Structure and dynamics of Type III periplasmic proteins VcFhuD and VcHutB reveal molecular basis of their distinctive ligand binding properties

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Molecular mechanisms of xenosiderophore and heme acquisitions using periplasmic binding protein (PBP) dependent ATP-binding cassette transporters to scavenge the essential nutrient iron are elusive yet in *Vibrio cholerae*. Our current study delineates the structures, dynamics and ligand binding properties of two Type III PBPs of *V. cholerae*, VcFhuD and VcHutB. Through crystal structures and fluorescence quenching studies we demonstrate unique features of VcFhuD to bind both hydroxamate and catecholate type xenosiderophores. Like *E. coli* FhuD, VcFhuD binds ferrichrome and ferri-desferal using conserved Tryptophans and R102. However, unlike EcFhuD, slightly basic ligand binding pocket of VcFhuD could favour ferri-enterobactin binding with plausible participation of R203, along with R102, like it happens in catecholate binding PBPs. Structural studies coupled with spectrophotometric and native PAGE analysis indicated parallel binding of two heme molecules to VcHutB in a pH dependent manner, while mutational analysis established the relative importance of Y65 and H164 in heme binding. MD simulation studies exhibited an unforeseen inter-lobe swinging motion in Type III PBPs, magnitude of which is inversely related to the packing of the linker helix with its neighboring helices. Small inter-lobe movement in VcFhuD or dramatic twisting in VcHutB is found to influence ligand binding.

Iron is an essential nutrient that pathogenic bacteria acquire to support the function of a number of enzymes crucial for survival, such as ribonucleotide reductase for the synthesis of DNA precursors, cytochromes for electron transport or tricarboxylic acid (TCA) cycle enzymes for energy production1. Since the concentration of free iron in tissues is extremely low (10−18 M)2, bacteria use various strategies to acquire Fe3+. Many infectious diseases, such as gonorrhea, malaria, tuberculosis and diarrheal infections depend on the expression of microbial acquisition mechanisms capable of competing with the host’s iron scavenging systems. The main strategies for iron acquisition include direct extraction of Fe3+ from specific iron-containing complexes synthesized by their hosts, such as lactoferrin, transferrin, haemoglobin or hemin3,4 and/or production of siderophores, small Fe3+ -chelating molecules with high affinity and selectivity for Fe3+ ion5,6. The mechanisms of such ‘iron thievery’ of pathogenic bacteria have drawn special attention in recent years since these are being exploited for ‘Trojan Horse’ mechanism of antibiotic delivery to reduce permeability-mediated drug resistance7.

The iron uptake mechanism of *Vibrio cholerae*, the causative agent of diarrheal disease cholera, is currently insufficiently understood. Among the two major classes of siderophores, hydroxamates and catecholates, *V. cholerae* acquires iron through an indigenous catecholate siderophore, vibriobactin8. Some strains of non-epidemic *V. cholerae* responsible for wound infections and septicemia in susceptible people obtain enough iron from blood using iron acquisition systems other than vibriobactin9. Additionally, both vibriobactin synthesis and transport

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antibiotic albomycin2,18. Interestingly, despite a high degree of structural conservation, PBPs such as hydroxamate siderophores such as ferrichrome, desferal, fungal siderophore coprogen and the structurally related of Type III PBPs that lead to efficient and diverse ligand binding.

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tinctive features of heme binding by identifying the differences in binding mechanism of two heme molecules by studies through native PAGE and spectrophotometry on the wild type and mutant proteins have illuminated dis- implication of which has been addressed here. Structure of apo-

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Types I and II PBPs have three and two inter-lobal β-strands or extended elements respectively, Type III PBPs are characterized by a single α-helical linker connecting the N- and C-terminal lobes15. Mostly, the siderophore and heme binding PBPs belong to Type III. Unlike Type I and II PBPs, which undergo large "Venus flytrap" conformational changes upon ligand binding, Type III PBPs are known as relatively more rigid molecules, although in some proteins like FepB of E. coli or HtsA of Gram-positive bacteria Staphylococcus aureus, ligand binding is accompanied by significant loop movements16,17. FhuD of Escherichia coli (EcFhuD), another Type III PBP, binds hydroxamate siderophores such as ferrichrome, desferal, fungal siderophore coprogen and the structurally related antibiotic albomycin18,19. Interestingly, despite a high degree of structural conservation, PBPs such as EcFhuD, YfY of Bacillus cereus or FhuD2 of Staphylococcus aureus have evolved with critical siderophore binding residues on either the C-lobe, N-lobe or both19.

Available structural evidences on heme binding PBPs, on the other hand, suggest variable mode of heme binding in different bacteria. While ShuT of Shigella dysenteriae (SdShuT) and PhuT of Pseudomonas aerugi-

nos (PaPhuT) share common architecture and bind one heme molecule in a narrow cleft between the N- and C-terminal lobes, HmuT of Yersinia pestis (YPHmuT), having similar overall structure, contains a significantly wider central cleft capable of accommodating two stacked heme molecules20,21.

In V. cholerae, the molecular details of siderophore binding by the PBPs are restricted to the structures of catecholate specific ViuP22 and VctP23 while mechanisms of hydroxamate siderophores or heme binding are unknown yet. Here we report the structures, dynamics and ligand binding properties of (i) a putative ferrichrome binding PBP, VcFhuD (Accession code: A0A0H6E7H0) that belongs to the ABC transporter genes fhuBCD24 and (ii) the heme binding PBP, VcHutB (Accession code: A5F05S) belonging to hutBCD of V. cholerae genome25. The crystal structures of VcFhuD and VcHutB, together with Molecular Dynamics (MD) simulations and biochemical analysis, provide first quantitative overview of xenosiderophores and heme binding by the PBPs in V. cholerae. VcFhuD has shown a unique property of binding both hydroxamate and catecholate siderophores, mechanistic implication of which has been addressed here. Structure of apo-VcHutB coupled with heme-protein interaction studies through native PAGE and spectrophotometry on the wild type and mutant proteins have illuminated distinctive features of heme binding by identifying the differences in binding mechanism of two heme molecules by VcHutB and YPHmuT. Additionally, MD simulation results have identified distinguishing inter-lobe movements of Type III PBPs that lead to efficient and diverse ligand binding.

Results

Growth revival of V. cholerae O395 by exogenous siderophores and heme. We have investigated the capability of V. cholerae O395 strain to utilize Fe²⁺ bound to the siderophores and heme using growth assay. For this purpose, we have used EDTA (ethylene diamine tertaacetic acid), the versatile chelator that binds both Fe⁴⁺ and Fe²⁺ ions. 150 μg of EDTA per ml at 1 × 10⁴ bacteria per ml caused complete inhibition of growth of V. cholerae (Fig. 1a). Precise zones of growth were, however, observed around the disks soaked with heme, ferrichrome and Fe²⁺ treated deferoxamine mesylate (ferri-desferal) in a metal ion depleted plate which establish that these iron compounds can efficiently be used by V. cholerae O395 strain to restore growth (Fig. 1b).

Figure 1. (a) Effect of EDTA on growth of V. cholerae O395 strain; (b) Distinct zones of growth observed around filter paper disks soaked with (1) heme, (2) ferrichrome and (3) ferri-desferal in the presence of EDTA.
Overall structure of VcFhuD in apo and ferri-Desferal bound states. VcFhuD shares 30% sequence identity with its nearest structural homolog EcFhuD where the hydroxamate siderophore binding residues are found to be conserved (Fig. 2a). Tris-catecholate siderophore binding protein FeuA from Bacillus subtilis, YfiY from Bacillus cereus and BsFeuA from B. subtilis show about 27% sequence identity with VcFhuD. However, overall sequence identity among these PBPs is as low as 3.6% (Fig. 2a). Interestingly, vibriobactin binding PBP ViuP of V. cholerae exhibits only 15% identity with VcFhuD.

We have solved the crystal structure of VcFhuD in apo and ferri-desferal bound states. Two polypeptide chains of the asymmetric unit of apo-VcFhuD are almost identical with an rmsd of 0.3 Å. VcFhuD has a typical bi-lobal structure where the two lobes are connected by a 25-residue linker helix designated as α6 (Fig. 2b). The N-terminal lobe is comprised of a twisted five-stranded parallel β-sheet connected by the α-helices whereas the

![Figure 2.](image-url)
C-terminal lobe has a mixed β-sheet surrounded by the α-helices (Fig. 2b). Helix at the end of N-terminal lobe (α5) and the extreme C-terminal helix (α12) provide stability to the linker helix through hydrophobic and polar interactions. The siderophore binding site is located in the shallow cleft between the N- and C-terminal lobes (Fig. 2b).

Superposition of holo-VcFhuD on the apo structure produces an rmsd of 0.4 Å and no significant change, either in the overall structure or at the ligand binding pocket, is observed between the two. The conformation of the ligand binding residues almost remains unaltered upon ferri-desferal binding, as observed in Fig. 2c. All four molecules of holo-VcFhuD of the asymmetric unit show unambiguous electron density of the ligand, ferri-desferal (Fig. S1). Similar to ferri-desferal bound EcFhuD structure, the mesylate (OSO₂CH₃) portion of the siderophore could not be located in the electron density probably because of its disordered nature.

Properties of ligand binding site and ferri-desferal binding. The ligand binding cleft of VcFhuD is primarily hydrophobic and made of W61, W86 of the N-lobe helices, W239 and W293 from the loop regions of the C-lobe and F211 and I241 of the C-lobe helices (Fig. 2a,c). R102 remains pre-oriented for the ligand binding by hydrophobic packing with W61 (Fig. 2c) and the salt bridge interactions with E79 (Fig. 2d). Corresponding residue, R84 of EcFhuD, is found to be salt bridged with E42. The ferrioxamine portion of ferri-desferal snugly fits in the binding cleft of VcFhuD (Fig. 2d). Two hydroxamate carbonyl oxygens of ferri-desferal are hydrogen bonded by R102 (Fig. 2c). The other hydroxamate carbonyl oxygen solely serves the purpose of coordinating Fe³⁺ of ferri-desferal. Y295 forms hydrogen bond with the backbone of ferri-desferal (Fig. 2c).

Ferri-desferal bound VcFhuD superposes on its E. coli counterpart (PDB code: 1K2V) with an rmsd of 1.6 Å. Although ferri-desferal binding residues are primarily conserved in these two PBPs, the loops surrounding the binding site differ significantly (Fig. 2d). In VcFhuD, the loop that connects α5 with the N-lobe is more inclined toward the ligand binding pocket by ~8 Å (Fig. 2d). As a result, the loop of the C-lobe that possesses W239 experiences a lateral shift towards the pocket and the ferri-desferal bound to VcFhuD is shifted outward compared to that bound to EcFhuD (Fig. 2d). However, because of similar lateral shift of the crucial residues, which are responsible for polar interactions and hydrophobic packing with ferri-desferal, the binding of this ligand to VcFhuD is not compromised (Fig. 2d). The contribution made by Y106 of EcFhuD in ferri-desferal binding is not expected in VcFhuD since P123 occupies the corresponding position of the later (Fig. 2d).

Important variations in surface electrostatics around the binding pocket are observed between EcFhuD and VcFhuD (Fig. 2e). The charge of the residues surrounding the binding site of EcFhuD is predominantly negative, making it unsuitable for binding the negatively charged catecholate siderophores²⁻¹⁸. Some of such negatively charged residues of EcFhuD namely, E42, E213, T181, D239, D241 are replaced by non-polar, zwitterionic and basic residues in VcFhuD (Fig. 2a). This, together with the adjacent shorter and non-polar loop makes the binding site of VcFhuD apolar with slightly basic nature and accessible both for the catecholate and hydroxamates siderophores (Fig. 2d.e).

Ligand binding affinity of VcFhuD. We have calculated the binding affinity of VcFhuD to (i) Fe³⁺ treated ferrichrome, (ii) Fe³⁺ treated deferoxamine mesylate and (iii) Fe³⁺ treated enterobactin by measuring the intrinsic fluorescence quenching (Δλexc = 280 nm, Δλem = 295–410 nm), accounting the contributions of tryptophans and tyrosines present at the ligand binding site. Ferrichrome and ferri-desferal have shown significant binding with VcFhuD with the dissociation constant (Kd) values of 1 ± 0.226 μM and 0.6 ± 0.015 μM respectively (Fig. 3a,b). Interestingly, considerable interaction is also observed between catecholate siderophore ferri-enterobactin and VcFhuD. Since binding of ferri-enterobactin to a hydroxamate siderophore specific PBP is an outstanding observation, we have verified this binding through near-UV CD. The gradual increase of the positive band at 268 nm with increasing amount of the ferri-enterobactin, keeping protein amount constant, indicates interactions between the two (Fig. S2). However, for quantitative measurements of affinity, fluorescence quenching data were used which produce a Kd of 5.5 ± 0.414 μM (Fig. 3c). Our results indicate that VcFhuD has the unique property of binding both hydroxamate and catecholate siderophores.

Structural uniqueness of VcHutB. Heme uptake from the host is another important aspect that contributes significantly to iron acquisition mechanisms of bacteria. VcHutB has been identified as the exclusive heme binding PBP of V. cholerae that shares 32%, 34% and 37% sequence identity with the other structurally characterized heme binding PBPs, SdShuT, PaPfuT and YpHmuT respectively (Fig. 4a).

The structure of apo-VcHutB has been determined up to 2.4 Å resolution. Although VcHutB has shown highest identity with YpHmuT (Fig. 4a), only apo-SdShuT (PDB code: 2RG7)⁰° produced an acceptable MR solution for VcHutB. Apo-VcHutB superposes on apo-SdShuT and apo-YpHmuT with an rmsd of 1.9 Å and 2.1 Å respectively showing significant differences in the loop regions around the ligand binding site (Fig. 4b,c). SdShuT bind one heme molecule in a narrow cleft using Y67⁰⁰. The wider cleft of YpHmuT, on the other hand, accommodates two heme molecules stacked in an anti-parallel fashion where Y70 and H167 anchor Fe³⁺ of two separate heme molecules. VcHutB also contains two potential heme binding residues Y65 and H164, located likewise on either side of the cleft with suitable micro-environment to bind heme (Fig. 4a,c). Additionally, the adjacent R67 is in a position to stabilize the tyrosinate formed by Y65. The heme binding pocket of VcHutB is even wider than YpHmuT with an average Cβ distance of 20 Å between Y65 and H164 which is 15 Å for YpHmuT.

Structural comparison of YpHmuT and VcHutB suggests that although the ligand binding pocket of VcHutB is suitable to accommodate two heme molecules, the mode of heme binding may differ from YpHmuT. Two loops of VcHutB which possess Y198 and V257 respectively are oriented differently to cause steric clashes if the heme molecules would bind in a manner similar to YpHmuT (Fig. 4d). Being already tightly packed, these loops of VcHutB are apparently incapable of changing conformation to accommodate heme. Our knowledge based docking of two heme molecules in the binding pocket of VcHutB proposes parallel mode of heme binding with...
probable participation of the unique basic residues R68, R167 and K223 which are poised to stabilize the propionate groups (Fig. 4a,e).

**Shift of Soret band of heme upon VcHutB binding.** We attempted to estimate the relative affinity of heme for VcHutB spectrophotometrically by performing hemin titration experiments. The absorption spectrum of hemin features a peak in the Soret band region around 385 nm. Hemin was incrementally titrated to free VcHutB at pH 8.0 and the spectral range between 250–700 nm was measured. VcHutB alone does not show absorption in Soret region but when it was mixed with hemin, the Soret band initially shifted towards right with maxima at 405 nm, followed by a blue-shifted peak at 371 nm, suggesting VcHutB mediated perturbation of the electronic structure of the hemin iron which is indicative of interactions between the two (Fig. 5a). The protein concentration for this experiment was 10 μM. The highest intensity at 405 nm was observed with hemin concentration of 6 μM and the 2nd peak at 371 nm was found to be saturated with 20 μM of hemin (Fig. 5a). The distinct absorption maxima, one at 405 nm and another at 371 nm, point towards two binding events with participation of H164 and Y65, as observed previously in case of YpHmuT21. Change in absorbance at 371 nm versus hemin concentration indicates saturable heme binding to VcHutB (Fig. 5b). Changes in the slope of this graph shows a plateau at the top with two apparent maxima at VcHutB:hemin molar ratios of ~1:1.5 and ~1:2.5 which probably points towards two binding events (Fig. 5b).

**Effect of mutations and pH on the shift of hemin Soret band.** We performed the same experiment with the mutants VcHutB-Y65A and Y65F. Almost no peak shift occurred up to 20 μM of hemin and then gradual red shift took place up to 415 nm with an increasing concentration of hemin up to 40 μM (Fig. 5c,d). No blue shifted peak was observed for either of these mutants. The results indicate that although H164 is capable of recognizing heme, interactions are not efficient enough in the absence of Y65. Furthermore, to investigate the effect
of pH on heme binding, same experiment was carried out with wild type VcHutB at pH 7.0 (Fig. 5e,f). We have obtained similar Soret shift as observed in case of pH 8.0. However, in contrast to pH 8.0 (Fig. 5a,b), the highest intensity at 405 nm was observed here with hemin concentration of 4 μM and a saturation of the 2nd peak at 16 μM. The molar ratio graphs indicate saturable heme binding to VcHutB, where the maxima at pH 7.0 are observed at protein:heme ratio of ~1:1.3 and ~1:1.6 (Fig. 5f). Presumably, incomplete deprotonation of H164 causes weaker heme binding at pH 7.0.

Native PAGE shows binding of two heme molecules to VcHutB. To obtain an affinity estimate, we studied the interactions between VcHutB and hemin by native PAGE, as it was employed previously. Hemin-loaded VcHutB, as judged by visual inspection of Coomassie blue-stained gels, migrates faster in native PAGE than the apo form, reflecting significant compactness of VcHutB upon heme binding, as observed in case of the heme binding PBP, HbpA of Haemophilus influenzae. Experiments were carried out at pH 8.0 with 0.2 mM of VcHutB and a complete shift from apo to holo band occurs with 0.4 mM of hemin further indicating 1:2 binding between VcHutB and heme (Fig. 6a). It was known that as the hemin concentration approaches its Kd value, the protein band splits up, with about half of it migrating faster because of complexation with hemin. Based on densitometric scanning of Coomassie blue-stained gels we have determined the relative amounts of apo-VcHutB as a function of increasing hemin concentrations. By plotting the optical densities of the bands on gel and fitting the data in non-linear curve of GraphPad Prism 7.01, the apparent dissociation constant (Kd) of 200 μM has been calculated for the hemin-VcHutB binding. The Kd value indicates that similar to ShuT/PhuT or HbpA, heme is
not very tightly held by VcHutB. We repeated the same experiments with VcHutB mutants Y65A and Y65F and the results suggest that the compactness acquired upon heme binding is the lowest for Y65A and little higher for Y65F although it is still much lesser than the wild type (Fig. 6b). Native PAGE at pH 7.0 showed a protein:heme stoichiometry that corroborates the observations of spectroscopic studies (Fig. 6c).

**Figure 5.** Hemin binding by VcHutB is observed by UV/VIS spectroscopy. (a) VcHutB at pH 8.0 (no protein in the reference cuvette) was titrated with increasing amounts of hemin. Difference spectra (reference subtracted from sample) are shown for selected heme concentrations (colours as indicated). (b) Saturable heme binding is shown by plotting change in absorbance at 371 nm versus heme concentration measured at pH 8.0. (c,d) Difference spectra for VcHutB-Y65A and VcHutB-Y65F at pH 8.0. (e) Titration of VcHutB at pH 7.0 with increasing amounts of hemin, by same method as of (a). (f) Change in absorbance at 370 nm versus heme concentration indicates saturable heme binding at pH 7.0.

**MD simulation and principal component analysis (PCA) reveal unique inter-domain movement in VcFhuD and VcHutB.** To understand the dynamic nature of VcFhuD and VcHutB, we carried out MD simulations on the apo structure of both the PBPs for 1000 ns. Both VcFhuD and VcHutB are found to be mechanically stable as seen from their RMSD values over the simulation time (Fig. 7a). B-factors, averaged over 1000 ns of MD simulation trajectories demonstrates that while VcFhuD is moderately stable, some of the segments of VcHutB, such as majority of the C-lobe including H164, the loop that connects linker helix with the C-lobe and the loop region adjacent to Y65 show significant dynamics (Fig. 7b). To bring out the collective internal motions in the proteins under consideration, we have carried out Principal Component Analysis (PCA) on the MD
trajectories for up-to 500 ns. For PCA, all the atoms in the protein were considered except the five residues from the terminals due to their excessive motion. The covariance matrix was calculated using the option “covar” and the eigenvectors were analyzed using ‘anaeig’ in Gromacs. Before the covariance matrix was built, the overall translational and rotational motion was eliminated as they are irrelevant for the analysis of the internal motions. The eigenvalues of the covariance matrix were calculated and arranged in order of decreasing value. The motion along the first five eigenvector directions is shown by projecting the trajectory onto these individual eigenvectors against time (Fig. S3). Since the eigenvalues are the average square displacements, the first eigenvectors represent the largest positional deviation, which is pictorially presented in Fig. 7c,d. To date, ‘open’ and ‘closed’ conformations related to the substrate binding mechanism was observed in Type III siderophore binding PBPs like HtsA of S. Aureus or FepB of E. coli16,17. In contrast to that, our results of MD trajectories and resultant PCA revealed a unique twisting movement between the N- and C-lobes, extent of which varies dramatically from one Type III PBP to the other.

Discussion

Blocking and interfering with the iron acquisition systems could disrupt bacterial iron homeostasis and thus suppress bacterial growth. Our observations of growth assay (Fig. 1) established that heme, ferrichrome and ferri-desferal act as efficient iron sources and are utilised by V. cholerae O395 to restore growth. This observation encouraged us to investigate the mechanism of uptake of these iron compounds by VcFhuD and VcHutB inside V. cholerae.

VcFhuD efficiently binds hydroxamate siderophores ferrichrome and ferri-desferal (Fig. 3a,b). Crystal structure showed that VcFhuD possesses a preformed pocket for accommodating hydroxamate siderophores. Binding of ferri-desferal in this pocket is incurred by hydrophobic packing using four conserved tryptophans, W61, W86, W239 and W293, supported by the polar interactions with R102 and Y295 (Fig. 2c). Gallichrome and albomycin binding to EcFhuD using R84 (which corresponds to R102 of VcFhuD) and four conserved tryptophans2,18 suggest that these ligands are expected to fit in the binding pocket of VcFhuD as well. MD simulations and the resultant PCA indicated that although the overall inter-lobe movement is negligible in case of VcFhuD (Fig. 7c), variable local and subtle movements of different crucial amino acids occur here (Fig. 7e). R102 executes minimum fluctuations with retention of the salt bridge with E79 almost in all the frames. On the other hand, variable fluctuations are executed by the tryptophans: while movement is negligible for W61 and W293, relatively higher fluctuations are exhibited by W86 and W239 (Fig. 7e). Presumably, variable ligand binding by VcFhuD is facilitated by the movement of key residues by creating enough room to accommodate different hydroxamate siderophores.
So far, most of the reported PBPs only accept single type (either hydroxamate or catecholate) siderophore. Contrary to that, along with binding the cognate hydroxamate siderophore, VesFhuD also binds catecholate xeno-siderophore ferri-enterobactin with substantial affinity (Kd of 5.5 ± 0.414 μM) (Fig. 3c). The intriguing question raised is if the hydroxamate siderophore binding residues are predominantly conserved in EcFhuD and VesFhuD, then what is making the later capable of binding ferri-enterobactin? Interestingly, unlike EcFhuD, the binding pocket of VesFhuD is primarily made of non-polar and positively charged residues that exert no repulsion to the catecholate siderophores (Fig. 2e). Ferri-enterobactin bound structure of EcFepB shows that along with some hydrophobic residues, three diagonally opposite Arginines critically interact with ferri-enterobactin16. VesFhuD possesses a unique R203 located opposite to R102 (Fig. 7f). Notably, R203 of VesFhuD corresponds to T181 of EcFhuD which is known to add negative charge to the binding site of EcFhuD18. On the other hand, R203 corresponds to R180 of BsFeuA (Fig. 2a) which is seen to play a predominant role in binding ferri-enterobactin (PDB code: 2XUZ)20. MD simulation trajectories showed that although R203 forms hydrophobic lining for
ferri-desferal in the ligand binding pocket of VcFhuD and its terminal basic group hydrogen bonds with the side chain hydroxyl group of S294, R203 has enough potential to orient towards the ligand binding pocket during dynamics. In fact, a concerted movement of W86 creates space for such re-orientation of R203 (Fig. 7e). Docking of ferri-enterobactin in the ligand binding pocket revealed that the non-negative nature of the binding site, together with the contributions of R102 and R203, render VcFhuD suitable for ferri-enterobactin binding (Fig. 7g).

With the rapid rise in bacterial resistance to antibiotics, the cooperative behaviour in microbial communities needs to be urgently understood for the development of novel drugs to control infections caused by resistant bacteria and utilization of ferri-enterobactin by *V. cholerae* as iron source is an important phenomenon in this respect. No PBP was identified in *V. cholerae* so far which is capable of efficiently binding ferri-enterobactin. Therefore, the broader specificities of VcFhuD towards hydroxamate siderophores like ferrichrome and ferri-desferal and the catecholate siderophores ferri-enterobactin, as revealed in our study, may provide *V. cholerae* with competitive advantage for survival. In turn, because of this property, VcFhuD may emerge as a potential drug target.

Additionally, our observations help to better understand the mechanism of variable heme binding by the PBPs in different bacterial species. Binding of two heme molecules with efficient participation of both Y65 and H164 provides compactness to the structure of VcHutB leading to faster migration in native PAGE (Fig. 6). Moreover, heme binding to VcHutB is found to be pH dependent and heme affinity reduces with the decrease in pH from 8.0 to 7.0 (Figs 5 and 6). Such pH dependence may be attributed to the incomplete deprotonation of H164 at low pH causing inefficient heme binding. Appearance of major Soret peak at 405 nm in low heme concentration, which shifts to 371 nm at high heme concentrations, together with the results of native PAGE at different pH, suggest that even though H164 is capable of recognising heme, efficient binding must require Y65. Although binding of two heme molecules in anti-parallel fashion was structurally established in *Y. pestis* HutB, our docking study revealed that the micro-environment of the heme binding site induce in binding of two hemes in parallel fashion to VcHutB with additional participation of the unique basic residues R68, R167 and K223 to stabilize the propionate groups (Fig. 4d).

The physiological significance of binding of two heme molecules *in-vitro* to VcHutB (or to *Yp*HutU) remains to be established. The size of the periplasmic entrance to the translocation pathway of HutUV of *Yersinia pestis*, whose sequence is highly similar to VcHutCD, is narrower than BtuCD of *E. coli*30 in which in turn points towards plausible translocation of one heme molecule by VcHutCD during single reaction cycle. The binding of two heme molecules by VcHutB, may thus be attributed to storage purpose which occurs in a pH and concentration dependent manner. Notably, heme storage is observed in heme degrading protein MhuD of *Mycobacterium tuberculosis* in non-iron depleted situation31.

Most siderophore and heme transporters are Type III PBPs. Although most of the Type III PBPs have a relatively stable bi-lobe arrangement with a linker helix, the sequence diversity within the superfamily is so high that most of the PBPs acquire significantly different non-superimposable structural folds, as observed in case of VcFhuD and VcHutB, the apo structures of which superpose with an rmsd of 15 Å. In contrast to the ‘open’ to ‘close’ lobe movement observed previously for Type III PBPs upon ligand binding, our MD simulation results and the resultant PCA revealed an inter-lobe twisting motion, magnitude of which vary dramatically. VcHutB undergoes dramatic inter-lobe swinging which is more vigorous for the C-lobe. While an ‘in-out’ movement is observed for Y65, movement of H164 seems to be more chaotic (Fig. 7h). During MD simulations, the distance between the Cα atoms of Y65 and H164 varies between 15 to 26 Å with a maximum at 22 Å (Fig. 7i). Variation in proximity of these two crucial residues might be the determinant factor for efficient heme binding. Critical analysis of the simulation trajectories suggests that dynamics of these two Type III PBPs are primarily controlled by the interactions of the linker helix with its two neighbouring helices. Interestingly, the linker helix, along with the two adjacent helices shows maximum sequence variations (Figs 2a and 4a). The interactions among these helices are quite extensive and robust in case of VcFhuD because of which VcFhuD has emerged as a relatively rigid molecule with negligible inter-lobe movement (Fig. 7c). On contrary, the linker helix of VcHutB is one turn shorter than VcFhuD with a highly flexible loop at the C-terminal part of the linker helix (Fig. 7b). Furthermore, in VcHutB, the neighbouring helices are not properly parallel to the linker helix, leading to less extensive packing among them. As a result, VcHutB is more dynamic with drastic inter-lobe twisting and variable proximity between Y65 and H164, which is expected to be an important factor in regulation of heme binding and/or exchanging. Collectively, our current study produces first mechanistic view of xenosiderophores and heme binding by the PBPs in *V. cholerae* including the relationship between the dynamic behaviour of Type III PBPs and ligand binding properties. Further investigations in this direction with more Type III PBPs would also be of interest to enrich the ‘Trojan Horse’ mechanism of drug delivery.

**Methods**

**Growth assay.** Growth assays were performed to test the ability of *V. cholerae* to utilize various compounds as iron sources. Mid-log-phase LB broth culture was plated onto LB agar - streptomycin plate containing 150 μg of EDTA per ml. Filter paper disks (3 mm in diameter) containing 460 μM of hemin chloride, 200 μM of ferri-desferal and 250 μM of ferrichrome were placed on the plates and revival of growth around the disks was examined after 16 h of incubation at 37°C.

**Cloning, overexpression and purification.** Gene sequence of VcFhuD without the signal peptide(N-terminal 54 amino acids) was cloned into kanamycin resistant pET28a+ (Novagen) vector using specific primers (forward primer: 5′-GGAATTCGATATGCAGTGTTCTGAATCTGGGATC-3′, and reverse primer: 5′-CATTGGGATCCCTCATGATGGCAGCCACGCCCAGC-3′). The primers were synthesized (NeoProCell) with Ndel and BamiHI restriction enzyme sites. Chromosomal DNA of *V. cholerae* strain O395,
Isolated using the protocol described in the Molecular Biology Laboratory Manual of UMBC (http://userpages.umbc.edu/~jwollf/methods.html), was used as the template to amplify the region encoding fhuD. The 795 bp fhuD PCR amplicon and the pET28a+ vector with Ndel and BamHI restriction sites were ligated using T4 DNA ligase and the appropriate clones were selected using E. coli XL1-Blue cells with kanamycin resistance. The construct was verified by restriction digestion analysis and commercial DNA sequencing.

Protocol of cloning of VcHutB (residues 24–277 excluding signal peptide; 254 aa) in E. coli XL1-Blue cells in presence of the antibiotic, kanamycin, as a fusion protein having 6x His-tag at the N-terminus has been described in Agarwal et al.25. Matured VcHutD (residues 55–319 excluding signal peptide; 265 aa) and VcHutB were overexpressed in E. coli BL21 (DE3) cells in the presence of antibiotic, kanamycin, by Isopropyl β-D-1-thiogalactopyranoside (IPTG) induction and then purified to homogeneity by Ni-NTA affinity chromatography as per protocol reported in Agarwal et al.25. Mutants of VcHutB, Y65A and Y65F were prepared by two-step PCR. The mutants were verified by commercial sequencing and purifications were done following the same protocol as wild type VcHutB.

Crystallization and diffraction data collection. Crystals of the PBPs were grown at 293 K using the hanging drop vapour diffusion method. Apo- VcHutD crystals were obtained when 2 μl of the protein (45 mg/ml) and 2 μl of precipitant solution consisting of 18% (w/v) PEG 6000, 0.1 M MES (pH 6.0) were equilibrated against a reservoir solution of 34% (w/v) PEG 6000, 0.1 M NaCl and 0.1 M Tris (pH 7.0) for one month. Holo-VcHutD was prepared in a ratio of 1:3 by mixing 2.5 mM of the protein solution with 7.5 mM of ferri-desferal and incubated for 30 minutes in ice. Holo-VcHutD crystals grew similarly with the precipitant solution consisting of 0.8 M ammonium sulfate, 0.1 M HEPES (pH 7.0) after incubation for 20 days against 1.6 M ammonium sulfate, 0.1 M NaCl and 0.1 M Tris (pH 7.0). Crystals were then soaked in cryoprotectant containing 20% (v/v) glycerol, and mother liquor and flash frozen. Crystallization of apo-VcHutB was done as per protocol reported previously25.

Diffraction data of apo and holo VcHutD were collected at 100 K at the PX-BL213 beamline of Indus-2 synchrotron, India with resolutions up to 2.4 Å and 3.4 Å respectively. The data were indexed and integrated using XDS33 and subsequently scaled using AIMLESS from CCP434. Data-collection and processing parameters are given in Table 1.

Phasing and Model Refinement. The initial phases for the PBPs were obtained by molecular replacement (MR) using Phaser35. Since, during phasing experiments of apo-VcHutD no apo structure was available for any such PBP, we used the coordinates of the Gallichrome bound structure of EcFhuD (PDB code: 1EFD) for MR calculations. Two molecules of apo-VcHutD were obtained in the asymmetric unit (with RFZ = 4.4, TFZ = 11.4 and LLG = 144), in the space group P2_12_12_1, using data between 47 Å and 2.5 Å resolutions. Few cycles of refinement by PHENIX36, systematic model building using WinCoot37, gradual inclusion of the solvents individual B-factor and TLS refinement produced a final R cryst of 18.9% (Rmerge = 23.7%) (Table 1). Non-crystallographic symmetry (NCS) was not used during the refinement of apo-VcHutD structure.

Ferri-desferal bound VcHutD was crystallized in the hexagonal space group P3_21 (Table 1). MR with the apo-VcHutD model identified four molecules of ferri-Desferal bound VcHutD in the asymmetric unit. Four folds of NCS were used during refinement. After few cycles of positional refinement and simulated annealing, ferri-desferal was fitted (using the Desferal coordinates of PDB code 1K2V) in the positive density observed in the 2Fobs−Fcalc map at the ligand binding cleft of all four polypeptide chains. The structure of holo-VcHutD was refined well with final R cryst of 19.2% (Rmerge = 22.8%) (Table 1).

Coordinates of apo-SdShuT (PDB code: 2RG7; 20) produced acceptable MR solution for VcHutB. The search model was modified with removal of water molecules, truncation of certain loops around the ligand binding pocket and truncation of the non-identical amino acids to Alanine. One molecule of this modified search model produced the rotation function Z-score (RFZ) of 4.2, translation function Z-score (TFZ) of 10.0 and a convincing log-likelihood gain (LLG) of 88.0 in the space group P4 3 212 using data between 37.0 Å and 2.4 Å resolutions. Refinement was done in a process similar to apo-VcHutD with gradual inclusion of the missing regions. Final refinement produced Rwork and Rfree of 19.7% and 25.5% respectively (Table 1).

Fluorescence quenching studies. Fluorescence measurement was carried out using a spectrophotometer, Hitachi F-7000 following the protocol described in Raines et al.38. Changes in fluorescence of tryptophan and tyrosine residues were measured at an excitation wavelength of 280 nm and the emission spectra were recorded between 295 nm and 410 nm with slit widths of 5 nm for both excitation and emission. All reactions were carried out at 298 K. The reactions were performed in a buffer containing 50 mM Tris-HCl (pH 7.0) and 300 mM NaCl. Equilibrium titration of VcHutD was carried out individually with ferrichrome, ferri-desferal and ferri-enterobactin. Desferal, ferrichrome (iron-free) and enterobactin were purchased from Sigma-Aldrich. Purity was of ≥92.5% (TLC grade), ≥98% (TLC grade) and ≥98% (HPLC grade) for desferal, ferrichrome and enterobactin respectively. 15 mM stock solutions of desferal and ferrichrome were prepared in buffer with composition mentioned above while a 15 mM stock solution of enterobactin was prepared in 90% (v/v) acetonitrile. These ligands were treated with FeCl3 in 1:2 molar ratio to get Fe²⁺ bound ligands. Ferrichrome, ferri-desferal were then diluted to 300 μM in the same buffer while ferri-enterobactin was diluted to 500 μM in 90% (v/v) acetonitrile to use as stock solutions during experiments.

The concentration of VcHutD was 1 μM and ligand concentrations varied from 0 to 231 μM. The dissociation constant, Kd was determined using nonlinear curve fitting analysis as per equations [1] and [2]38. All experimental points for the binding isotherms were fitted by the least-squares method:

\[
K_d = \frac{[C_o] - (\Delta F/\Delta F_{max}) C_p}{C_p} = \frac{(\Delta F/\Delta F_{max}) C_o}{(\Delta F/\Delta F_{max}) C_o + K_d}
\]  

(1)
\[ C_0 \left( \frac{\Delta F}{\Delta F_{\text{max}}} \right)^2 - \left( C_0 + C_p + K_d \Delta F/\Delta F_{\text{max}} \right) + C_p = 0 \]  

(2)

\( C_0 \) and \( C_p \) denotes the input concentrations of the ligands and VcFhuD respectively. \( \Delta F \) is the change in fluorescence intensity at 337 nm (\( \lambda_{\text{ex}} = 280 \) nm) for each point of titration curve and \( \Delta F_{\text{max}} \) is the same parameter when ligand is totally bound to the protein. A double-reciprocal plot of \( 1/\Delta F \) against \( 1/(C_p - C_0) \), as shown in equation [3] was used to determine the \( \Delta F_{\text{max}} \).

\[ 1/\Delta F = 1/\Delta F_{\text{max}} + K_d [\Delta F_{\text{max}}(C_p - C_0)] \]  

(3)

\( \Delta F_{\text{max}} \) was calculated from the slope of the best-fit line corresponding to the above plot. All experimental data points of the binding isotherm were fitted by linear fit analysis using Microsoft EXCEL and Origin 8.0.

**Circular Dichroism.** Circular Dichroism spectroscopy was carried out using a MOS-450 spectrometer, Bio-Kine. The instrument was operated in the near-UV range to study the ferri-enterobactin binding to VcFhuD. The instrument was operated with the following parameters: wavelength range, 250–500 nm; scanning mode, continuous; scanning speed, 100 nm/min; response, 0.5 s; accumulation, 3; pathlength, 1 mm. The experiment was carried out at 298 K. The reactions were performed in a buffer containing 50 mM Tris-HCl (pH 7.0) and 300 mM NaCl. The concentration of VcFhuD was 33 \( \mu \)M and ligand concentration varied from 20 to 300 \( \mu \)M. The blank spectrum was subtracted from all spectra.

**Spectroscopic titration assays.** For hemin titrations, 10 \( \mu \)M purified 6 \times His-tagged VcHutB in 300 mM NaCl and 50 mM Tris–HCl (pH 7.0 and 8.0) and its mutants VcHutB-Y65A and Y65F in pH 8.0 were subjected...
to UV-Visible spectroscopy. A 3 mM hemin stock solution was prepared in 100% DMSO. Stock concentration was determined spectrophotometrically (ε 385 = 58,400 M⁻¹ cm⁻¹). Hemin concentration was successively added from 2 μM to 40 μM, into 10 μM of protein. Samples were equilibrated for 10 min after addition of each hemin aliquot and UV/Vis spectra between 250 nm and 700 nm were recorded with a dual-beam spectrophotometer Hitachi U2900 at 293 K. The resulting difference spectra were generated by subtracting the free hemec spectra from the heme-titrated VcHutB spectra. Reproducibility of the results was confirmed by performing the experiments in triplicate. The mean values are reported here with errors representing the standard error of the mean at less than 0.05%.

**Native PAGE heme-binding gel shift assay.** A 3 mM hemin stock solution was prepared as described above. Purified 6 × His-tagged VcHutB (0.2 mM) in buffer containing 300 mM NaCl, 50 mM Tris (pH 8.0) was incubated with hemin of gradually increasing concentration from 0.025 mM to 1.2 mM or with buffer alone for 45 min at 293 K and subjected to native PAGE. The VcHutB mutants Y65A and Y65F (0.2 mM) in the same buffer were individually incubated with 0 mM, 0.1 mM, 0.2 mM and 0.4 mM hemin for 45 min at 293 K and subjected to native PAGE. Similar incubation experiment was carried out with 6 × His-tagged VcHutB at pH 7.0. Continuous native PAGE was performed as reported previously and all the experiments were done at least in triplicate. The hemin-complexed protein species migrated faster compared to the apo-forms as was already described for other hemin binding proteins. Therefore, the disappearance of apo-VcHutB as a function of hemin concentration was quantified densitometrically, after Coomassie staining, using Quantity One Software. The fold-densitometric decrease of apo-HutB-band intensity was plotted against hemin concentration and K₅ was calculated using GraphPad Prism software (www.graphpad.com).

**Molecular dynamics simulation.** We carried out MD simulations for 1000 ns on VcHutB and VcFluD in explicit solvent using Gromacs-5.0.4 program with CHARMM27 all atom force field. The simulation box was a dodecahedron with minimum protein-edge distance of 12 Å and periodic boundary condition was applied on all the three (xyz) directions. The protein was solvated with TIP3P waters and the system was neutralized by adding suitable numbers of ions (5 Na⁺ for VcHutB and 4 Na⁺ for VcFluD) replacing the solvent molecules at random locations. After the energy minimization of the whole system using the steepest descent algorithm, the system was gradually heated to 300 K using NVT ensemble. The system was then equilibrated using NPT ensemble. During the equilibrations, the protein backbone was restraint with a harmonic potential of force constant 1000 kJ/mol. The leap-frog integrator with a time-step of 2 fs was used. The Parrinello-Rahman algorithm was employed to control the pressure at 1 bar with a coupling constant of 2 ps and the modified Berendsen (V-rescale) thermostat was used to control the temperature of the system at 300 K with a time constant of 0.1 ps. The Particle Mesh Ewald (PME) method was used to compute the electrostatic interactions with a real space cut-off distance of 12 Å. The same cut-off value was used for calculations of the van der Waals interactions. After 5 ns of equilibration using position restraints on the protein, the protein MD simulation run was carried out for 1000 ns. The position and the velocity of all the atoms were recorded in the trajectory file at every 20 ps for analysis of the dynamics. To avoid terminal motion excessively dominate the results, the first and last five residues were not considered for the analysis.

**Equipment and settings.** Sequence and structural alignments were carried out using the programs Clustal Omega and PyMOL respectively. Structure figures were prepared using PyMOL. Coulombic potential of surfaces were calculated using the program UCSF-chimera. Figures 3 and 5 were prepared using Microsoft EXCEL and Origin 8.0. Native Gels were scanned using MEGA-CAPT software of Gel Documentation System. All the figures were paneled using Adobe Photoshop CS3.

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Author Contributions
S.A. has carried out most of the experiments with the help of S.D. B.G. has done molecular dynamics simulation and also helped in diffraction data collection. MB was involved in cloning expression and purification. J.D. has conceived the idea of the project, analysed results and has written the manuscript.
Additional Information

Accession Codes: The atomic coordinates and structure factors of apo-VcFhuD, holo-VcFhuD and apo-VcHutB have been deposited in the Protein Data Bank (http://wwpdb.org/) with PDB codes 5GGY, 5GGX, 5KHL respectively.

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