Unique Heme-Iron Coordination by the Hemoglobin Receptor IsdB of *Staphylococcus aureus*

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**ABSTRACT:** Iron is an essential requirement for life for nearly all organisms. The human pathogen *Staphylococcus aureus* is able to acquire iron from the heme cofactor of hemoglobin (Hb) released from lysed erythrocytes. IsdB, the predominant Hb receptor of *S. aureus*, is a cell wall-anchored protein that is composed of two NEAT domains. The N-terminal NEAT domain (IsdB-N1) binds Hb, and the C-terminal NEAT domain (IsdB-N2) relays heme to IsdA for transport into the cell. Here we present the 1.45 Å resolution X-ray crystal structure of the IsdB-N2–heme complex. While the structure largely conforms to the eight-strand β-sandwich fold seen in other NEAT domains such as IsdA-N and uses a conserved Tyr residue to coordinate heme-iron, a Met residue is also involved in iron coordination, resulting in a novel Tyr-Met hexacoordinate heme-iron state. The kinetics of the transfer of heme from IsdB-N2 to IsdA-N can be modeled as a two-step process. The rate of transfer of heme between the isolated NEAT domains (82 s⁻¹) was found to be similar to that measured for the full-length proteins. Replacing the iron coordinating Met with Leu did not abrogate high-affinity heme binding but did reduce the heme transfer rate constant by more than half. This unusual Met-Tyr heme coordination may also bestow properties on IsdB that help it to bind heme in different oxidation states or extract heme from hemoglobin.
Isd proteins have been further demonstrated to mediate either heme binding\(^{10-16}\) or hemoprotein binding.\(^{14,17,18}\)

IsdB contains two NEAT domains\(^9\) with 12% pairwise sequence identity and acts as the primary Hb receptor for the cell.\(^{19}\) Deletion of isdB virtually abolishes Hb binding by the \(S.\) aureus cell and significantly impacts disease in a mouse infection model.\(^{19}\) IsdB is also highly immunogenic during infection and is the basis for a monovalent vaccine currently undergoing clinical trials.\(^{20}\) IsdB extracts the heme from Hb at the cell surface for the transfer to IsdA or IsdC, which then transfers it to the membrane transporter for internalization.\(^{21-23}\) Recent evidence strongly implies that each NEAT domain of IsdB has a different binding specificity; note that NEAT domain numbering proceeds from the N-terminus to the C-terminus (Figure 1). IsdB-N1 binds Hb but not heme.\(^{22}\) Conversely, IsdB-N2 binds heme\(^{15,16}\) but not Hb,\(^{22}\) and its sequence contains a conserved Tyr residue that is known to coordinate the heme-ion in all other heme binding NEAT domains.\(^{10-12}\) IsdB-N2 alone is able to transfer heme to IsdA-N1 and IsdC-N1;\(^{15}\) however, full-length IsdB transfers heme to full-length IsdA at 8 times the rate for full-length IsdC,\(^{15}\) and protease digestion suggests that IsdB may be physically closer to IsdA than IsdC on the cellular surface, suggesting the physiological route of heme transfer is likely from IsdB-N2 to IsdC-N1 via IsdA-N1.

We sought to characterize heme binding by IsdB-N2 as a first step in understanding Hb reception and heme transfer by the Isd system. We determined the 1.45 Å resolution crystal structure of heme-bound IsdB-N2 and discovered a new mode of heme-ion coordination, with protein ligands provided by both a Met and a Tyr. We further showed that iron coordination by the Met residue is dispensable for high-affinity heme binding but has a role in facilitating the transfer of heme to IsdA-N1.

## Experimental Procedures

### Cloning of NEAT Domains and Variants

The coding region of the second, heme binding NEAT domain (IsdB-N2, Lys341—Thr459) was amplified from \(S.\) aureus N315 chromosomal DNA and cloned into the pET28a(+) expression vector for expression with an N-terminal His\(_6\) tag and thrombin cleavage site. Site-directed variants were created by subcloning from the IsdB-N2 clone, using a modified whole plasmid polyacrylamide gel electrophoresis (PAGE) apoprotein was dialyzed against either 20 mM Tris (pH 8.0) for crystallization or 50 mM Tris (pH 8.0) and 100 mM NaCl for spectroscopic and kinetic studies.

Selengomethionine-labeled (Se-Met) IsdB-N2 was prepared by the method previously described by Van Duyne et al.\(^{25}\) and purified as described for native protein.

### Heme Reconstitution

Purified apoprotein was incubated for 1 h at 4 °C with 1.5 molar equiv of hemin dissolved in 0.1 M NaOH and diluted in 0.1 M phosphate buffer (pH 7.4). Excess hemin was removed by centrifugation, and nonspecifically bound hemin was removed by gel filtration chromatography on a Sephadex G-50 column (1 cm \(\times\) 6 cm). The concentration of the holoprotein was then determined by the pyridine hemochrom assay using an \(ε_{418}\) extinction coefficient of 191.5 mM\(^{-1}\) cm\(^{-1}\) as previously described.\(^{26}\)

### Crystal Structure Determination

Native holo-IsdB-N2 and Se-Met IsdB-N2 crystals were grown by hanging drop vapor diffusion at room temperature at a 1:1 ratio of protein to well solution (composed of 0.1 M Tris (pH 8.0), 0.1 M MgCl\(_2\), and 25% polyethylene glycol 3350). Crystals were briefly washed in mother liquor and flash-frozen in liquid nitrogen. Data were collected at the Stanford Synchrotron Radiation Lightsource on beamline 9-2. Data were processed and scaled using HKL2000.\(^{27}\) Crystals grew in space group \(P2_12_12_1\) with four molecules in the asymmetric unit.

Se-Met crystals were generated initially for anomalous phasing, but in the interim, a suitable model was deposited in the Protein Data Bank (PDB) (IsdH-N3, 60% identical; PDB entry 2E7D) and molecular replacement with a single chain (with no heme) from that structure using MolRep\(^{29}\) from the CCP4 program suite\(^{29}\) yielded interpretable phases. Se-Met IsdB-N2 was built using ARP/WARP,\(^{30}\) and manual building was completed using Coot.\(^{31}\) A Ramachandran plot revealed that 92.5% of residues were in the most favored regions with the remaining 7.5% in additional allowed regions. Subsequently, native holo-IsdB-N2 crystals were generated, and MolRep was again used for phasing by molecular replacement with the Se-Met structure as the search model. The structure was modified using Coot\(^{31}\) and refined with Refmac5.\(^{22}\) Multiple conformations of side chains were modeled by visual examination of \(F_o\) – \(F_c\) maps. Waters

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**Figure 1.** Schematic of IsdB sequence features. Amino acids are numbered according to the \(S.\) aureus N315 IsdB sequence (NP_374246). The black bar near the N-terminus signifies the signal sequence, which directs the protein to be secreted onto the surface. The white bar near the C-terminus signifies the LPQQTG sortase signal, which directs the protein to be covalently anchored to the peptidoglycan by sortase A. The two NEAT domains are shown in different shades of gray to indicate their dissimilar pairwise sequence identity (12%). The mature protein on the staphylococcal surface encompasses Ala41–Thr613.
Table 1. X-ray Data Collection and Refinement Statistics

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*Values in parentheses represent those for the highest-resolution shell.

were added using the ARP/Waters$^{30}$ function in Refmac5, and model quality parameters were assessed using Procheck.$^{33}$ A Ramachandran plot revealed that 94.3% of residues were in the most favored regions with the remaining 5.7% in additional allowed regions. Figures are of the native structure and were generated using PyMOL.$^{34}$ Data collection and refinement statistics for both the native and Se-Met IsdB-N2 structures are listed in Table 1.

**Determination of the IsdB-N2 Heme Binding Stoichiometry.** Heme binding was tracked by difference absorption spectroscopy in the Soret region (around 400 nm) at room temperature. Aliquots of hemin (1.3 μL) solubilized in 0.1 M NaOH and diluted in 50 mM Tris (pH 8.0) and 100 mM NaCl to 1 mL of a 5 μM solution of apo-IsdB-N2 or a reference cuvette containing buffer alone. Spectra were recorded 10 min after addition of heme, and titrations covered a heme concentration range from 0.5 to 12 μM. Saturation was defined as a plateau in the absorption difference between the reference cuvette and the experimental cuvette. For reference, absorption spectra (250–650 nm) of purified reconstituted proteins were measured. All spectra were recorded using a Cary 50 Bio UV—visible spectrophotometer (Agilent Technologies, Mississauga, ON) with an optical path length of 1 cm in a quartz cell at room temperature.

**Trp Fluorescence Quenching by Heme.** Heme binding was monitored by fluorescence quenching of the tryptophan residue (Trp392) at the base of the heme pocket, based on the method described by Eakanunkul et al.$^{35}$ Fluorescence emission spectra from 300 to 450 nm were recorded at 20 °C with excitation at 295 nm in 50 mM Tris (pH 8) and 100 mM NaCl with an apo-IsdB-N2 concentration of 1 μM in a 1 mL volume using a Cary Eclipse fluorescence spectrophotometer (Agilent Technologies). Heme (prepared as described above, in various dilutions) was added to the buffered protein solution in 0.5−1.5 μL aliquots and allowed to reach equilibration (which ranged from 6 to 12 min depending on the variant) before readings were taken. The titrations covered a heme concentration range of 0.1−15 μM. The dissociation constant ($K_D$) was calculated from the decrease in the fluorescence intensity between 300 and 450 nm as a function of increasing heme concentration, using an equation for equilibrium binding that accounts for ligand depletion as described by Stein et al.$^{36}$ All reactions were conducted in triplicate.

**Determination of Rates of Transfer of Heme to Apomyoglobin.** The rates of dissociation of heme from IsdB-N2 were measured by single-wavelength stopped-flow spectroscopy with apomyoglobin (apoMb) as a heme scavenger.$^{37}$ ApoMb was prepared from myoglobin (Sigma-Aldrich, St. Louis, MO) as previously described.$^{38}$ Heme dissociation reactions were conducted with 10 μM holo-IsdB-N2 (reconstituted as described above) in one syringe (final concentration of 5 μM) and 100 μM apomyoglobin (final concentration of 50 μM) in the second syringe, both in 50 mM Tris (pH 8) and 100 mM NaCl at room temperature in an SX.18MV stopped-flow reaction analyzer (Applied Photophysics, Leatherhead, UK). Reactions were monitored by recording the absorbance at 408 nm for 1000 s (wild-type; ~7 half-lives) or 600 s (M362L variant; ~11 half-lives); 1000 readings at logarithmic intervals were acquired using Pro-Data SX software, regardless of the time frame. The change in absorbance at 408 nm (maximal absorbance for holomyoglobin) was plotted versus time and fit by a double-exponential equation to yield the first-order rate constants for heme dissociation using GraphPad Prism version 5.02 for Windows (GraphPad Software, La Jolla, CA). All reactions were conducted in triplicate.

**Determination of Rates of Transfer of Heme to IsaDA-N1.** Differences in spectral characteristics between holo-IsdB-N2 and holo-IsdA-N1 were exploited to monitor the transfer of heme from holo-IsdB-N2 to apo-IsdA-N1 by stopped-flow spectroscopy, conducted using an SX.18MV stopped-flow reaction analyzer (Applied Photophysics) equipped with a photodiode array detector. The reactions were conducted with 4 μM holo-IsdB-N2 in one syringe (final concentration of 2 μM) and concentrations of apo-IsdA-N1 ranging from 20 to 100 μM (final concentrations of 10−50 μM) in the second syringe; a minimal 5-fold excess of the IsaD acceptor was used to attain pseudo-first-order conditions. One hundred spectra were recorded at logarithmic intervals from 0 to 10 s from approximately 300 to 650 nm, using Xscan (Applied Photophysics). The drive syringe chamber and optical cell were maintained at 25 °C by a circulating water bath. The change in absorbance over time at the wavelength of maximal change (418.9 nm for the wild type and 425.3 nm for the M362L variant) for a given concentration was fit by a single-exponential equation to determine the observed transfer rate ($k_{obs}$). All reactions were conducted in quadruplicate.

**RESULTS**

**Crystal Structure of IsdB-N2.** The holo structure of IsdB-N2 was determined to 1.45 Å resolution. The structure of Se-Met IsdB-N2 was also determined to 1.7 Â but was inappropriate for interpretation of heme binding due to the artifact introduced by
the selenomethionine substitution, discussed in a later section. There were four molecules in the asymmetric unit that overlaid with an average root-mean-square deviation (rmsd) of 0.49 Å over all C\(^\alpha\) atoms, with the main differences occurring at the N-termini. The final model consists of residues 341–452 for chain A, 341–458 for chains B and C, and 341–456 for chain D and includes an N-terminal Gly-Ser cloning artifact left over from the thrombin cleavage site.

IsdB-N2 adopts the characteristic eight-strand immunoglobulin-like \(\beta\)-sandwich fold observed in other NEAT domains whose structures are known (Figure 2A).\(^{10-12,14,39}\) In addition to short \(\alpha\)-helices between \(\beta\)1 and \(\beta\)2 and between \(\beta\)3 and \(\beta\)4, there is a C-terminal \(\alpha\)-helix of approximately 1.5 turns immediately following \(\beta\)8 that is not observed in other NEAT domain structures.

**Structure of the Heme Binding Pocket.** Heme is bound in a highly hydrophobic pocket and is modeled at equal occupancy in two orientations when rotated by 180° along the C\(\alpha\)–C\(\gamma\) axis (Figure 2B). Approximately 330 Å\(^2\) (38%) of the heme surface area was found to be exposed to solvent as calculated using ArealMol.\(^{29}\) Crystal packing interactions occurred across the face of the heme pocket, resulting in vinyl groups of heme molecules being ~3.3 Å apart (Figure 1 of the Supporting Information). The heme propionates are extended out from the pocket, and one forms a hydrogen bond with the hydroxyl group of Tyr444 and the backbone nitrogen of Met362. Tyr440, Tyr444, and Phe366 contribute to \(\pi\)-stacking with a buried heme pyrrole ring, whereas Val431, Val433, Val446, Val435, Tyr391, Trp392, and Met363 contribute to hydrophobic contacts.

The heme iron is coordinated by Tyr440, which forms a hydrogen bond with the phenolate of Tyr444, an interaction that is absolutely conserved among heme binding NEAT domains of known structure.\(^{10-12}\) The heme-iron exists in a mixture of hexacoordinate and pentacoordinate states, with the sulfur from Met362 occupying the sixth-coordinate position. The electron density is best modeled with Met362 in both coordinating and noncoordinating conformations in the structure, and the occupancy of the conformations is unequal and dependent upon the molecule examined. In chains A and D, inspection of \(F_o - F_c\) maps indicates that Met362 exists in a coordinating position at 75% occupancy (sulfur atom 2.5–2.6 Å from heme-iron) and 25% occupancy in a noncoordinating position (sulfur atom

**Figure 2.** Crystal structure of the heme-bound C-terminal NEAT domain of IsdB. (A) Overall structure of holo-IsdB-N2 (chain A) viewed down the heme binding pocket. The backbone is shown as a cartoon with helices colored purple, \(\beta\)-strands cyan, and loops gray. Heme is shown protruding from the pocket as sticks, with carbon atoms colored dark blue and oxygen atoms red. One conformation of heme is shown for the sake of clarity. The N- and C-termini and helices are labeled. (B) Close-up of the heme pocket. Residues directly involved in binding the heme molecule (Tyr440, Met362, and Ser361) or indirectly involved in binding (Tyr444) are shown as sticks and labeled. Protein carbon, oxygen, and sulfur atoms are colored purple, red, and yellow, respectively. Heme carbon, oxygen, nitrogen, and iron atoms are colored blue, red, dark blue, and dark red, respectively. (C and D) \(F_o - F_c\) omit maps (contoured at 3\(\sigma\)) for Tyr440, Met362, and heme of chains A and B.
4.9–5.0 Å from heme-iron) (Figure 2C). The opposite is true for chains B and C, where the sulfur atom models away from the pocket at 70% occupancy (4.8 Å from heme-iron) and the sulfur atom is directed toward the heme iron at 30% occupancy (2.6 Å from heme-iron) (Figure 2D). Furthermore, these conformations appear to be complementary to one another, such that the heme pocket containing a mainly coordinating Met362 (chain A or D) is tightly packed against a heme pocket containing a mainly noncoordinating Met362 (chain B or C). Lastly, the heme-iron is pulled out of the plane of the porphyrin ring and closer to Tyr440 in chains B and C (2.1–2.2 Å Fe–O bond length, 0.4–0.5 Å from planarity), where Met362 is mainly noncoordinating, whereas the iron is pulled away from Tyr440 and lies closer to the porphyrin ring plane in chains A and D (2.2–2.3 Å Fe–O bond length, less than 0.2 Å from planarity), where Met362 is mainly coordinating.

The variability in the Met362 coordination state was first observed in the 1.7 Å Se-Met structure, motivating the determination of the native structure. Although it is well-documented that substitution of a Met heme-iron ligand for Se-Met has little effect on heme iron coordination, we were concerned that the slightly larger van der Waals radius of selenium or an incomplete substitution of Se-Met for Met might have caused the mixture of coordination states observed in the structure.

Visible Absorption Spectra of Heme Pocket Variants. On the basis of the residues that were seen to directly participate in heme binding in the IsdB-N2 crystal structure, several variants of the wild-type (WT) protein were constructed using site-directed mutagenesis: Y440A, Y444A, S361A, and M362L. The visible absorption spectra of the variants reconstituted with equimolar hemin were compared to that of the WT protein, as the spectrum is indicative of the heme environment. Mutation of any one of these residues was found to significantly affect the shape, height, and wavelength of the Soret peak, indicating a significant change in the environment experienced by the heme (Figure 3A). Furthermore, the spectra of the Y444A, Y440A, and S361A variants more closely resembled that of free heme than that of WT IsdB-N2, indicating that loss of Tyr444, Tyr440, or Ser361 resulted in severe heme binding disruption. The spectrum of M362L resembled most closely that of the wild-type (WT) protein (Figure 3B). However, the Soret peak shifted from 404 nm for WT to 398 nm for M362L, and the Soret shape and height were also significantly different, indicating different heme environments.

Heme Titration of IsdB-N2 and Variants. Although a single heme molecule is bound per monomer of IsdB-N2 in the crystal structure, investigators found that a monomer could bind up to four molecules of heme in solution. The crystal structure of IsdB-N2 also bound one heme per monomer, to confirm that 1:1 stoichiometry was recapitulated in solution apoprotein was titrated with increasing amounts of heme and difference spectra were generated by subtracting spectra from that of the buffer titrated with the same amounts of heme. WT IsdB-N2 and M362L (Figure 4A,B) both demonstrated a binding stoichiometry of approximately 1:1. However, Y444A, Y440A, and S361A difference spectra did not follow a single-site, specific binding curve and resembled that of a non-heme binding protein, transferrin, indicating aberrant or weak heme binding (data not shown). Because of the apparent weak binding for these variants, they were deemed unsuitable for performing transfer assays.

IsdB-N2 contains one Trp residue (W392), which resides at the base of the heme binding pocket. We were able to exploit this fact to develop a fluorescence-based measurement of heme binding as heme addition causes quenching of Trp fluorescence. We observed a concentration-dependent and saturable quenching of Trp fluorescence by both WT IsdB-N2 and the M362L variant (Figure 4C). At a concentration of 1 μM, IsdB-N2 WT and M362L variant had very similar K_D values (0.38 ± 0.06 and 0.49 ± 0.09 μM, respectively) when fit to an equation accounting for ligand depletion; however, as we are working at a protein concentration close to the calculated K_D (necessarily high for a reproducible fluorescence signal), these values can be considered only an upper bound, with the actual K_D likely being lower. We can conclude that both the native form and M362L variant bind heme with high nanomolar or better affinity. By comparison, the variant proteins that exhibited aberrant or weak heme binding by visible spectroscopy required much greater (at least 5-fold higher) heme concentrations for maximal quenching, and curve fitting suggested dissociation constants at least 5-fold higher (data not shown).

Rate of Transfer of Heme to Apomyoglobin. The transfer of heme from the binding pockets of IsdB-N2 WT and M362L...
variant to apomyoglobin (apoMb) was followed by visible stopped-flow spectroscopy at 408 nm, the Soret maximum for holomyoglobin. ApoMb has an affinity for heme in the picomolar range, does not directly bind to IsdB, and has been used to characterize heme binding by Isd proteins previously. The rate of transfer of heme to apoMb is independent of the concentration of apoMb; therefore, the observed rate is assumed to be the rate of release of heme (off rate) from IsdB-N2. The transfer rate data are best fit by a double-exponential curve, as judged by the greater randomness of the residual plot than that for a single-exponential curve. The initial fast phase accounted for only 2.8% of the curve for WT and 3.5% for M362L and may represent the presence of a minor extraneous species in solution, such as improperly folded holo-IsdB-N2 or IsdB-N2 with nonspecifically bound surface heme. The slow phase, describing the vast majority of the absorption change, yielded an off rate of 4.8 s⁻¹ for WT IsdB-N2 (Figure 5A), and the substitution of Met362 with Leu (Figure 5B) increased the off rate by 2.7-fold, to 1.3 s⁻¹. A summary of kinetic and equilibrium binding characteristics can be found in Table 2.

**Rate of Transfer of Heme to IsdA-N1.** Using stopped-flow spectroscopy, the observed rates (kₜₐₜ) of transfer of heme from holo-IsdB-N2 to apo-IsdA-N1 were determined by the difference in the Soret region of their visible spectra at the wavelength of maximal change over ~0.3 s (for WT) or ~1.0 s (for M362L), equivalent to a minimum of 15 half-lives. The change in absorbance at 418.9 nm (WT) or 425.3 nm (M362L) was plotted versus time and fit by a single exponential to determine a value for the observed rate constant (kₜₐₜ). The kₜₐₜ for 2 μM IsdB-N2 was determined under pseudo-first-order conditions for
Concentrations of IsdA-N1 from 10 to 50 μM. The \( k_{\text{obs}} \) values vary hyperbolically with respect to the concentration of IsdA-N1, suggesting a two-step transfer mechanism. A model for heme transfer has been proposed by Liu et al., which states that the transfer of heme from the holo-NEAT domain to the apo-NEAT domain has two observable steps characterized by rapid formation of the protein–protein complex followed by rate-limiting heme transfer between the proteins. Using this model, the rate constant \( k_1 \) was found to decrease by more than 2-fold, to 34 ± 3 s\(^{-1} \) (Figure 6A). The calculated \( k_1 \) for the protein–protein complexes was similar for both WT and M362L transfer reactions (18 and 16 μM, respectively).

### DISCUSSION

Investigating the molecular basis for the uptake of heme by *S. aureus* is a crucial step toward understanding host–pathogen interactions. IsdB is the dominant Hb-binding protein of *S. aureus*. We have found that the heme binding NEAT domain (IsdB-N2) of IsdB adopts a fold similar to those observed in other NEAT domain structures. Additionally, heme-iron is coordinated by a conserved Tyr residue as predicted by multiple-sequence alignments and observed in other heme binding NEAT domains in *S. aureus*. However, a unique feature of IsdB-N2 is a distal heme-iron ligand, Met362. The equivalent position in IsdA-N1 is taken by a His, which is a common iron ligand but does not coordinate to the heme-iron in the crystal structure or in solution in the absence of a strong reductant.

In the heme binding NEAT domains of IsdC (IsdC-N1) and IsdH (IsdH-N3), there are an Ile and a Val, respectively, neither of which has the capacity for heme-iron coordination. As well as being unprecedented in the Isd system of *S. aureus*, Tyr-Met heme-iron coordination has not been reported in the literature for any heme binding protein to date.

The protein Shp from *Streptococcus pyogenes* uses an unusual bis-methionyl heme-iron coordination as part of a heme uptake system with some components distantly related to the Isd system of *S. aureus*. The crystal structure of the NEAT-like domain of Shp, denoted Shp180, superposes on IsdB-N2 with a core rmsd of 2.5 Å over 94 Cα atoms using SSM Superposition. The average Fe–S bond length between the sulfur of the coordinating Met residues of Shp and the heme-iron was found to be 2.4 Å, shorter than the Fe–S bond length of 2.5–2.6 Å found here. By comparison, the Fe–S bond length seen in IsdB, which coordinates heme-iron using Met and His, is shorter at 2.3 Å. The Fe–O bond length (Tyr–heme-iron) observed in IsdB-N2, which ranges from 2.1 to 2.3 Å depending on the molecule, is typical of those seen in other NEAT domain structures: 2.1 Å for IsdA-N1, 2.2 Å for IsdH-N3, and 2.0–2.1 Å for IsdC-N1.

It is noteworthy that previous work using electronic and magnetic circular dichroism (MCD) spectroscopy did not identify Met as a heme-iron ligand in IsdB-N2, though Tyr was accurately predicted to be a ligand. The discrepancy may be explained by photoreduction of the heme-iron to Fe(II) in the crystal by the X-ray radiation, resulting in a preference for ligation by methionine; beamline photoreduction is a well-known issue when studying hemoproteins and can result in conformational changes as well as changes in coordination state. However, IsdA-N1 contains a potential distal heme-iron ligand (His83) that was not observed to coordinate heme-iron in the crystal structure. Moreover, it has been shown that reduction of the
heme-iron in IsdA results in heme iron coordination solely by the distal His83. Given that Met362 clearly occupies both coordinating and noncoordinating conformations in the IsdB-N2 crystal structure, we suggest that it may do the same in solution, consistent with spectra indicating a predominantly five-coordinate heme-iron with the sixth position readily available to bind exogenous CN⁻. Aside from Tyr440 and Met362, two other residues form important interactions to secure heme: Ser361, which hydrogen bonds to one propionate, and Tyr444, which hydrogen bonds with Tyr440. All four residues were mutated separately to probe their involvement in stable heme binding by IsdB-N2, resulting in the Y440A, Y444A, M362L, and S361A variants. Of these four, only the M362L variant was proficient at heme binding: a gel filtration step intended to purify hemoprotein after heme reconstitution resulted in mainly apoprotein in the cases of the Y440A, Y444A, and S361A variants (data not shown). The apparent deficiency in heme binding of these variants was further characterized by examining their visible electronic spectra, which more closely resembled that of free heme than that of hemoprotein. The loss of high-affinity heme binding by the S361A variant suggests that the interaction with the heme propionate is crucial for the stability of the loop that includes Met362 and forms one side of the heme binding pocket. Similarly, Tyr444 is absolutely conserved in all NEAT domains of the Isd system, heme binding or not; the data suggest that it is a critical heme binding residue as well, although it does not directly interact with the heme. Tyr440 may be required to stabilize the fold of NEAT domains as well as to position Tyr440 for heme-iron coordination or to mediate the phenolate–iron bond.

To investigate the significance of Met362 with regard to potential functional roles in the heme pocket, we conducted a series of spectroscopic and kinetic characterizations of the WT IsdB-N2 protein in comparison with an M362L variant. We found that the variant retained at minimum low micromolar affinity for heme, as measured by tryptophan fluorescence quenching. The 1:1 stoichiometry of heme binding was not altered either; however, the off rate for removal of heme from the pocket increased by 2.7-fold. This may be explained by considering that loss of a heme-iron ligand weakens heme binding and may also destabilize the distal loop, allowing an increased level of solvation of the heme in the pocket. In contrast, the M362L substitution causes a decrease in the first-order rate constant for the transfer of heme to IsdA-N1 by half, from 81 to 34 s⁻¹. The combined increase in the heme off rate by the M362L variant paired with a decreased rate of transfer to IsdA-N1 results in a ratio of catalyzed heme transfer to heme release of 6.4-fold less for the M362L variant (transfer rate ~2600-fold greater than the off rate) compared to the wild-type protein (transfer rate ~17000-fold greater than the off rate). The implication is that Met362 plays a role in the passage of heme between IsdA and IsdN NEAT domains. The transfer of heme between the NEAT domain-containing hemophore IsdX1 and cell wall-anchored IsdC of Bacillus anthracis was observed to be biphasic and slightly slower than that seen here (fast phase of 13 s⁻¹) but once again was at least 10000-fold faster than the observed off rate. The mechanism of inter-NEAT domain heme transfer is not yet known; however, in the case of IsdB-N2, Met362 may act to pull the heme-iron away from Tyr440, weakening the bond and thus facilitating transfer to IsdA. The observed bond lengthening between Tyr440 and the heme-iron in chains A and D, where Met362 mainly coordinates, supports this hypothesis.

A comparison with the streptococcal protein Shp again reveals surprising parallels. As previously mentioned, Shp uses two methionines for heme-iron coordination, namely, Met66 and Met153, located in positions structurally analogous to IsdB-N2 Met362 and Tyr440, respectively. Mutation of the Met66 or Met153 heme-iron ligand to Ala was found to have dramatically different effects on the heme binding and kinetic parameters of the protein. An M66A variant caused minimal change in the heme dissociation constant (22 µM) compared with the wild-type protein (22 µM); conversely, mutation of the other heme iron ligand, Met153, resulted in a 3-fold increase in the heme dissociation constant (62 µM). An investigation of the effect of Met153 and Met66 mutation on rates of transfer to streptococcal HtsA, the cognate lipoprotein receptor for Shp-heme, also revealed striking differences between the two variants. Whereas mutation of Met66 to Ala resulted in a 7.5-fold decrease in the rate-limiting transfer step (0.4 s⁻¹ vs 2.9 s⁻¹ for the wild type), mutation of Met153 to Ala resulted in very little change in the rate-limiting transfer step (2.5 s⁻¹). Parallels between IsdB-N2 and Shp reveal that abolishing a heme-iron ligand can have highly variable effects on the heme binding and transfer characteristics of the protein.

Nonetheless, the extent of the effects of the loss of Met362 as a heme iron ligand in IsdB is likely not yet fully characterized. As ferric iron (Fe³⁺) displays a preference for the phenolate of Tyr and ferrous iron (Fe²⁺) prefers Met or His, possessing both Met and Tyr heme-iron ligands may provide IsdB the flexibility to bind heme-iron in both oxidation states. For instance, S. aureus secretes hemolysins during infection that lyse red blood cells, releasing the stored Hb in a reduced state. Hb heme-iron quickly oxidizes in the bloodstream and is subsequently lost from Hb at an accelerated rate. If heme-iron is not completely oxidized, being able to bind ferrous heme-iron through Met362 may be advantageous for S. aureus. It is also possible that Met362 plays a role in extraction of heme from Hb; however, we have found that mixing IsdB-N2 with Hb did not result in observable heme transfer (data not shown), which is consistent with published observations that IsdB-N2 alone does not interact with Hb.

Though IsdB-N2 and IsdH-N3 (the heme binding NEAT domain of IsdH) are 56% identical in sequence and both IsdB and IsdH are known to bind Hb, the loss of IsdB significantly hampers the cells’ utilization of Hb as a sole iron source. Furthermore, a solution of IsdH-N1 (Hb binding NEAT domain) and IsdH-N3 takes up heme from Hb at a rate slightly faster than the rate of dissociation of heme from Hb, 11 h⁻¹, whereas full-length IsdB takes up heme from Hb at a rate of 100 times faster at a rate of 0.31 s⁻¹. While regions outside the individual NEAT domains may be required for efficient extraction of heme by IsdH, it is also possible that the unique Met362 residue somehow participates in Hb recognition or heme extraction and expedites the process. An M362L variant of full-length IsdB is thus planned to explore this possibility.
what that may be, as each NEAT domain heme pocket is built to coordinate one heme at a time. An investigation of transfer kinetics between IsdH-N3 and IsdA or IsdC may help to resolve this question.

A comparison of spectroscopic and kinetic results between full-length IsdB and IsdB-N2 reveals highly similar heme binding and transfer characteristics. Zhu et al. were able to determine an off rate of $1.3 \times 10^{-3} \text{s}^{-1}$ for recombinant full-length IsdB by transfer of heme to apoMb, close to our value of $4.8 \times 10^{-3} \text{s}^{-1}$. Transfer of heme between IsdB and IsdA also appears to be closely mirrored by the NEAT domains alone. A solution of 3 μM full-length holo-IsdB transferred heme to a solution of 30 μM full-length apo-IsdA at a rate of 114 s$^{-1}$, whereas using 2 μM holo-IsdB-N2, we observed a rate of transfer to 30 μM apo-IsdA-N1 of $\sim 50 \text{s}^{-1}$. Because the buffers, temperatures, and protein concentrations of the experiments differ, these rates are not directly comparable; nonetheless, they are on the same order of magnitude, supporting the hypothesis that NEAT domains alone are sufficient for heme transfer. Finally, the close similarity in the rate of transfer to IsdA or IsdA-N1 demonstrates the likelihood that the unusual Tyr-Met heme-iron coordination observed in the NEAT domain crystal structure is recapitulated in the full-length IsdB protein.

In summary, we have defined the structural basis for heme binding by IsdB. The second NEAT domain of IsdB is structurally similar to other heme binding NEAT domains, and heme-iron is coordinated by a tyrosine residue as predicted; however, we have also shown that Met362 coordinates the heme-iron on the distal side, resulting in an unprecedented mode of heme-iron coordination. We also demonstrate that Met362 plays a role in the distal side, resulting in an unprecedented mode of heme-iron-iron coordination. Zhu et al. were able to determine an off rate of $1.3 \times 10^{-3} \text{s}^{-1}$ for recombinant full-length IsdB by transfer of heme to apoMb, close to our value of $4.8 \times 10^{-3} \text{s}^{-1}$. Transfer of heme between IsdB and IsdA also appears to be closely mirrored by the NEAT domains alone. A solution of 3 μM full-length holo-IsdB transferred heme to a solution of 30 μM full-length apo-IsdA at a rate of 114 s$^{-1}$, whereas using 2 μM holo-IsdB-N2, we observed a rate of transfer to 30 μM apo-IsdA-N1 of $\sim 50 \text{s}^{-1}$. Because the buffers, temperatures, and protein concentrations of the experiments differ, these rates are not directly comparable; nonetheless, they are on the same order of magnitude, supporting the hypothesis that NEAT domains alone are sufficient for heme transfer. Finally, the close similarity in the rate of transfer to IsdA or IsdA-N1 demonstrates the likelihood that the unusual Tyr-Met heme-iron coordination observed in the NEAT domain crystal structure is recapitulated in the full-length IsdB protein.

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