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# The structure of haemoglobin bound to the haemoglobin receptor IsdH from *Staphylococcus aureus* shows disruption of the native $\alpha$ -globin haem pocket

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Staphylococcus aureus is a common and serious cause of infection in humans. The bacterium expresses a cell-surface receptor that binds to, and strips haem from, human haemoglobin (Hb). The binding interface has previously been identified; however, the structural changes that promote haem release from haemoglobin were unknown. Here, the structure of the receptor–Hb complex is reported at 2.6 Å resolution, which reveals a conformational change in the  $\alpha$ -globin F helix that disrupts the haem-pocket structure and alters the Hb quaternary interactions. These features suggest potential mechanisms by which the *S. aureus* Hb receptor induces haem release from Hb.

#### 1. Introduction

Staphylococcus aureus is a Gram-positive bacterial pathogen that is carried asymptomatically by approximately one third of the human population, and yet has the capacity to cause fatal invasive infection (Kluytmans et al., 1997). Although S. aureus strains that are resistant to  $\beta$ -lactam antibiotics have traditionally been associated with hospital-acquired infections, antibiotic-resistant strains have now become commonplace in the general community (Dukic *et al.*, 2013). Understanding the mechanisms that S. aureus uses to survive inside the human host is an important step towards the discovery of new treatments that neutralize pathogen virulence. Iron-uptake pathways are of interest in this regard because iron is required by many essential bacterial enzymes, and access to iron is necessary for bacterial infection (Nairz et al., 2010). The most abundant form of iron in the human body is haem (iron protoporphyrin-IX), and the majority of haem is found in haemoglobin (Hb). S. aureus has evolved a pathway, encoded by nine iron-regulated surface determinant (isd) genes (Mazmanian et al., 2003; Morrissey et al., 2002), that is specialized for the uptake of haem/iron from human Hb (Pishchany et al., 2010).

Two of the Isd proteins, IsdB and IsdH, are cell-surface proteins that bind to Hb and remove haem groups by a process that is not completely understood (Dryla *et al.*, 2003, 2007; Torres *et al.*, 2006; Pilpa *et al.*, 2009; Pishchany *et al.*, 2010; Krishna Kumar *et al.*, 2011; Dickson *et al.*, 2014). IsdB and IsdH supply two haem chaperones, IsdA and IsdC, which relay haem to the substrate-binding subunit (IsdE) of a haem transporter that spans the bacterial cell membrane (IsdF) (Zhu *et al.*, 2008; Liu *et al.*, 2008; Muryoi *et al.*, 2008;

Tiedemann *et al.*, 2012; Grigg *et al.*, 2007*a*, 2010; Pluym *et al.*, 2007; Tiedemann & Stillman, 2012). IsdH binds more tightly to iron(III) haem than to iron(II) haem (Moriwaki *et al.*, 2011), suggesting that the Hb receptors target ferric metHb, which forms rapidly following erythrocyte lysis. Inside the bacterial cytoplasm the porphyrin macrocycle is cleaved by either of two haem oxygenase enzymes, IsdG or IsdI, to release iron (Wu *et al.*, 2005).

IsdA/B/C/H proteins are all anchored *via* a C-terminal amide linkage to the peptide cross-bridge of the cell-wall peptidoglycan (Mazmanian *et al.*, 2002), and all bind haem *via* 



#### Figure 1

(a) Domain structure of *S. aureus* IsdB and IsdH, and sequence homology between these domains. The construct crystallized in this study contains the domains highlighted in red (IsdH<sup>N2N3</sup>). (b) Crystal of IsdH<sup>N2N3</sup>- metHb. (c) The crystallographic asymmetric unit contained an Hb  $\alpha\beta$  dimer with a single molecule of IsdH<sup>N2N3</sup> bound to the  $\alpha$  subunit. (d) Detail of the haem-pocket region of the  $\alpha$  subunit from IsdH<sup>N2N3</sup>-metHb (orange) overlayed with the metHb structure (PDB entry 3p5q; slate blue)

a near iron transporter (NEAT) domain, which is an antiparallel  $\beta$ -barrel domain of 15–20 kDa with an immunoglobulin-like topology (Vu et al., 2013; Moriwaki et al., 2011; Watanabe et al., 2008; Pilpa et al., 2006; Grigg et al., 2007b, 2011; Villareal et al., 2008; Sharp et al., 2007; Gaudin et al., 2011). Related NEAT domains that function in haem acquisition have been described in Streptococcus pyogenes (Aranda et al., 2007), Listeria monocytogenes (Malmirchegini et al., 2014) and Bacillus anthracis (Ekworomadu et al., 2012; Honsa et al., 2013), highlighting the importance of NEAT domains for iron uptake in Gram-positive pathogens. The haem-binding site in the NEAT domain is located between a short 310-helix and the  $\beta$ -hairpin formed by  $\beta$ -strands 6 and 7 (numbered according to homology with the immunoglobulin domain topology). The haem iron is typically five-coordinate iron(III) (Pluym et al., 2008; Vermeiren et al., 2006) and is bound to the protein through the first Tyr side chain from a conserved YxxxY motif in  $\beta$ -strand G. IsdB and IsdH contain one or two additional NEAT domains, respectively, that lack haembinding activity and instead bind to Hb chains at a site comprising parts of the A and E helices of the globin (Dryla et al., 2003, 2007; Pilpa et al., 2009; Pishchany et al., 2010; Krishna Kumar et al., 2011; Dickson et al., 2014).

A three-domain region of IsdH comprising the Hb-binding second NEAT domain (IsdH<sup>N2</sup>) and haem-binding third NEAT domain (IsdH<sup>N3</sup>) separated by an  $\alpha$ -helical linker domain (IsdH<sup>Linker</sup>) is required to capture haem from Hb (Spirig et al., 2013; Dickson et al., 2014). This three-domain region of IsdH, referred to hereafter as IsdH<sup>N2N3</sup> (Fig. 1*a*), shares 62% sequence homology with IsdB. Haem transfer from metHb to IsdB occurs with a rate constant  $k = 0.31 \text{ s}^{-1}$  at 25°C (Zhu et al., 2008). Haem transfer from metHb to Isd $H^{N2N3}$  is biphasic, with rate constants of 0.85 and 0.099 s<sup>-1</sup> (Sjodt et al., 2015). These rates are 2-3 orders of magnitude faster than the thermal dissociation of haem from the  $\beta$  (k = 0.003 s<sup>-1</sup>) or  $\alpha$  (k = 0.0002 s<sup>-1</sup>) sites of metHb under similar solution conditions (Hargrove et al., 1996), indicating that the receptors activate haem release from metHb. To investigate the mechanism of haem extraction, we previously determined the structure of IsdH<sup>N2N3</sup> bound to metHb by X-ray crystallography at a resolution of 4.2 Å (PDB entry 4ij2; Dickson et al., 2014). In this structure, IsdH<sup>N2N3</sup> carried an Ala mutation of the haem-ligating Tyr642 (located in the IsdH<sup>N3</sup> domain), resulting in crystallization of the IsdH<sup>N2N3</sup>-metHb complex in a state prior to haem transfer. The crystal structure with PDB code 4ij2 contained a metHb tetramer with one IsdH<sup>N2N3</sup> receptor bound to each of the four globin chains. IsdH<sup>N2N3</sup> bound  $\alpha$  or  $\beta$  subunits in a similar manner, with the haemaccepting IsdH<sup>N3</sup> domain positioned precisely over the entrance to the globin haem pocket. However, the data resolution was not sufficient to identify the tertiary-structure changes in the globin that are presumed to facilitate haem transfer.

In this work, we have engineered mutations into the Isd $H^{N2}$  domain of Isd $H^{N2N3}$  that effectively blocked binding to Hb  $\beta$  subunits, but left binding to Hb  $\alpha$  unchanged. Crystals grown using this mutant gave dramatically improved diffraction

quality. We report the crystal structure of metHb with IsdH<sup>N2N3</sup> bound only at the Hb  $\alpha$  subunits at a resolution of 2.6 Å. The structure reveals a disturbance of the globin haem pocket and provides a potential mechanism for haem release.

#### 2. Materials and methods

#### 2.1. Mutagenesis

The IsdH expression vector used in this study was a modified version of the pRM208 plasmid, which contained the second and third NEAT domains and the intervening IsdH<sup>Linker</sup> domain of IsdH (IsdH<sup>N2N3</sup> residues 326–660) cloned between the BamHI and XhoI sites of the pHis-SUMO vector (Spirig *et al.*, 2013). The modified QuikChange method of Liu & Naismith (2008) was used to introduce IsdH mutations FYHYA(365–369) $\rightarrow$ YYHYF and Y642A into the pRM208 plasmid.

#### 2.2. Protein production

IsdH<sup>N2N3</sup> carrying the FYHYA(365–369) $\rightarrow$ YYHYF and Y642A mutations was transformed into E. coli strain Rosetta2 (DE3) pLysS and expression was performed at 37°C in LB broth containing 34  $\mu$ g ml<sup>-1</sup> chloramphenicol and 50  $\mu$ g ml<sup>-1</sup> kanamycin with addition of 1 mM IPTG to induce expression. The expressed protein comprised the N-terminal sequence MGSSHHHHHHSSGLVPRGSHMAS followed by the 11.3 kDa ubiquitin-like protein SMT3 from Saccharomyces cerevisiae followed by the IsdH region of interest. This protein was purified at 4°C over IMAC resin (HIS-Select Nickel Affinity gel, Sigma). Fractions containing the fusion protein were pooled and mixed in a 100-fold molar excess over His-tagged ubiquitin-like protein-1 protease for 1 h at room temperature. Following dialysis, the mixture was reapplied onto the IMAC column to remove the His-SUMO tag and Histagged ubiquitin-like protein-1 protease. Additional purification steps were performed by anion-exchange (Q Sepharose, GE Healthcare Life Sciences) and gel-filtration (Superose 12HR, GE Healthcare Life Sciences) chromatography.

Hb was purified from blood as reported previously (Gell et al., 2002). During Hb purification the globin was maintained in the carbon monoxide-liganded state to inhibit autooxidation. Although we required oxidized metHb for structural work, purification in the HbCO form was beneficial to inhibit the autooxidation and subsequent irreversible denaturation that otherwise occurred. Briefly, red blood cells, freshly collected from a human volunteer, were washed in saline solution and then lysed under hypotonic conditions. Hb was purified from the haemolysate by cation-exchange chromatography (SP Sepharose, GE Healthcare Life Sciences) followed by anionexchange chromatography (Q Sepharose, GE Healthcare Life Sciences). HbCO was converted to HbO<sub>2</sub> by passing a pure stream of oxygen over a protein solution held on ice and illuminated with a high-intensity light source. HbO<sub>2</sub> was converted to ferric metHb by incubation with a fivefold molar excess of potassium ferricyanide at  $\sim 10^{\circ}$ C in 20 mM sodium phosphate pH 7.0. The reaction was monitored to completion

#### Table 1

Data-collection and refinement statistics.

Values in parentheses are for the highest resolution shell.

Data collection	
Wavelength (Å)	0.95370
Space group	P6 <sub>5</sub> 22
Unit-cell parameters (Å, °)	a = b = 92.22, c = 365.12,
•	$\alpha = \beta = 90, \gamma = 120$
Resolution (Å)	33.39-2.55 (2.66-2.55)
Observed reflections	342621
Unique reflections	33107
$R_{\rm merge}$ † (%)	13.3 (78.4)
$\langle I/\sigma(I)\rangle$	9.3 (2.0)
$CC_{1/2}$ ‡	0.996 (0.766)
Completeness (%)	99.9 (100)
Multiplicity	9.6 (9.7)
Refinement	
Resolution (Å)	33.39-2.55 (2.61-2.55)
Reflections in working set	29544
Reflections in test set	1581
$R_{ m work}$ (%)	25.9 (36.4)
$R_{\rm free}$ § (%)	28.7 (39.2)
No. of non-H atoms	
Protein	4775
Ligand/ion	94
Water	4
B factors $(Å^2)$	
Protein	63.4
Ligand/ion	77.7
Water	40.0
R.m.s. deviations	
Bond lengths (Å)	0.004
Bond angles (°)	0.8
Ramachandran plot <sup>††</sup>	
Most favoured (%)	96.6
Allowed (%)	3.1
Outliers (%)	0.3
PDB code	4xs0

 $\begin{array}{l} \dagger \ R_{\rm merge} = \sum_{hkl} \sum_i |I_i(hkl) - \langle I(hkl) \rangle | / \sum_{hkl} \sum_i I_i(hkl). \\ \ddagger \ Values \ of \ CC_{1/2} \ according \ to \\ Karplus \ \& \ Diederichs (2012). \\ \$ \ R_{\rm free} \ was \ calculated \ using 5\% \ of \ the \ reflections, which \\ were \ chosen \ at \ random \ and \ excluded \ from \ the \ data \ set. \\ \P \ Deviations \ from \ ideal \ values \\ (Engh \ \& \ Huber, \ 1991). \\ \dagger^{\dagger} \ Calculated \ using \ MolProbity \ (Chen \ et \ al., \ 2010). \end{array}$ 

by UV–visible spectroscopy and metHb was isolated over G-25 Sepharose (GE Healthcare Life Sciences). The concentration of Hb was determined by UV–visible absorption spectroscopy using the molar absorption coefficients  $\varepsilon_{419} = 192\ 000\ M^{-1}\ {\rm cm}^{-1}$  and  $\varepsilon_{539} = 13\ 900\ M^{-1}\ {\rm cm}^{-1}$  for HbCO and  $\varepsilon_{405} = 169\ 000\ M^{-1}\ {\rm cm}^{-1}$  and  $\varepsilon_{500} = 9000\ M^{-1}\ {\rm cm}^{-1}$  for metHb (Eaton & Hofrichter, 1981)

#### 2.3. Haemoglobin-binding assays

Size-exclusion chromatography (SEC) experiments were performed on Superose 12 (GE Healthcare Life Sciences) equilibrated in 150 mM sodium phosphate pH 7.0. Proteins were mixed at the concentrations indicated and 100  $\mu$ l samples were loaded onto the column. Elution of protein samples was monitored at 280 or 410 nm with a Jasco UV2070 detector.

#### 2.4. Crystallization

IsdH<sup>N2N3</sup> mutant protein and metHb were bufferexchanged into 50 m*M* HEPES pH 7.5, mixed in a 2:1 molar ratio to give a final protein concentration of 11.1 mg ml<sup>-1</sup> and subjected to sparse-matrix crystallization screening at 21°C. Hexagonal crystals grew reproducibly in 0.2 *M* potassium/ sodium tartrate, 0.1 *M* trisodium citrate pH 5.6, 2 *M* ammonium sulfate. In a fine screen, crystals of identical morphology were obtained over a pH range of 5.4–5.8 and a precipitant concentration of 2.0–2.2 *M* ammonium sulfate. The crystals were cryoprotected in up to 30% glycerol before being flash-cooled in liquid nitrogen.

#### 2.5. Data collection and processing

Diffraction data were collected on the MX1 beamline at the Australian Synchrotron to a resolution of 2.55 Å. The data were integrated with *iMosflm* (Leslie & Powell, 2007) and scaled using *AIMLESS* (Evans & Murshudov, 2013). Table 1 contains the collection and refinement statistics.

#### 2.6. Structure solution and refinement

The structure was solved by molecular replacement. The search model was derived from the 4.2 Å resolution complex of IsdH bound to metHb (PDB entry 4ij2) and included an Hb  $\alpha\beta$  dimer with one IsdH<sup>N2N3</sup> bound through the  $\alpha$  subunit. Molecular replacement performed with *Phaser* (McCoy *et al.*, 2007) gave rise to a single solution. A single round of rigid-body refinement followed by multiple rounds of restrained refinement were performed with *REFMAC* 5.7 (Murshudov *et al.*, 2011) and model building was completed with *Coot* (Emsley & Cowtan, 2004). In later rounds of refinement,

translation/liberation/screw refinement was introduced, with groups defined as single domains (where the IsdH<sup>Linker</sup>–IsdH<sup>N3</sup> unit was considered to be a single group). The final model was validated using *MolProbity* (Chen *et al.*, 2010). Ramachandran statistics for the final model are as follows: 96.6% of residues were found in favoured regions and 3.1% in allowed regions. The structure factors and coordinates have been deposited in the Protein Data Bank as entry 4xs0.

#### 2.7. Analysis of Hb tertiary and quaternary structure

To understand quaternary-structural changes in Hb, Baldwin and Chothia identified a set of residues in the  $\alpha 1\beta 1$ dimer with unchanged positions in the liganded relaxed (R) and ligand-free tense (T) states that act as a reference frame for tertiary- and quaternary-structural transitions, referred to as the BGH frame and comprising residues  $\alpha 30-\alpha 36$ ,  $\alpha 101-\alpha 113$ ,  $\alpha 117-\alpha 127$  and  $\beta 30-\beta 36$ ,  $\beta 51-\beta 55$ ,  $\beta 107-\beta 132$  (Baldwin & Chothia, 1979). The reference set was updated by Park and coworkers based on higher resolution (1.25 Å) structures to comprise residues  $\alpha 23-\alpha 42$ ,  $\alpha 57-\alpha 63$ ,  $\alpha 101-\alpha 111$ ,  $\alpha 118-\alpha 125$ and  $\beta 51-\beta 57$ ,  $\beta 110-\beta 116$ ,  $\beta 119-\beta 132$  (Park *et al.*, 2006). In this study, we perform overlays using the original BGH frame, but note that the results are not significantly altered when using the reference frame of Park and coworkers. Relative rotations/ translations of  $\alpha\beta$  dimers were calculated using *DynDom* 



IsdH<sup>N2N3</sup> carrying the FYHYA(365–369)  $\rightarrow$  YYHYF mutation binds Hb  $\alpha$  but not Hb  $\beta$ . (a) An equimolar mixture of  $\alpha$  and IsdH<sup>N2N3</sup> eluted predominantly as a single peak from SEC (grey dashed–dotted line), indicating the formation of a protein complex. Elution profiles for free  $\alpha$  (solid line) and IsdH<sup>N2N3</sup> (dotted line) are shown for comparison. (b) An equimolar mixture of  $\alpha$  and IsdH<sup>N2N3</sup> carrying the FYHYA $\rightarrow$ YYHYF mutation eluted predominantly as a single peak from SEC (grey dashed–dotted line), indicating the formation of a complex with a similar elution time to the wild-type complex in (a). (c) An equimolar mixture of  $\beta$  and IsdH<sup>N2N3</sup> (grey dashed–dotted line) produced a peak with an earlier elution time than free  $\beta$  (solid line) or IsdH<sup>N2N3</sup> (dotted line), indicating the formation of a complex between  $\beta$  and IsdH<sup>N2N3</sup>. (d) An equimolar mixture of  $\beta$  and IsdH<sup>N2N3</sup> carrying the FYHYA $\rightarrow$ YYHYF mutation yields an elution profile (grey dashed–dotted line) as expected for a mixture of free  $\beta$  (solid line) and IsdH<sup>N2N3</sup> (dotted line); thus, no interaction was detected between  $\beta$  and IsdH<sup>N2N3</sup> carrying the FYHYA $\rightarrow$ YYHYF mutation. Note that  $\alpha$  is predominantly a monomer, whereas  $\beta$  is predominantly a tetramer under the conditions of the assay, explaining the substantial difference in the elution time of the free Hb chains.

 Table 2

 Hydrogen-bonding and salt-bridge interactions at the IsdH<sup>N2N3</sup>-metHb interfaces

$IsdH^{N2N3}$ (chain C)	Hb $\alpha$ subunit (chain A)	Distance (Å)		
Tyr443 OH	Asn9 OD1	2.71		
Lys391 NZ	Ala71 O	3.13		
Tyr421 OH	Asp74 OD2	2.52		
Tyr495 OH	Ser81 O	3.90		
Lys503 NZ	Ser81 OG	3.81		
Tyr495 OH	Asp85 OD2	2.72		
Thr392 OG1	Lys11 NZ	2.98		
Ser370 OG	Lys11 NZ	2.83		
Glu449 OE2	Lys16 NZ	2.92		
Tyr365 OH	Ala19 N	3.39		
Tyr646 OH	His45 NE2	3.24		
Glu562 OE2	Gln54 NE2	3.87		
Val564 O	Gln54 NE2	3.58		
Ser563 O	Gln54 NE2	3.45		
Lys499 NZ	Asp85 OD2	3.63		
Glu449 OE1	Lys16 NZ	3.38		
Glu449	Lys16 NZ	2.92		
Haem O1D	Tyr646 OH	2.95		
Haem O2D	Tyr646 OH	2.84		



#### Figure 3

Structural features of the IsdH<sup>N2N3</sup> subunit. (a) Structure of the aromatic helix carrying the FYHYA $\rightarrow$ YYHYF mutation (blue) at the interface with Hb  $\alpha$  (orange). An  $F_o - F_c$  density map contoured at  $3\sigma$  (green mesh) was generated by omitting residues 362–369 before refinement. (b) Structure of the aromatic helix carrying the FYHYA $\rightarrow$ YYHYF mutation (blue) superposed with the structure of the corresponding loop from IsdH<sup>N1</sup> with sequence YYHFF (pink; PDB entry 3szk). (c) Structure of the native aromatic helix in IsdH<sup>N2</sup> with sequence FYHYA (green) at the interface with Hb  $\alpha$  (PDB entry 4fc3). (d) Structure of the loop linking the IsdH<sup>Linker</sup> and IsdH<sup>N3</sup> domains, showing OMIT electron density for the region not previously observed in PDB entry 4ij2 or structures of the isolated domains ( $F_o - F_c$  OMIT map contoured at  $3\sigma$ , where residues 532–543 were omitted from the model prior to refinement).

(Hayward & Berendsen, 1998). To emphasize the structural features of the haem pocket that are conserved between the  $\alpha$  and  $\beta$  subunits, and across other globins with divergent sequences, amino-acid residues are identified (in parentheses) according to their corresponding helix, and position in the helix, in the related structure of sperm whale myoglobin, which is the standard reference structure by convention. For example,  $\alpha$ His87(F8) indicates a residue with structural similarity to the eighth residue in helix F of sperm whale myoglobin, which is the haem-ligating proximal His.

#### 3. Results and discussion

3.1. Engineering of IsdH to remove weak binding to Hb  $\beta$  chains

In the crystal structure with PDB code 4ij2, electron density was missing for the haem-acceptor IsdH<sup>N3</sup> domain belonging to one of the four IsdH<sup>N2N3</sup> receptors in the crystallographic asymmetric unit, suggesting disorder in the position of this

IsdH<sup>N3</sup> domain. The 'missing' domain belonged to an IsdH<sup>N2N3</sup> receptor that was bound to an Hb  $\beta$  subunit. We hypothesized that steric factors might interfere with the simultaneous binding of IsdH to all four globin subunits of a single Hb tetramer and, therefore, that selectively removing binding to either the  $\alpha$  or  $\beta$  subunits might allow the growth of crystals that are better ordered. Based on comparison of the globin-interaction face of IsdH<sup>N2</sup>, which binds to both  $\alpha$  and  $\beta$  subunits, with the globin-interaction face of IsdH<sup>N1</sup>, which binds to  $\alpha$  but not  $\beta$  (Krishna Kumar *et* al., 2011), we identified a short  $\alpha$ -helix, rich in aromatic side chains, as a site that potentially influences  $\alpha/\beta$  binding selectivity. We constructed an IsdH<sup>N2N3</sup> variant in which residues in the IsdH<sup>N2</sup> domain, FYHYA (365-369; interfacial residues are underlined), were replaced with the corresponding interfacial residues from IsdH<sup>N1</sup> to give YYHYF. IsdH<sup>N2N3</sup> carrying the FYHYA(365- $369) \rightarrow YYHYF$  mutation bound to Hb  $\alpha$  with similar affinity as the wild-type IsdH<sup>N2N3</sup> sequence, as judged by complex formation assayed by gel filtration (compare Fig. 2a with Fig. 2b). However, compared with the wildtype Isd $H^{N2N3}$  (Fig. 2c), Isd $H^{N2N3}$  carrying the FYHYA(365–369) $\rightarrow$ YYHYF mutation showed a dramatic loss of binding to  $\beta$ Hb (Fig. 2d). These mutations, which are located in the N-terminal NEAT domain, preserve biologically

relevant binding to the  $\alpha$  subunit and do not impede interpretation of the haem-transfer interface mediated by the C-terminal NEAT domain.

For crystallography, the FYHYA(365–369) $\rightarrow$ YYHYF mutation was combined with a Y642A substitution of the haem-ligating side chain in the IsdH<sup>N3</sup> domain, which was also present in the 4ij2 structure, to prevent haem transfer and minimize disassembly of the apo Hb complex (Spirig *et al.*, 2013). The engineered IsdH<sup>N2N3</sup> protein was mixed with Hb in a 2:1 molar ratio. Hexagonal crystals grew in 0.2 *M* potassium/ sodium tartrate, 0.1 *M* trisodium citrate over a pH range of 5.4–5.8 and a precipitant concentration range of 2.0–2.2 *M* ammonium sulfate (Fig. 1*b*).

#### 3.2. Overall structure of the IsdH<sup>N2N3</sup>-metHb complex

structure of IsdH<sup>N2N3</sup>-metHb carrying The the FYHYA(365-369)→YYHYF and Y642A mutations was determined to a resolution of 2.55 Å and refined to an  $R_{\text{cryst}}$ and  $R_{\text{free}}$  of 0.259 and 0.287, respectively (Table 1). The crystallographic asymmetric unit contains an Hb  $\alpha\beta$  dimer, with a single IsdH<sup>N2N3</sup> receptor bound to the  $\alpha$  subunit (Fig. 1c), suggesting that our protein-engineering approach to improve crystal order by removing binding to the  $\beta$  subunit was successful. Hydrogen bonds and salt bridges at the IsdH<sup>N2N3</sup>metHb interface are shown in Table 2. Symmetry-related  $\alpha\beta$ dimers form a Hb tetramer in the crystal; however, the relative positioning of  $\alpha\beta$  dimers does not correspond to the canonical T (low O<sub>2</sub> affinity) or R (high O<sub>2</sub> affinity) quaternary structures of Hb (Perutz et al., 1960; Muirhead & Perutz, 1963) and is more similar to two recently determined structures of fully liganded Hbs with a T-like quaternary structure (Safo et al., 2013; Balasubramanian et al., 2014; discussed further below).

Local contacts between the Hb  $\alpha$  subunit and the aromatic helix of IsdH<sup>N2</sup> carrying the FYHYA(365-369)→YYHYF mutation are identical to those observed at the IsdH<sup>N1</sup>- $\alpha$ interface, showing that these contacts were successfully transplanted (Figs. 3a, 3b and 3c). The arrangement of domains in the IsdH receptor and their positions on Hb  $\alpha$  are the same as in PDB entry 4ij2, suggesting that the FYHYA(365–369) $\rightarrow$ YYHYF mutation does not disturb the orientation of the receptor on  $\alpha$ . Electron density and anomalous signal were detected at the globin haem iron sites but not in the receptor haem pockets, suggesting that the complex is trapped in a state prior to haem transfer. A striking feature of the complex is a localized change in the conformation of the Hb  $\alpha$  subunit F helix, which carries the haemligating  $\alpha$ His87(F8), and the FG loop, which participates in quaternary interactions in the Hb tetramer (Fig. 1d). Features of the tertiary and quaternary structure are discussed in the following sections.

#### 3.3. Interdomain linkers of IsdH

The residues linking the three domains of the IsdH<sup>N2N3</sup> receptor, which were absent or unstructured in the isolated domains, were not modelled in the earlier structure (PDB entry 4ij2) owing to the low data resolution. In the current

structure, the polypeptide segment joining the IsdH<sup>Linker</sup> domain to IsdH<sup>N3</sup> (residues Val531-Gln543) is clearly visible in the electron density (Fig. 3d). Helix 3 of the IsdH<sup>Linker</sup> domain is extended by one turn. Val531-Glu535, compared with the solution NMR structure of the free IsdH<sup>Linker</sup> (Spirig et al., 2013). The side chain of Phe534 projects into a pocket between the two sheets of the IsdH<sup>N3</sup>  $\beta$ -sandwich; the pocket is typically occupied by a glycerol molecule in crystal structures of the free IsdH<sup>N3</sup> domain, for example PDB entries 2e7d (Watanabe et al., 2008) and 3qug (Moriwaki et al., 2011). Residues 536-544 adopt an extended conformation across the surface of the IsdH<sup>N3</sup> domain and make two backbone hydrogen bonds with the edge of  $\beta$ -strand 4 (numbered according to homology with the immunoglobulin domain topology). In contrast, the sequence connecting IsdH<sup>N2</sup> to the IsdH<sup>Linker</sup> domain (residues 464-472) together with an adjacent loop of IsdH<sup>N2</sup> (residues 378-383) are not sufficiently well resolved in the electron density for unambiguous interpretation, consistent with some disorder in this region. In summary, the IsdH<sup>Linker</sup> and IsdH<sup>N3</sup> domains pack together across a well ordered interface, suggesting that they behave as a single structural unit, whereas interdomain contacts between the IsdH<sup>Linker</sup> and the IsdH<sup>N2</sup> domain are more limited. This conclusion is consistent with a recent NMR study of the free IsdH<sup>N2N3</sup> receptor, which indicated that the IsdH<sup>N2</sup> domain is able to reorient with respect to IsdH<sup>Linker</sup>–IsdH<sup>N3</sup>, making excursions from its average position of up to 10 Å (Sjodt et al., 2015).

3.4. Interactions between  $IsdH^{N2N3}$  and the Hb *a* haem pocket

Together, the IsdH<sup>N3</sup> and IsdH<sup>Linker</sup> domains make multiple contacts surrounding the entrance to the Hb  $\alpha$  pocket. The positioning of the IsdH<sup>N3</sup> domain means that the haembinding sites in Hb  $\alpha$  and IsdH<sup>N3</sup> are approximately coplanar. Movement of haem from Hb  $\alpha$  to the IsdH<sup>N3</sup> domain requires a translation of  $\sim 12$  Å through the entrance to the globin haem pocket, together with a rotation of  $\sim 115^{\circ}$  in the plane of the porphyrin ring (Fig. 4*a*). The Isd $H^{N3}$  loop containing the  $3_{10}$ -helix makes van der Waals contacts across the CE loop and the N-terminal region of helix E of Hb  $\alpha$ , which together form one side of the haem-pocket entrance and retain a native conformation (Fig. 4b). On the opposite side of the haempocket entrance, Tyr495 and Lys499 from the IsdH<sup>Linker</sup> domain make a hydrogen bond and a salt bridge to  $\alpha$ Asp85(F6), while the Tyr495 ring is packed against the side chain of  $\alpha$ Leu86(F7), which adopts a non-native position (Fig. 4*b*).

The  $\beta$ -hairpin of IsdH<sup>N2N3</sup> carrying Ala642 (Tyr642 in the wild-type complex; Fig. 4*b*) accounts for the majority of contacts with the distorted region of  $\alpha$  (the F helix and FG corner) and forms one wall of a haem-transfer chamber that connects the haem-binding sites of both domains. Other contacts between IsdH<sup>N3</sup> or IsdH<sup>Linker</sup> and Hb  $\alpha$  involve regions of  $\alpha$  with native structure; these contacts may brace the IsdH<sup>N3</sup> domain against stable regions of  $\alpha$  to facilitate



Figure 4

Structural changes in the Hb  $\alpha$  haem pocket. (*a*) Detail of the haem-transfer interface formed between IsdH<sup>N3</sup> (dark blue) and  $\alpha$  (orange, yellow); haem is bound through  $\alpha$ His87(F8). The expected position of the haem group post-transfer is illustrated by superposing the structure of holo IsdH<sup>N3</sup> (PDB entry 2z6f; only the  $\beta$ -hairpin, the Tyr642 side chain and haem group are shown; purple). (*b*) Detail of (*a*) showing interactions at the haem-transfer interface. A region of  $\alpha$  with non-native conformation is coloured yellow. The mutation, Y642A, is highlighted in red. His and Tyr side chains with capacity to act as ligands to the haem iron are shown. (*c*) Stereo diagram of the  $\alpha$  subunit from the IsdH<sup>N2N3</sup>-metHb complex (orange). An  $F_o - F_c$  map generated by omitting residues 80–91 of the  $\alpha$  F helix is shown contoured at  $3\sigma$  (green).  $2F_o - F_c$  electron density, contoured at  $1.5\sigma$ , is also shown for the  $\alpha$  F helix and haem group (*d*) The haem pocket of the haem-coordinating histidine,  $\alpha$ His87(F8), against hydration. (*e*) The haem pocket of Hb  $\alpha$  in the IsdH<sup>N2N3</sup>-metHb complex, showing that the hydrophobic side chains of  $\alpha$ Leu86(F7) and  $\alpha$ Leu91(FG3) are displaced from their native positions. The region of  $\alpha$  with non-native conformation in the IsdH<sup>N2N3</sup>-metHb is coloured yellow.

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conformational exchange of residues surrounding the haem. Structures of the free IsdH<sup>N3</sup> domain in the apo and haem-bound states show that the Tyr642  $\beta$ -hairpin also undergoes a small movement upon haem ligation (Vu *et al.*, 2013; Moriwaki *et al.*, 2011; Watanabe *et al.*, 2008), suggesting that concerted changes in the structure of the IsdH<sup>N3</sup> Tyr642  $\beta$ -hairpin and the Hb  $\alpha$  F helix/FG corner could accompany haem transfer.

We note that Isd proteins of B. anthracis lack the three-domain structure that is conserved in S. aureus IsdB/IsdH, and instead some of the isolated haem-binding domains are able to acquire haem from metHb (Ekworomadu et al., 2012; Honsa et al., 2011). Mutations on the surface of the  $3_{10}$ -helix in the first NEAT domain of *B*. anthracis IsdX1, and the single NEAT domain of B. anthracis IsdX2, disrupt Hb binding or haem uptake from metHb (Ekworomadu et al., 2012; Honsa et al., 2013) and thus it is possible that these NEAT domains associate with globin chains in a similar manner to the IsdH<sup>N3</sup> domain.

3.5. Altered structure of the Hb  $\alpha$  haem pocket

The most striking feature of the IsdH<sup>N2N3</sup>-metHb complex is the altered structure of the  $\alpha$ -subunit haem pocket (Fig. 4). These tertiary-structural changes are described in comparison with the 2.0 Å resolution structure of R-state human metHb (PDB entry 3p5q; Yi et al., 2011). In IsdH<sup>N2N3</sup>metHb, the largest perturbation from the native Hb  $\alpha$  structure occurs in helix F and the FG corner (residues αLeu86–  $\alpha$ Arg93; Figs. 1*d* and 4). The final turn of the F helix ( $\alpha$ Leu86– $\alpha$ His89) is unwound, resulting in a different set of intramolecular contacts for these residues. The side chain of  $\alpha$ His89(FG1) is displaced by  $\sim 13$  Å from a position facing away from the haem group on the outside of helix F to an inwards-facing position, where it forms a hydrogen bond to the propionate at the  $\beta$ -pyrrolic position 6 (Fischer & Orth, 1934) of the haem group (Figs. 1d and 4). Despite these changes,  $\alpha$ His87(F8) retains

Table 3 Ouaternary-structure comparisons between PDB entry 4xs0 and reference Hbs.								
Hb structure <sup>†</sup>	PDB code	$\alpha 1\beta 1$ r.m.s.d. (Å), all atom pairs ( $C^{\alpha}$ atom pairs)‡	$\alpha 2\beta 2$ r.m.s.d (Å), all atom pairs ( $C^{\alpha}$ atom pairs)‡	Rotation§ (°)	Translation§ (Å)			
T state	2dn2	1.7 (1.0)	4.1 (3.9)	9.2	0.7			
R state	2dn1	1.9 (1.1)	7.4 (7.2)	17.0	4.2			
R2 state	1bbb	2.3 (1.4)	11.5 (11.4)	27.9	6.0			
RR2 state	1mko	2.1 (1.4)	9.7 (9.5)	22.1	6.2			
R3 state	1yzi	1.9 (1.3)	9.0 (8.9)	20.1	4.4			
Human $\zeta_2 \beta_2^{\rm S}$	3w4u	1.5 (1.0)	1.8 (1.4)	3.5	0.6			
Cat $\alpha_2\beta_2$	3gqp	1.5 (1.0)	1.5 (1.0)	1.1	0.1			

† The T-state and R-state reference structures are human deoxy-Hb and HbCO from Park *et al.* (2006), respectively, the R2 structure is from Silva *et al.* (1992), the RR2 and R3 structures are from Safo & Abraham (2005), the human variant Hb comprising ζHb and  $\beta^8$  (carrying the sickle-cell mutation) is from Safo *et al.* (2013) and the low-affinity Hb from *F. silvestris catus* is from Balasubramanian *et al.* (2014). ‡ Comparison with the 4xs0 structure after superposing  $\alpha 1\beta 1$  dimers using residues of the BGH frame. § Rotation/translation needed to align  $\alpha 2\beta 2$  of the test structure with  $\alpha 2\beta 2$  of the 4xs0 structure after first superposing the  $\alpha 1\beta 1$  dimers according to the BGH frame.

coordination of the haem iron. The hydrophobic side chains of  $\alpha$ Leu86(F7) and  $\alpha$ Leu91(FG3), which in the native structure sit under the porphyrin *meso*-position  $\gamma$  and pyrrole ring III, respectively (Fig. 4*d*), are rotated outwards (Fig. 4*e*). The positions vacated by Leu86 and Leu91 are partially filled by the  $\alpha$ His87(F8) main chain and  $\alpha$ Ala88(F9). Other side chains in the haem pocket retain their native positions (r.m.s.d. of 0.32 Å over 105 C<sup> $\alpha$ </sup> atom pairs with PDB entry 3p5q, excluding residues  $\alpha$ Val73– $\alpha$ Arg92).

The tertiary-structural changes seen in Hb  $\alpha$  suggest three factors that might contribute to haem release. Firstly, disturbance of side-chain packing in the proximal haem pocket could lead to an increased probability of water entry. Water promotes autooxidation (Brantley et al., 1993) and competes with the proximal His as a ligand for iron(III) haem, leading to accelerated rates of haem loss (Liong et al., 2001). To prevent this, the  $\alpha$  and  $\beta$  subunits share a common feature of six bulky hydrophobic side chains that form a hydrophobic 'cage' shielding the iron-His(F8) bond from water; in  $\alpha$  these residues are aLeu83(F4), aLeu86(F7), aLeu91(FG3),  $\alpha$ Val93(FG5),  $\alpha$ Phe98(G5) and  $\alpha$ Leu136(H19) (Fig. 4d). Movement of the  $\alpha$ Leu86(F7) and  $\alpha$ Leu91(FG3) side chains out of the haem pocket may have a significant effect on haem loss through permitting the ingress of water molecules (Fig. 4e). Secondly, unwinding of helix F may result in loss of a physical barrier to haem movement; the  $\alpha$  and  $\beta$  haem groups are embedded deep between the E and F helices and make van der Waals contacts over an extensive surface. Consequently, haem binding and dissociation is thought to occur in conjunction with local protein folding/unfolding. In apo myoglobin the F helix and the adjacent EF and FG corners undergo local unfolding on an approximately microsecond time scale to allow haem entry (Hughson et al., 1990; Cocco & Lecomte, 1990; Lecomte & Cocco, 1990; Eliezer & Wright, 1996), and similar unfolding is presumably required during haem loss. In the case of Hb,  $\alpha\beta$  dimers lacking one haem group have reduced  $\alpha$ -helical content, suggesting that haem binding/extraction involves localized folding/unfolding (Kawamura-Konishi et al., 1992). In the IsdH<sup>N2N3</sup>-metHb crystal, the  $\alpha$  F helix, although not disordered, adopts some nonregular secondary structure, with consequent loss of the cooperative hydrogenbonding network that confers comparative rigidity to the native  $\alpha$ -helical structure. Thirdly, alternative His/Tyr ligands may compete with  $\alpha$ His87(F8) for haem binding. A number of side chains with haem-binding capability lie at positions between the haem-coordinating side chains of Hb  $\alpha$  and the IsdH<sup>N3</sup> domain, namely  $\alpha$ His58(E7), aHis89(FG1), aHis45(CE3) and IsdH Tyr646 (Fig. 4b). The position of  $\alpha$ His89 is particularly

intriguing, as this residue has undergone the largest displacement from its native location and sits with its  $N^{\epsilon 2}$  atom ~4.5 Å away from  $N^{\epsilon 2}$  of  $\alpha$ His45, leading us to speculate that these two side chains could potentially form a hexacoordinate haem as an intermediate during haem transfer. A potential role for intermediate haem ligands needs to be investigated in future studies of haem transfer.

Finally, it is not clear how contacts between IsdH and Hb  $\alpha$  lead to local conformational changes in the globin, but we speculate that the *S. aureus* bacterium is exploiting an intrinsic feature of the F helix that makes it susceptible to strain. A similar effect has been observed during interaction of  $\alpha$  with the chaperone  $\alpha$ -Hb stabilizing protein (AHSP): AHSP binding to  $\alpha$  at a distant site induces a localized disturbance in the F helix that precedes a larger conformational rearrangement of  $\alpha$  (Dickson *et al.*, 2013).

#### 3.6. Altered Hb quaternary structure

Using mass spectrometry, Spirig and coworkers have shown that IsdH<sup>N2N3</sup> induces a shift from tetrameric to dimeric Hb species (Spirig et al., 2013). Dissociation of Hb tetramers into  $\alpha\beta$  dimers accelerates autooxidation (Zhang *et al.*, 1991) and haem dissociation from Hb chains (Hargrove et al., 1997), suggesting that IsdH may exploit this mechanism to promote haem release. Following erythrocyte lysis in vivo,  $\alpha\beta$  dimers are formed naturally by the dilution of Hb tetramers and are scavenged by the host serum protein haptoglobin. IsdH can bind haptoglobin-Hb complexes (Visai et al., 2009; Dryla et al., 2003, 2007; Pilpa et al., 2009), although haem uptake from this complex has not been demonstrated. However, certain conditions, such as when the haptoglobin capacity is exceeded, may bring S. aureus into contact with the Hb tetramer state and the influence of IsdH on Hb quaternary structure is therefore of interest.

In native adult human Hb the  $\alpha 1\beta^2$  interface undergoes a reorganization that is coupled to oxygen binding, involving a  $\sim 15^{\circ}$  rotation and  $\sim 1$  Å translation of the  $\alpha 2\beta^2$  dimer relative to  $\alpha 1\beta^1$  (Baldwin & Chothia, 1979) that brings the two  $\beta$  subunits closer together during the T $\rightarrow$ R transition

Table 4Iron-iron distances (Å).

Hb structure	PDB code	$\alpha 1\beta 1$	α1β2	α1α2	$\beta 1 \beta 2$
T state	2dn2/1hgb	36.4/36.1	24.3/24.4	34.1/34.4	39.5/39.7
R state	2dn1/3p5q	34.8/35.0	25.5/25.4	34.1/34.5	34.9/34.9
Human $\zeta_2 \beta_2^{\rm S}$	3w4u	34.9	24.6	34.1	38.7
Cat $\alpha_2 \beta_2$	3gqp	34.4	24.4	32.7	38.6
IsdH <sup>N2N3</sup> -metHb	4xs0	34.5	24.7	33.9	38.3

(Muirhead & Perutz, 1963). In addition to the R structure, fully liganded human Hb has been crystallized in a number of other quaternary structures, including the R2 (Silva *et al.*, 1992), RR2 and R3 states (Safo & Abraham, 2005). To determine the similarity of IsdH<sup>N2N3</sup>–metHb to the T, R, R2, RR2 and R3 quaternary structures, we superposed  $\alpha 1\beta 1$ dimers over a region of invariant structure (the 'BGH' frame established by Baldwin & Chothia, 1979) and determined the r.m.s.d. for the non-aligned  $\alpha 2\beta 2$  dimer (Table 3). We then determined the rotation/translation required to align the  $\alpha 2\beta 2$ dimers for each of the T, R, R2, RR2 and R3 structures (Table 3). These comparisons showed that of these five quaternary structures, IsdH<sup>N2N3</sup>-metHb was most similar to the T state  $(\alpha 2\beta 2 \text{ r.m.s.d.} \text{ of } 3.9 \text{ Å}; \alpha 2\beta 2 \text{ rotation/translation of } 9.2^{\circ}/$  0.7 Å); however, IsdH<sup>N2N3</sup>-metHb was clearly different from all of the test structures. The iron-iron distance between  $\beta$ haems of IsdH<sup>N2N3</sup>-metHb (Table 4; 38.3 Å) also indicated closer similarity to the T state (39.5 Å) than to the R state (34.9 Å). Using the same criteria (Tables 3 and 4), two other Hb structures in the PDB show much greater levels of similarity to IsdH<sup>N2N3</sup>-metHb, namely a human Hb variant comprising embryonic  $\zeta$  chain and  $\beta$  chain carrying the sicklecell mutation ( $\beta^{S}$ ) crystallized in the fully CO-ligated form (Safo et al., 2013) and metHb from cat (Felis silvestris catus; Balasubramanian *et al.*, 2014). Both the Hb  $\zeta_2 \beta_2^{\rm S}$  and cat metHb structures exhibit an unusual T state-like quaternary structure with several local tertiary and quaternary features characteristic of the R state (Safo et al., 2013; Balasubramanian et al., 2014).

The presence of specific side-chain interactions that characterize the T and R quaternary states (Baldwin & Chothia, 1979), and the functional importance of C-terminal salt bridges of the T state (Perutz, 1970), were established in early



#### Figure 5

Hb quaternary features of the IsdH<sup>N2N3</sup>-metHb complex (PDB entry 4xs0; yellow), O<sub>2</sub>-ligated R-state Hb (PDB entry 2dn1; red), deoxy T-state Hb (PDB entry 2dn2; blue), CO-ligated  $\zeta_2 \beta_2^S$  (PDB entry 3w4u; cyan) and cat metHb (PDB entry 3gqp; green). (*a*) Detail of the switch region of the  $\alpha 1\beta 2$  interface, showing the relative positions of the  $\alpha 1$  subunit helix C and the  $\beta 2$  subunit FG corner. (*b*) Detail of the joint region of the  $\alpha 1\beta 2$  interface, showing the relative positions of the  $\beta 2$  helix C and the  $\alpha 1$  FG corner. In the R and T states the side chain of  $\beta 2$ Trp37 contacts the side chain of  $\alpha 1$ Pro95. In the IsdH<sup>N2N3</sup>-metHb complex, the side chain of  $\alpha 1$ Arg92 separates  $\beta 2$ Trp37 and  $\alpha 1$ Pro95 and makes salt bridges with  $\alpha 1$ Asp94 and the terminal carboxyl of the  $\alpha 1$  subunit (van der Waals contacts are represented by dashed green lines; hydrogen-bonding interactions are represented by dashed magenta lines). (*c*) Comparison of the joint region of the  $\alpha 1\beta 2$  interface from IsdH<sup>N2N3</sup>-metHb.

#### Table 5

Hydrogen-bond and salt-bridge interactions characteristic of Hb quaternary states.

Values in parentheses are for the pseudo-symmetrical contact where there are two  $\alpha\beta$  dimers in the crystallographic asymmetric unit.

Residue 1	Residue 2	T deoxyHb (2dn2)	T aquometHb (1hgb)	$\begin{array}{c} R\\ HbO_2\\ (2dn1) \end{array}$	R aquometHb (3p5q)	R2 HbCO (1bbb)	RR2 HbCO (1mko)	R3 HbCO (1yzi)	$\zeta_2 \beta_2^{S}$ HbCO (3w4u)	Cat aquometHb (3gqp)	IsdH <sup>N2N3</sup> – aquometHb (4xs0)
Hydrogen bonds at 1	the $\alpha 1\beta 2$ interface										
β2Asp99 OD1/2	α1Tyr42 OH	2.5 (2.5)	2.5 (2.5)	_	_	_	_	_	2.4 (2.4)	2.8 (2.5)	2.5
β2Asp99 OD1/2	α1Asn97 ND2	2.8 (2.8)	3.9 (3.1)	_	_	_	_	_	2.9 (2.8)	2.8 (2.7)	2.7
α1Asp94 OD1/2	β2Trp37 NE1	2.9 (2.8)	3.2 (3.1)	3.7	3.7	3.8 (3.8)	3.6 (3.4)	3.5		_ ` `	_
α1Asp94 OD1/2	β2Asn102 ND2	_ ` `	_ ` `	2.8	2.8	2.7 (2.8)	2.8 (2.8)	2.8	_	_	_
β2Trp37 NE1	β2Asn102 OD1	_	_	2.9	3.0	3.0 (3.0)	3.1 (3.1)	3.3	3.1 (3.2)	(2.9)	3.5
C-terminal salt bridg	ges										
α1Arg141 NH1/2	α2Asp126 OD2	2.8 (2.7)	2.5 (2.4)	_	Disordered	_	_	_	_	(2.9)	2.9
α1Arg141 OXT	α2Lys127 NZ	2.8 (2.8)	2.6 (2.8)	_	Disordered	_	_	_	_	2.9	_
β2His146 NE2	$\beta$ 2Asp94 OD(1)	2.6 (2.8)	2.8 (2.6)	_	_	_	_	_	3.0 (2.6)	2.9 (3.0)	3.6
$\beta$ 2His146 OXT	α1Lys40 NZ	2.8 (2.6)	2.4 (2.7)	-	_	-	-	-	2.9 (2.7)	2.6 (2.5)	2.7

Table 6

Distances between  $C^{\alpha}$  atoms of selected residues across the switch and joint regions of the  $\alpha\beta$  dimer interface.

Values in parentheses are for the pseudo-symmetrical contact where there are two  $\alpha\beta$  dimers in the crystallographic asymmetric unit.

Residue 1	Residue 2	T deoxyHb (2dn2)	R HbO <sub>2</sub> (2dn1)	$\zeta_2 \beta_2^{\rm S}$ HbCO (3w4u)	Cat aquometHb (3gqp)	IsdH <sup>N2N3</sup> -aquometHb (4xs0)
α1Pro95(FG7)	β1Pro100(FG7)	21.6 (21.9)	18.9	19.7 (19.8)	18.6 (19.2)	19.1
α1Pro95(FG7)	β2Trp37(C3)	9.5 (9.4)	8.4	12.9 (13.1)	13.8 (13.0)	13.2
$\beta$ 1Pro100(FG7)	α2Thr38(C3)†	7.1 (7.2)	10.0	7.0 (7.0)	7.4 (7.0)	6.9
α2Thr38(C3)†	β2Trp37(C3)	29.0 (29.0)	29.1	29.9 (29.8)	28.9 (29.1)	29.2

† Replaced by α2Gln38(C3) in PDB entry 3w4u.

seminal works and provide a useful and robust definition of quaternary structure (Park *et al.*, 2006). For example, the hydrogen bonds that are characteristic of the R state are largely preserved in the R2, RR2 and R3 structures (Table 5), despite relative rotations of the  $\alpha\beta$  dimers of up to 12°. This pattern is consistent with NMR measurements made in solution, which indicate that CO-ligated Hb populates multiple low-energy states that average to a structure intermediate between the R and R2 conformations (Lukin *et al.*, 2003), and a sliding motion at the  $\alpha\beta$  dimer interface that does not



Figure 6

Schematic of one half of the symmetrical  $\alpha 1\beta 2$  interface, showing only the  $C^{\alpha}$  atoms of four residues to represent the switch and joint contacts, in the IsdH<sup>N2N3</sup>-metHb complex (PDB entry 4xs0; yellow), O<sub>2</sub>-ligated R-state Hb (PDB entry 2dn1; red) and deoxy T-state Hb (PDB entry 2dn2; blue), prepared by superposition of the  $\alpha 1\beta 1$  dimers in the BGH frame (Baldwin & Chothia, 1979). Dashed lines indicate approximately equal interatomic distances in the three structures (Table 6). Arrowheads indicate displacement of R-state atoms (red) or IsdH<sup>N2N3</sup>-metHb complex atoms (yellow) relative to the T-state structure (blue).

require substantial changes to subunit contacts has been proposed (Mueser *et al.*, 2000). The  $\alpha 1\beta 2$  interface in IsdH<sup>N2N3</sup>-metHb displays a number of hydrogen bonds and C-terminal salt bridges that are characteristic of the quaternary T state; however, the intrasubunit hydrogen bond between  $\beta$ Trp37 and  $\beta$ Asn102 is a tertiary R-state feature and the loss of a salt bridge to the terminal carboxyl of  $\alpha$ Arg141 is atypical of a T-state structure.

In addition to electrostatic interactions, steric interactions at the  $\alpha\beta$  dimer interface also characterize the T and R states (Baldwin & Chothia, 1979). The most extensive area of contact is the  $\alpha 1\beta 2$  interface, which includes interactions between helix C of the  $\alpha$ 1 subunit and the FG corner of the  $\beta$ 2 subunit (the 'switch' region) and between the FG corner of  $\alpha 1$ and helix C of  $\beta 2$  (the flexible 'joint') (Perutz *et al.*, 1998). In the  $T \rightarrow R$  quaternary transition the switch moves between two positions characterized by different inter-chain contacts, whereas the joint maintains similar side-chain packing in the T and R states. In the IsdH<sup>N2N3</sup>-metHb crystal the switch region is essentially identical to that of the T state (Fig. 5a). In contrast, the joint region in the IsdH<sup>N2N3</sup>-metHb structure has undergone a significant rearrangement (Fig. 5b). Thus, in normal R and T states the side chain of  $\beta$ Trp37 is packed against  $\alpha$ Asp94 and  $\alpha$ Pro95 (Fig. 5b; red and blue shading). In IsdH<sup>N2N3</sup>-metHb the side chain of  $\alpha$ Arg92 projects between  $\beta$ Trp37 and  $\alpha$ Pro95, disrupting the joint region, and makes salt-bridge interactions with the side chain of  $\alpha$ Asp94 and the carboxyl-terminus of the  $\alpha$  subunit. Interestingly, this highly unusual structure of the joint region is also observed in HbCO  $\zeta_2 \beta_2^{\rm S}$  and cat metHb, although in the former it is the side chain

of  $\alpha$ Arg141, rather than  $\alpha$ Arg92, that projects between  $\beta$ Trp37 and  $\alpha$ Pro95 (Fig. 5*c*).

The changes seen at the joint contact provide an explanation for how R-like tertiary features are accommodated in the T-like quaternary structure. In native human Hb, the tertiarystructural changes in the globin subunits associated with ligand binding cause the FG corners of the  $\alpha 1$  and  $\beta 1$  subunits to move approximately 2.5 Å closer together in the  $T \rightarrow R$ transition (Perutz et al., 1998), and this is accommodated by movement of the  $\beta$ 1 FG corner relative to the  $\alpha$ 1 helix C at the switch region (Fig. 6; red arrows). In the IsdH<sup>N2N3</sup>-metHb structure the distance between the  $\alpha 1$  and  $\beta 1$  FG corners is characteristic of the liganded R state (Table 6) and this is accompanied by opening of the joint region (Fig. 6; yellow arrows), whilst the switch region maintains the canonical T structure. The relative positions of switch and joint secondarystructure elements are essentially the same in IsdH<sup>N2N3</sup>metHb, cat metHb and HbCO  $\zeta_2 \beta_2^{\rm S}$  (Table 6). In the case of IsdH<sup>N2N3</sup>-metHb, the cause of the altered interactions seems likely to be the altered conformation of the  $\alpha$  F helix and FG corner induced by Hb receptor binding. Cat Hb has low oxygen affinity, even in the absence of 2,3-diphosphoglycerate (DPG; Balasubramanian *et al.*, 2014), whereas Hb  $\zeta_2 \beta_2^{\rm S}$  is a high-affinity variant (Safo et al., 2013); thus, the contribution of the T-like quaternary structure to oxygen affinity is not clear in the background of amino-acid sequence differences between Hb  $\alpha$ , Hb  $\zeta$  and cat Hb. IsdH<sup>N2N3</sup> induces a shift from tetrameric to dimeric Hb species (Spirig et al., 2013), and it is possible that the altered  $\alpha\beta$  dimer interface observed in the crystal structure is the underlying mechanism.

In conclusion, the IsdH<sup>N2N3</sup>–Hb complex shows substantial deformation of the  $\alpha$ Hb haem pocket that is predicted to weaken the globin–haem interaction and disrupt the allosteric  $\alpha 1\beta 2$  interface, potentially leading to weakening of the Hb quaternary structure.

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