged fusion proteins were purified according to the manufacturer’s instructions. All his-IsdH\textsuperscript{N1} mutants were checked for proper folding using 1D NMR (data not shown).

Two proteins were used as heme receptors in the heme transfer assay, histidine tagged IsdC (his-IsdC, residues Ser\textsuperscript{25} to Gly\textsuperscript{150}) and histidine tagged C-terminal NEAT domain from IsdH (his-IsdH\textsuperscript{N3}, residues Thr\textsuperscript{540} to Gln\textsuperscript{664}). Both proteins were expressed from pET15b plasmids (Novagen) and contained the amino acid sequence Met-Gly-Ser-His-His-His-His-Ser-Gly-Leu-Val-Pro-Arg-Gly-Ser-His-Met at their N-termini. Plasmid construction and protein purification procedures were identical to those used for his-IsdH\textsuperscript{N1} (described above). The amount of heme associated with his-IsdC and his-IsdH\textsuperscript{N3} was determined using a pyridine hemochrome assay (30). After purification, ~2-3% of his-IsdH\textsuperscript{N3} and his-IsdC was bound to heme.

**Enzyme-Linked Immunoabsorbent Assay (ELISA).** An enzyme-linked immunoabsorbent assay (ELISA) was used to test his-IsdH\textsuperscript{N3} binding to a variety of proteins. ELISA plate wells (Nunc microwell plates, Fisher Scientific) were coated with a 150 µL solution containing 20 µg/ml of bovine serum albumin solution in PBS buffer for one hour with nutation. The wells were then washed three times with PBS to remove excess blocking solution, and blocking solution was added to each well (0 to 30 picomoles per well). Interactions between his-IsdH\textsuperscript{N3} and protein ligands were then detected by adding nickel-labeled horseradish peroxidase according to the manufacturer’s directions (Express Detector Nickel-HRP, KPL). After incubating for one hour at 37 °C, the wells were washed with PBS to remove unbound nickel-labeled horseradish peroxidase, and the binding of his-IsdH\textsuperscript{N3} was quantified by measuring the conversion of the substrate, TMB (3,3',5,5'-tetramethylbenzidine, Sigma) to a yellow color at 450 nm.
tetramethylbenzidine (Sigma-Aldrich) dissolved in citric phosphate buffer, pH 5) to a colored product using an automated ELISA reader (Molecular Devices, SpectraMax M5). As a positive control similar experiments were performed using his-IsdH3 protein instead of his-IsdH3N.

**Metalloprotoporphyrin affinity measurements.** UV and fluorescence spectroscopy were used to determine the affinity of IsdH3N for metalloprotoporphyrins. In the UV assay the soret band of his-IsdH3N at 402 nm was monitored as a function of heme (iron protoporphyrin XI) added. The concentration of his-IsdH3N was held constant at 1 µM in buffer A (50mM Tris HCl, pH 7.5; 100 mM NaCl) and hemin was added in small aliquots from a 1 mM stock solution (hemin dissolved in 0.1 M NaOH). After pH adjustment, the UV absorbance was measured and plotted as a function of heme concentration to yield a binding curve. All UV measurements were performed using a PharmaSpec UV-1700 spectophotometer (Shimadzu). The affinity of IsdH3N for the heme analog zinc protoporphyrin IX (ZnPPIX) was determined using fluorescence spectroscopy. In this assay the concentration of ZnPPIX was held constant and its fluorescence was monitored as a function of protein added. Binding reactions contained 150 µL of 7.5 µM ZnPPIX (Sigma-Aldrich) dissolved in binding buffer (50mM potassium phosphate; 100 mM NaCl, pH 7.5). Aliquots of his-IsdH3N from a 125 µM stock solution were added to a final concentration of 50 µM. Fluorescence was measured using a Quanta Master Model QM5 spectrofluorimeter (Photon Technology International) at room temperature. The excitation and emission wavelengths were 365 nm and 585 nm, respectively. Binding data were fit to the equation 

\[ A = A_t + (A_b-A_t)(K_a[L]/1+K_a[L]), \]

where A is the fluorescence emission, A_t is the fluorescence emission of free ZnPPIX, A_b is the fluorescence of bound ZnPPIX, and K_a is the association constant (31). Data were fit using the program SigmaPlot 2000.

**Surface plasmon resonance (SPR) affinity measurements.** Binding of IsdH1N to immobilized MetHb was measured by SPR. MetHb was immobilized on a CM5 chip using the following procedure. First, a 0.1 mg/ml stock solution of MetHb was prepared by dissolving commercially available MetHb (MetHb, Sigma) in 50 mM sodium phosphate pH 6.2, 100 mM NaCl. The stock solution was then diluted to 1µg/ml in Biacore binding buffer (10 mM sodium acetate pH 5.5) and immobilized on a CM5 chip using N-ethyl-N-(dimethylaminopropyl)carboimide (EDC)/N-hydroxysuccinimide (NHS) according to the manufacturers instructions. After MetHb immobilization, the final measured response units (RU) of the CM5 chip was ~1000. Measurement of IsdH1N binding was performed at 10 °C. Wild-type and mutant IsdH1N was flowed over the MetHb conjugated surface at a rate of 85 µL/min with a contact time of 150 seconds and a dissociation time of 300 seconds. Surface plasmon resonance (SPR) measurements were taken at concentrations of 0, 50, 100, 200, 300, 400, 500, 1000, 2000, and 4000 nM IsdH1N using Buffer A (50 mM sodium phosphate, pH 6.2; 100mM NaCl) as the flow buffer. A similar procedure was used to measure IsdH1N binding to Hp. Haptoglobin from pooled human plasma (Athens research) was immobilized on a CM5 chip using the aforementioned methods. This protein consists of two alleles of haptoglobin, α-1 and α-2. These proteins differ in their oligomerization state as a result of differences in their amino-acid sequence. A 100 µg/ml solution of Hp dissolved in Biacore binding buffer was immobilized so as to yield a haptoglobin conjugated chip with a RU of ~1000. SPR experiments were performed on a Biacore T100 instrument and the data was analyzed using the program BIAevaluation 4.01 (Amersham Pharmacia Biotech).

**Heme transfer assay.** A total of 0.3 µmol of either his-IsdH3N or his-IsdC was incubated with 750 µl of Talon beads (Clontech) dissolved in Buffer A (50 mM sodium phosphate pH 6.0; 100 mM NaCl) for one hour. The beads were then centrifuged, washed and equilibrated in Buffer B (50 mM Tris HCl pH 7.2; 300 mM NaCl; 0.8% Triton X-100 (Sigma-Aldrich)). UV measurements and SDS-PAGE analysis of the wash and beads confirmed that the protein was bound. The beads were then incubated on a rotary with a 5 µM solution of MetHb dissolved in 10 ml of Buffer B. The incubation time was varied from 0 to 300 minutes. The beads were then centrifuged and the absorbance of the supernatant at 411 nm was measured. For transfer experiments that included
IsdH^{N1}, MetHb was first incubated with three molar equivalents of IsdH^{N1} for 90 min. The program SigmaPlot 2000 was used to fit the transfer data to the equation \( A = (A_f - A_0)(1 - e^{-k_{obs}}) + A_0 \), where \( A_0 \) is the initial absorbance of the MetHb solution prior to incubating with IsdH^{N3}, and \( A_f \) is the final absorbance of MetHb in the eluate.

Under the conditions we have used \( k_{obs} \) is approximately equal to the rate of heme release from the \( \beta \) subunit. This is because at the end of the assay ~50% of the MetHb dimers contain a single heme molecule and previous studies have shown that the \( \beta \) chain within MetHb binds heme less tightly that the \( \alpha \) chain (32). Under these conditions the heme transfer reaction is therefore described by the following equilibrium equation:

\[
MH_2 \xrightleftharpoons[k_{-H}][k_H] MH + H \xrightarrow[k_N][k_{-N}] N \rightleftharpoons NH
\]

Eq. 1

, where \( MH_2 \) is MetHb containing two heme molecules; \( MH \) is MetHb with a single heme molecule bound to its \( \alpha \) chain; \( N \), is the IsdH^{N3} NEAT domain; \( H \), is free heme, and \( NH \) is IsdH^{N3} bound to a single heme molecule. As described by Hargrove et al., a simple expression for the rate of change of \( MH_2 \) can be derived by assuming that \( d[H]/dt \sim 0 \) and that the concentration of free heme is much smaller than the concentration of heme bound to protein (i.e. \([H] \ll [MH_2] + [MH] + [NH] \sim [H]_{total}\)) (32). When these conditions are met the observed rate constant shows the following dependence:

\[
k_{obs} = \frac{k_{H} [MH]}{k_{N} [N] + k_{H} [MH]} \frac{1}{1 + \left( \frac{k_{H} [MH]}{k_{N} [N]} \right)}
\]

Eq. 2

This equation indicates that the initial kinetics of transfer are described by a single exponential where the observed rate constant is equal to the rate at which heme is released from the \( \beta \) subunit of MetHb (\( k_{obs} \sim k_H \)) because \([MH]/[N] \sim 0\). In principle, more complex binding behavior will be observed in which the data is biphasic. For example, as the ratio of \([MH]/[N]\) increased, the reaction could accelerate if \( k_{N} > k_H \), decelerate if \( k_{N} < k_H \), or remain the same if \( k_{N} = k_H \). However, these effects are expected to be small and difficult to detect experimentally.

**RESULTS**

The same site on IsdH^{N1} binds to MetHb and the structurally unrelated Hp protein. Previous studies have shown that the IsdH^{N1} and IsdH^{N2} domains within IsdH bind methemoglobin (MetHb) and haptoglobin (Hp) (19,23,25). Despite the determination of the 3D structure of IsdH^{N1}, the surface that mediates binding to Hp and MetHb remains unknown because samples of the IsdH^{N1}:Hb and IsdH^{N1}:Hp complexes have thus far proved unsatisfactory for chemical shift mapping by NMR (23,25). Therefore, we used alanine scanning mutagenesis combined with surface plasmon resonance (SPR) to probe the surface of IsdH^{N1} that binds MetHb and Hp. A total of 22 single amino acid mutants were constructed that alter surface exposed side chains in IsdH^{N1}. The majority of amino acid substitutions are located in the loops that connect strands \( \beta_1b \) to \( \beta_2 \) (\( \beta_1-\beta_2 \) loop), \( \beta_3-\beta_4 \) (\( \beta_3-\beta_4 \) loop) and \( \beta_7 \) to \( \beta_8 \) (\( \beta_7-\beta_8 \) loop) (Fig. 5A). The rationale behind altering these sites is based on the solution structure of IsdH^{N1} and sequence alignments of IsdH^{N1} with other NEAT domains known to bind MetHb (described in the Discussion section). Each single amino acid mutant contained a histidine tag to facilitate purification and was shown to be properly folded by 1D NMR (data not shown).

Fig. 2A shows representative SPR data of wild-type IsdH^{N1} binding to MetHb adhered to a CM5 chip. This data was interpreted quantitatively by plotting the maximal SPR response units (RU) obtained at varying concentrations of IsdH^{N1} (Fig. 2B). Curve fitting indicates that wild-type histidine IsdH^{N1} binds MetHb with a \( K_D \) of \( 17 \pm 10 \) nM. Importantly, the presence of the histidine tag does not affect protein binding as similar affinities for MetHb have been measured for GST-tagged
(23) and untagged variants of IsdH\textsuperscript{N1} (data not shown). Binding affinities for the mutants were obtained in a similar manner and are listed in Table 1. The most significant changes in affinity occur when residues within the β1-β2 loop are altered, since mutation of five of the amino acids in this loop result in a greater than 40 fold reduction in affinity. Alanine substitutions in adjacent loops also cause affinity reductions, albeit to a lesser extent. An interpretation of this data is presented in the Discussion section.

The surface mutants of IsdH\textsuperscript{N1} were also tested for their ability to bind to immobilized haptoglobin (Hp) using SPR. As shown in Fig. 2C, the SPR data acquired for Hp contains more noise than the MetHb data (~0.5 and ~3 RU for Hp and MetHb, respectively). The increase in noise is likely caused by heterogeneity in the commercially acquired Hp protein, which consists of distinct allelic forms that differ in their oligomerization state. However, the binding affinity of IsdH\textsuperscript{N1} for Hp could still be measured quantitatively from plots of the maximal signal intensity of each sensogram as a function of IsdH\textsuperscript{N1} concentration (Fig 2D). The results of this work are reported in Table 1 and reveal that the wild-type protein binds Hp with a K\textsubscript{D} of ~35 ± 10 nM. Alanine substitution of Y126, H127, F128 and F129 within the β1-β2 loop completely abrogates binding to Hp, while a Y125A mutation causes a 21 fold reduction in affinity. All other substitutions had a negligible effect on Hp binding. These same five residues within the β1-β2 loop are also important for MetHb binding, indicating Hp and MetHb bind to the same site on IsdH\textsuperscript{N1}.

IsdH contains a third NEAT domain that selectively binds heme. Inspection of IsdH’s amino-acid sequence suggests that it contains a third NEAT domain at its C-terminus (IsdH\textsuperscript{N3}, residues T540 to Q664) (Fig. 1A). This putative domain has not been characterized previously and shares only ~15% sequence identity with the IsdH\textsuperscript{N1} and IsdH\textsuperscript{N2} domains. Moreover, it shares limited sequence homology with all other NEAT domains of known function (23 and 15% sequence identity with IsdA and IsdC, respectively, Fig. 1C). To investigate its function, a histidine tagged variant of IsdH\textsuperscript{N3} was overexpressed and purified (his-IsdH\textsuperscript{N3}, residues T540 to Q664 containing a 21 residue histidine tag at its N-terminus). Centrifuged cell pellets overexpressing his-IsdH\textsuperscript{N3} were colored red, a phenomena characteristic of many hemoproteins (data not shown). Heme binding was further substantiated by the UV-visible spectrum of purified his-IsdH\textsuperscript{N3}, which exhibits a Soret band at 402 nm along with additional Q-band peaks at 511, 548, and 625 nm (Fig. 3A). The amount of heme bound to his-IsdH\textsuperscript{N3} was quantified using a pyridine hemochrome assay, and revealed that after purification from \textit{E. coli}, ~2-3 % of the protein contains heme (assuming 1:1 binding of heme to protein). NMR spectra of \textsuperscript{15}N labeled IsdH\textsuperscript{N3} indicate that it is folded and that it binds heme, but are of insufficient quality for detailed structural studies using this technique.

The affinity of his-IsdH\textsuperscript{N3} for metalloprotoporphyrins was measured using two techniques. Initially, binding of his-IsdH\textsuperscript{N3} to heme (iron-substituted protoporphyrin IX in its ferric form) was measured by monitoring changes in the Soret band of the protein-heme complex after the addition of varying amounts of hemin (data not shown). Analysis of this data yielded an approximate dissociation constant (K\textsubscript{D}) of ~1-20 µM, however, a more quantitative analysis could not be performed because heme aggregates at higher concentrations needed to saturate this protein. Therefore, a fluorescence-based assay was used to monitor the binding of his-IsdH\textsuperscript{N3} to a close analog of hemin that replaces the iron atom with zinc (ZnPPiX, zinc-substituted protoporphyrin IX). In this assay the concentration of ZnPPiX is held constant at 7.5 µM and changes in its fluorescence spectrum are monitored after adding varying amounts of his-IsdH\textsuperscript{N3}. As shown in Fig. 3B, the addition of protein caused a concentration dependent increase in the fluorescence of ZnPPiX that saturates when 10 µM of his-IsdH\textsuperscript{N3} is present. A plot of the fluorescence change as a function of the protein:ligand ratio revealed a binding stoichiometry of 1:1 (data not shown), while curve fitting of the data in Fig. 3B yielded a K\textsubscript{D} of 2.8 ± 0.1 µM. Similar experiments performed using an untagged variant of IsdH\textsuperscript{N3} yielded a K\textsubscript{D} of 2.3 ± 0.4 µM, indicating that the presence of the histidine tag does not affect metalloprotoporphyrin binding.
We also performed an enzyme linked immunoabsorbent assay (ELISA) to determine whether IsdH\textsuperscript{N3} was capable of binding to proteins previously demonstrated to interact with other NEAT domains. These included: holo-Transferrin, apo-Transferrin, fetuin, Fibronectin, Fibrinogen, Hp and MetHb (19,22). When probed by ELISA, IsdH\textsuperscript{N3} did not interact with MetHb (Fig. 3C, closed squares), nor did it bind to any of the other proteins tested (data not shown). This is in marked contrast to IsdH\textsuperscript{N1}, which as expected, binds MetHb when tested by ELISA (Fig. 3C, closed diamonds). Thus far, we have only shown that IsdH\textsuperscript{N3} binds heme.

**IsdH\textsuperscript{N3} and IsdC capture heme from MetHb.** Since the IsdH\textsuperscript{N1} and IsdH\textsuperscript{N2} NEAT domains within IsdH binds to MetHb, we wondered whether the IsdH\textsuperscript{N3} domain could capture heme from MetHb. To measure heme transfer, his-IsdH\textsuperscript{N3} was adhered to a cobalt resin (Talon beads, Clontech) and then incubated with MetHb (6:1 molar ratio of his-IsdH\textsuperscript{N3} to MetHb). The two proteins were then separated by washing the resin to elute MetHb. Maximum absorbance of the eluate at 411 nm is diagnostic for the presence of MetHb bound to heme, since free heme does not absorb maximally at this wavelength, and IsdH\textsuperscript{N3} and the IsdH\textsuperscript{N3}-heme complex should remain bound to the beads. When a solution of heme saturated MetHb is incubated with only the cobalt resin, ~96% of the heme bound MetHb is recovered in the eluate (Fig. 4A). In contrast, incubation of MetHb with apo-IsdH\textsuperscript{N3} for 60 minutes significantly reduces the amount of MetHb-bound heme that is obtained in the eluate when the resin is washed (Fig. 4A). Two control experiments verify that heme loss from MetHb is a result of transfer to IsdH\textsuperscript{N3}. First, SDS-PAGE analysis of the eluate indicates that similar levels of MetHb are recovered in either the presence or absence of IsdH\textsuperscript{N3} on the cobalt resin (data not shown). This indicates that the proteins do not interact with each another, consistent with the ELISA data (Fig. 3C). Second, when his-IsdH\textsuperscript{N3} is eluted from the resin with imidazole after incubation with MetHb the UV spectrum of the eluate indicates that IsdH\textsuperscript{N3} has acquired heme (data not shown). Taken together these results indicate that the IsdH\textsuperscript{N3} domain is capable of acquiring heme from MetHb and that heme transfer to IsdH\textsuperscript{N3} does not require stable protein-protein interactions.

The rate of heme transfer from MetHb to IsdH\textsuperscript{N3} was determined by varying the amount of time the proteins were incubated with one another before separation (Fig. 4B, open circles). The transfer data can be adequately fit to the single exponential equation \( A = (A_f - A_0)(1-e^{-k_{obs}t}) + A_0 \), where \( A_0 \) is the initial absorbance of the MetHb solution prior to incubating with IsdH\textsuperscript{N3}, and \( A_f \) is the final absorbance of MetHb in the eluate. This yields a rate constant \( (k_{obs}) \), which, under the experimental conditions used, describes the rate at which a single heme molecule is released from the \( \beta \) subunit of the MetHb heterodimer (detailed in the Materials and Methods section). The value of \( k_{obs} \) is 4.4 ± 0.33 hour\(^{-1}\), which is similar to the previously reported rate of spontaneous heme release from the \( \beta \) chain of MetHb (7.8 ± 2.0 hour\(^{-1}\) at pH 7.0, 37° C) (32). We next investigated whether IsdH\textsuperscript{N1} binding to MetHb accelerates the rate at which heme is transferred to IsdH\textsuperscript{N3}. The heme transfer assay was repeated where MetHb was first pre-incubated with a three-fold molar excess of IsdH\textsuperscript{N1}. At this ratio we have previously shown that a IsdH\textsuperscript{N1}:MetHb complex forms that has a stoichiometry of 2:1 (25). Thus, in this experiment the rate of transfer from the IsdH\textsuperscript{N1}:MetHb complex to his-IsdH\textsuperscript{N3} is measured. As shown in Fig. 4B (closed circles), IsdH\textsuperscript{N1} binding to MetHb increases the rate of hemin transfer to his-IsdH\textsuperscript{N3} by ~2-fold (\( k_{obs} = 10.9 ± 0.65 \) hour\(^{-1}\)).

To determine whether the IsdC NEAT domain also acquires heme from MetHb, the transfer assay was repeated using histidine-tagged IsdC (his-IsdC) as the receptor attached to talon beads instead of his-IsdH\textsuperscript{N3} (Fig. 4C, open squares). Again, the transfer data can be adequately fit to a single exponential equation with a rate constant of transfer equal to 3.8 ± 0.42 hour\(^{-1}\). Intriguingly, the rate of heme capture is similar to that observed for IsdH\textsuperscript{N3}, and is compatible with the rate at which heme is released from the \( \beta \) subunit of MetHb (vide supra). Similar experiments were performed after pre-incubating MetHb with IsdH\textsuperscript{N1}. The rate of hemin transfer from the MetHb-IsdH\textsuperscript{N1} complex to his-IsdC was 10.0 ± 0.84 hour\(^{-1}\). Similar to the case for capture by IsdH\textsuperscript{N3}, this rate is ~2-3-fold faster than from MetHb.
DISCUSSION

To cause disease, *Staphylococcus aureus* needs to acquire iron from its human host. Heme is the preferred source of iron and is harvested from hemoglobin using iron-regulated surface determinant (Isd) proteins (5-7,9). One of the first steps in this process is the capture of Hb or the Hb-Hp complex on the cell surface by the IsdH and IsdB proteins (6,19). These bacterial proteins are closely related to one another, yet structurally distinct from microbial Hb and Hp receptors found in Gram-negative bacteria (23,25). Thus far, only IsdH has been studied in detail *in vitro* (19,23,25). Analysis of its primary sequence reveals that it contains three NEAT domains: IsdH N1, IsdH N2 and IsdH N3 (Fig. 1A). Previously, the structure of IsdH N1 has been determined and the isolated IsdH N1 and IsdH N2 domains have been shown to bind Hb and Hp (19,23,25). However, the mechanism of Hb or Hp binding is not known because the structure of IsdH N1 was solved in isolation.

Since IsdH N1 and IsdH N2 are closely related to one another we hypothesized that they would use a common set of surface exposed side chains to mediate Hb and Hp binding. To identify these residues we compared the primary sequences of NEAT domains whose ligand specificities had been determined experimentally: IsdH N1, IsdH N2 and the NEAT domains from the IsdA and IsdC proteins, which do not bind Hb with high affinity (22,25). This comparison revealed the presence of an aromatic motif within IsdH N1 and IsdH N2 (Fig. 1B, boxed residues). A related motif was also found in the N-terminal domain of IsdB (IsdB N1). In IsdH N1, the motif consists of residues Y125-Y126-H127-F128-F129, which in the NMR structure are located within a loop that connects strands β1b to β2 (β1-β2 loop) (Fig. 5A). In conjunction with the surface loops that connect strands β3 to β4 (β3-β4 loop) and strands β7 to β8 (β7-β8 loop), it forms a large surface located at the end of the beta barrel structure. Intriguingly, the backbone atoms of many of the residues within this area of the protein are undetectable in the NMR data of IsdH N1, presumably because they undergo slow time scale motional fluctuations that are characteristic of ligand binding sites (23,25).

Our results indicate that residues within the aromatic motif and the surrounding surface loops are important for binding MetHb (the oxidized dimeric form of Hb that is prevalent in blood serum after erythrocyte cell lysis) (Fig. 5B). The most important binding surface is formed by the exposed side chains of Y125, Y126 and H127 within the aromatic motif. As compared to the wild-type protein, which binds MetHb with a KD of ~17 ± 10 nM, mutation of these residues causes more than a 100-fold decrease in binding affinity (Fig. 5B, in red). However, not all residues in the β1-β2 loop that houses the aromatic motif are important as amino acids immediately before the motif in the primary sequence can be altered without affecting binding (tan surface near the β1-β2 loop in Fig. 5B, N120A, T123A and Q124A mutations). In the structure, the β3-β4 and β7-β8 loops are in close proximity to the aromatic motif (Fig. 5A). Surface exposed residues within these sites also contribute to MetHb binding, although to a lesser extent. The β3-β4 loop appears to be more important as mutations in it reduce binding ~10 fold (Fig. 5B, colored dark purple) as compared to 3-4 fold reductions when residues within the β7-β8 loop are altered (Fig. 5B, colored pink). Since mutations that alter surface exposed side chains elsewhere in the protein have little effect on binding, we conclude that the primary interface for MetHb binding is located at the end of the β-barrel structure. IsdH N1 and MetHb form a complex that has a stoichiometry of 2:1 (25). This suggests that IsdH N1 interacts with both the α and β subunits of MetHb. Residues within the aromatic motif may adaptively recognize the distinct binding surfaces of these subunits, since NMR studies suggest that residues in the motif are mobile in the apo-state (26-28).

IsdH is also the sole haptoglobin (Hp) receptor in *S. aureus*, since isdh strains are unable to bind Hp (19). Because Hp binds MetHb with a KD ~ 10⁻¹⁵ M, IsdH binding to Hp may enable *S. aureus* to coat itself with this high affinity MetHb receptor. It is also possible that Hp coating of the cell surface enables the bacterium to disguise itself immunologically to prevent targeting by the host immune response (23). Our results indicate that IsdH N1 binds Hp with a KD of 35 ± 10 nM. This affinity is higher than previously reported values determined by SPR (23). Presumably this
difference is caused by the way we performed the binding assay. In our assay, Hp is conjugated to a CM5 chip and binding is measured by passing different concentrations of IsdH\textsuperscript{N1} over it. In contrast, previous studies have used the reverse approach; IsdH\textsuperscript{N1} was adhered to the chip and Hp was passed over it (23). With this prior approach a biphasic association curve was observed that contained both a fast and slow association phase. The slower phase describes a weaker binding event with a measured \( K_D \) of \(~5\ \mu\text{M}, \) while the faster phase characterizing a stronger interaction was not quantified. As our sensograms have single phase behavior they enable us to determine the dissociation constant for the high affinity interaction, and reveal that IsdH\textsuperscript{N1} is capable of binding Hp and MetHb with similar nanomolar affinities (35 nM and 17 nM respectively).

Interestingly, residues in the aromatic motif required for Hb binding also play an essential role in Hp recognition (Table 1 and Fig. 5C). The most important residues are Y126, H127, F128 and F129, which when mutated completely disrupt binding. Surprisingly, amino acid mutations in the surrounding \( \beta_{3-4} \) and \( \beta_{7-8} \) loops, as well as in other regions of IsdH\textsuperscript{N1}, have no detectable effect on Hp binding (Fig 5C, colored tan). This suggests that the interface between IsdH\textsuperscript{N1} and Hp is largely restricted to residues in the aromatic motif, in contrast to the MetHb binding surface, which also includes residues located in the surrounding loops (compare Figs. 5B and 5C). Interestingly, the residues in the aromatic motif of IsdH\textsuperscript{N2} and IsdB\textsuperscript{N2} are identical (see Fig. 1B, boxed sequence). However, only IsdH\textsuperscript{N2} binds Hp, whereas the full-length IsdB protein does not (19,23,25). This argues against the \( \beta_1-\beta_2 \) loop being the sole binding surface for Hp. It is possible that more drastic non-conservative amino acid substitutions are required to identify residues important for Hp that are located outside of the aromatic motif.

Our finding that the same surface on IsdH\textsuperscript{N1} contacts MetHb and Hp is surprising considering that these proteins are structurally unrelated to one another. Although their names imply that they both adopt a globin fold, only the \( \alpha \) and \( \beta \) subunits within the MetHb heterodimer possess a globin structure. In contrast, the structure of Hp is not known, however, based on primary sequence homology, its \( \beta \)-chain may adopt a fold related to the serine proteases (33). Previous modeling studies of Hp have suggested that it contains two hydrophobic surfaces that may enable it to function as a protein chaperone (34). To further investigate how IsdH\textsuperscript{N1} might interact with MetHb and Hp we used the program ClusPro to model the structures of the IsdH\textsuperscript{N1}:MetHb and IsdH\textsuperscript{N1}:Hp complexes (35). The docking calculations made use of the NMR structure of IsdH\textsuperscript{N1}, the crystal structure of carboxyhemoglobin (pdb 1IRD), and a model of the structure of Hp generated using the structure of the human complement C1S protease as a template (pdb 1ELV, which shares 33\% sequence identity with residues 91 to 406 of Hp) (36,37). In both sets of docking calculations, the predominant solution obtained positioned the aromatic motif within IsdH\textsuperscript{N1} at the protein-protein interface. In addition, this structural element tended to be inserted between the \( \alpha \) and \( \beta \) domains of Hb, and the protease-like and complement-like modules in Hp (data not shown). However, a molecular level understanding of recognition will require the determination of the three dimensional structures of the IsdH\textsuperscript{N1}:MetHb and IsdH\textsuperscript{N1}:Hp complexes by either NMR spectroscopy or x-ray crystallography.

Our results indicate that IsdH contains a third, previously uncharacterized heme binding NEAT domain at its C-terminus (IsdH\textsuperscript{N3}) (Fig. 3C). Its mode of heme binding can be predicted from the recently determined structures of the metalloprotoporphyrin complexes of the IsdA and IsdC NEAT domains (26-28). Although IsdH\textsuperscript{N3} shares only 15-23\% sequence identity with IsdA and IsdC, residues that form the heme binding pocket in these proteins are generally conserved (residues underlined in Fig. 1C). Of particular interest is the presence of an invariant tyrosine residue that coordinates the central metal ion of the heme; Tyr\textsuperscript{166} (IsdA), Tyr\textsuperscript{132} (IsdC) and Tyr\textsuperscript{642} (IsdH\textsuperscript{N3}) (26-28). Assuming that IsdH\textsuperscript{N3} adopts a conventional NEAT domain fold, it seems likely that it will bind heme in a similar manner through an exposed pocket located at the end of a \( \beta \)-barrel structure. The walls of the heme binding pocket would be formed by residues within strands \( \beta_7 \) and \( \beta_8 \), as well as residues within the loops that connect strands \( \beta_1 \) to \( \beta_2 \) (\( \beta_1-\beta_2 \) loop) and strands \( \beta_3 \) to \( \beta_4 \) (\( \beta_3-\beta_4 \) loop). Our data indicates that IsdH\textsuperscript{N3} binds metalloprotoporphyrins with a
The *IsdC* proteins from *S. aureus* and *B. anthracis* also bind heme weakly (12,28). This is compatible with *IsdH* and *IsdC*'s physiological role in heme transport, since they presumably need to rapidly bind and release heme as it is transferred across the cell wall (12,28). The weak affinity of these proteins is also consistent with the recently determined structures of *IsdA* and *IsdC*, since in them only ~35% of the solvent accessible surface area of the protoporphrin ring is contacted by the protein (26-28).

An overview of how the NEAT domains within *IsdH* might function in heme capture is shown in Fig. 6. Upon erythrocyte lysis large quantities of tetrameric ferrous Hb is released into the blood, which rapidly oxidizes and dissociates into dimeric MetHb (14). The *IsdH*\(^{N1}\) and *IsdH*\(^{N2}\) domains bind MetHb with nanomolar affinity tethering it to the cell surface (step 1). Heme is then released and captured by the adjacent *IsdH*\(^{N3}\) domain (step 2). This process is passive as the measured rate of transfer from MetHb to *IsdH*\(^{N3}\) is ~4.4 per hour (Fig. 4B), which is similar to the rate at which heme spontaneous dissociates from MetHb (12,28). Heme bound to *IsdH*\(^{N3}\) then presumably dissociates where it can be either recaptured by MetHb or transferred to either *IsdA* or *IsdC* within the cell wall (step 3). It is also conceivable that heme is directly transferred from *IsdH*\(^{N3}\) to *IsdA* and/or *IsdC* through activated protein-protein complexes, since this mechanism of transfer has been shown to occur between the *IsdA* and *IsdC* proteins (38). In addition, our results indicate that *IsdC* can also directly capture heme that is released from MetHb (Fig. 4C), however this process would seem to occur less frequently as *IsdC* is presumably positioned more distal to MetHb than *IsdH*\(^{N3}\). This model of heme transfer is compatible with recent studies by the Lei group, which have shown that the *IsdA* and *IsdC* proteins can acquire heme that is released from MetHb (39). The *IsdB* protein also functions as a MetHb and heme receptor in *S. aureus* (19,23,25). As shown in Fig. 1A, it contains two NEAT domains, *IsdB*\(^{N1}\) and *IsdB*\(^{N2}\). The *IsdB*\(^{N1}\) domain is most closely related to the *IsdH*\(^{N1}\) and *IsdH*\(^{N2}\) domains. In contrast, the C-terminal *IsdB*\(^{N2}\) domain is most closely related to *IsdH*\(^{N3}\) and it contains conserved residues that presumably enable it to bind heme (Fig. 1B). This suggests *IsdB* and *IsdH* capture heme from MetHb through a similar mechanism; MetHb is bound by *IsdB*\(^{N1}\) and released heme is scavenged by the *IsdB*\(^{N2}\) domain. Interestingly, Lei and colleagues have shown that heme transfer from MetHb to the full-length *IsdB* protein occurs at rates that are much faster than the rate of spontaneous heme release from MetHb (19,23,25). This suggests that MetHb binding to the full-length *IsdB* protein stimulates heme release. Our finding that binding of the isolated *IsdH*\(^{N1}\) domain to MetHb only modestly increases the rate of heme transfer to *IsdH*\(^{N3}\) may explain why *IsdB* has been proposed to play a more important role than *IsdH* in the *in vivo* capture of heme from hemoglobin (19,23,25). Alternatively, it suggests that the NEAT domains within *IsdH* and *IsdB* need to be part of the same polypeptide, that is, act in cis, in order to effectively capture heme. Unfortunately, we have not been able to test the degree of synergy between the NEAT domains within *IsdH* because the full-length protein degrades during purification.

In conclusion, we have shown that *IsdH* captures heme from MetHb using three NEAT domains that have different ligand binding specificities. The *IsdH*\(^{N1}\) and *IsdH*\(^{N2}\) bind MetHb and Hp, while a third C-terminal *IsdH*\(^{N3}\) domain captures released heme. Analogous NEAT domains are present in *IsdB* based on primary sequence homology, suggesting that it will capture heme from MetHb through a similar mechanism. We have also shown that *IsdH*\(^{N1}\) employs a common surface located at the end of its beta structure to interact with the structurally distinct Hp and MetHb proteins. Interestingly, structural studies of the *IsdA* and *IsdC* proteins indicate that they use the same site bind heme. This suggests that this part of the protein scaffold is acquiescent to change that ultimately determines the function of the NEAT domain.

ACKNOWLEDGEMENTS

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FIGURE LEGENDS

Figure 1. Comparison of the NEAT domains in S. aureus. (A) Schematic of the IsdB and IsdH proteins showing that they contain two and three Near Transporter (NEAT) domains, respectively. Domains are shaded based on their relatedness. Domains shaded grey share 46-65% primary sequence identity (IsdH N1, IsdH N2 and IsdB N1). Non-shaded domains share 49% primary sequence identity (IsdH N3 and IsdB N2). Based on studies of the isolated NEAT domains within IsdH, the grey- and non-shaded domains bind Hb and heme, respectively. Both proteins also contain a cell wall sorting signal motif at their C-termini which is covalently attached to the cell wall by the SrtA sortase enzyme. (B) Sequence alignment of related Hb binding NEAT domains within IsdH and IsdB: IsdH N1, IsdH N2 and IsdB N1. These domains are closely related to one another and share 46-65% sequence identity. Residues mutated in this study are indicated with an asterisk and amino acids are enclosed in a box if their mutation to alanine completely disrupts Hb binding and/or reduces the affinity of IsdH N1 for Hb by at least 50x fold. The aromatic motif diagnostic of NEAT domains that bind Hb is underlined. The secondary structure of IsdH N1 is shown above the primary sequence. Completely conserved residues are in bold. Only the IsdH N1 and IsdB N2 domains have thus far been shown to bind Hb, while the IsdB N1 domain alone has yet to be tested. (C) Sequence alignment of the IsdH N3 and IsdB N2 NEAT domains, which share 49% sequence identity. The sequences of the distantly related IsdA and IsdC domains are also shown because they bind to heme. Positions within the primary sequence that are within 3.5 Å of the heme in the NMR and crystal structures of the IsdC-heme and IsdA-heme complexes are underlined. The invariant tyrosine residues that coordinate the iron atom of the heme in these structures is enclosed in a box. Residues that are completely conserved in IsdH N3 and IsdB N3 are in bold.

Figure 2. Surface plasmon resonance (SPR) measurements of IsdH N1 binding to MetHb and Hp. (A) Representative SPR data of wild-type IsdH N1 binding to immobilized MetHb. The panel shows an overlay of several experiments in which the concentration of IsdH N1 was varied (0, 100, 200, 300, 400, 500, 1000, 2000 and 4000 nM). Each curve is a plot of the response units (RU) as a function of time. (B) Representative binding curve of the wild-type IsdH N1 for immobilized MetHb. The curve was generated from the data in panel (A) by plotting the RU value measured one second before dissociation for each concentration of IsdH N1. The solid line shows the best fit of this data to obtain the K D of binding. (C) Representative SPR data of wild-type IsdH N1 binding to immobilized Hp. The results of six independent experiments are superimposed and differ in the concentration of IsdH N1 (0, 100, 200, 300, 400, 500, and 1000 nM IsdH N1). (D) Representative binding curve of wild-type IsdH N1 for immobilized Hp by plotting the RU value measured one second before dissociation for each concentration of IsdH N1.

Figure 3. The IsdH N3 NEAT domain binds heme. (A) UV spectrum of purified IsdH N3 that show characteristics of a heme-binding protein. These include a Soret band at 402 nm along with additional Q-band peaks at 511, 548, and 625 nm. (B) Quantitative measurement of protoporphyrin binding using a fluorescence assay. The panel shows a plot of zinc protoporphyrin (ZnPPIX) fluorescence emission at 585nm as a function of IsdH N3 added. The solid line shows the best fit of this data which yields a K D of ZnPPIX binding to IsdH N3 of 2.8 ± 0.1 µM. (C) An ELISA assay showing that the IsdH N1 NEAT domain binds MetHb, while the IsdH N3 does not. IsdH N3 binding to a range of other proteins was also tested, but no interaction could be detected.

Figure 4. Transfer of heme from MetHb to IsdH N3 and IsdC (A) Bar graph showing the percentage of heme recovered after MetHb is incubation with talon beads for 120 minutes. The negative control indicates that the talon beads were naked. IsdH N3 and IsdC indicate that the talon beads used in this experiment were pre-incubated with the his-tagged versions of these proteins. (B) Quantitative measurement of the rate of heme transfer from MetHb to IsdH N3. The open circles represent heme transfer
from MetHb to IsdH\textsuperscript{N3} as a function of time. The closed circles indicate transfer from MetHb when it is bound to IsdH\textsuperscript{N1} in the IsdH\textsuperscript{N1}.MetHb complex. The solid line is the best fit of this data and was used to extract the rate of transfer (described in text). (C) Quantitative measurement of heme transfer from MetHb to IsdC. As described in panel (B), except that his-IsdC protein was used as the receptor for heme. The open and closed squares represent heme transfer from MetHb and the IsdH\textsuperscript{N1}.MetHb complex, respectively.

**Figure 5.** A common surface on the IsdH\textsuperscript{N1} NEAT domain binds haptoglobin and methemoglobin. (A) Ribbon drawing of the solution structure of IsdH\textsuperscript{N1}. Strands of $\beta$ sheet are colored grey and indicated by arrows. The three loops involved in protein binding are indicated. (B) Solvent exposed surface of IsdH\textsuperscript{N1} color coded to show the effects of amino acid mutations on MetHb binding. Color coding key: Red, $>50x$ reduction or no detectable binding; dark purple, 10-11x reduction; pink, 2-4x reduction; tan, no significant effect on binding ($<2x$ reduction). (C) Solvent exposed surface of IsdH\textsuperscript{N1} color coded to show the effects of amino acid mutations on MetHb binding. Color coded as described in panel (B). The view of the structure is identical in all of the panels.

**Figure 6.** Schematic showing how the IsdH protein may capture and transfer heme from hemoglobin. Step 1: MetHb is tethered to IsdH via the N-terminal IsdH\textsuperscript{N1} and IsdH\textsuperscript{N2} NEAT domains. MetHb is shown as two hexagons, representing the $\alpha$ and $\beta$ globin chains. Heme molecules are represented by 5-pointed stars. Step 2: Heme is released from MetHb. Our data indicates that this process is passive when NEAT domains are in isolation. Step 3: Heme diffuses and is captured by the C-terminal IsdH\textsuperscript{N3} NEAT domain. Step 4: Heme is released from IsdH\textsuperscript{N3} and subsequently captured by IsdC located within the cell wall. Our results also indicate that IsdC can directly capture heme that is passively released from MetHb (dashed line). Note, IsdH and IsdC are covalently attached to the cell wall by the SrtA and SrtB sortases, respectively.
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<td>57 ± 12</td>
<td>1.6</td>
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<td>G207A</td>
<td>32 ± 7.5</td>
<td>2</td>
<td>20 ± 7</td>
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<td>Surrounding mutants</td>
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<td>D136A</td>
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<td>K142A</td>
<td>16 ± 6.9</td>
<td>1.0</td>
<td>47 ± 16</td>
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<sup>a</sup>The wild-type protein consists of residues 86 to 229 of the IsdH protein connected at its amino terminus to a nineteen residue histidine tag. Each mutant is identical to the wild-type protein, but contains a single alanine residue substitution at the indicated position.

<sup>b</sup>Dissociation constants were measured twice using surface plasmon resonance (SPR) on a Biacore system. The reported errors are the average fits from the average dissociation constant.

<sup>c</sup>The fold decrease is the affinity of the mutant that is compared to the wild-type protein affinity for its substrate.
**Hemoglobin binding NEAT domains**

**Heme binding NEAT domains**

Fig. 1
Fig. 2
Fig. 4
Fig. 5
Fig. 6
References


Functionally distinct NEAT (NEAr transporter) domains within the staphylococcus aureus IsdH/HarA protein extract heme from methemoglobin

Rosemarie M. Pilpa, Scott A. Robson, Valerie A. Villareal, Melissa L. Wong, Martin Phillips and Robert T. Clubb

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