

Concerted Motions Networking Pores and Distant Ferroxidase Centers Enable Bacterioferritin Function and Iron Traffic

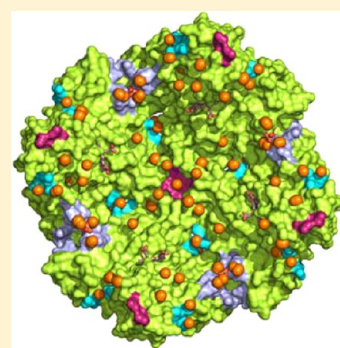
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S Supporting Information

ABSTRACT: X-ray crystallography, molecular dynamics (MD) simulations, and biochemistry were utilized to investigate the effect of introducing hydrophobic interactions in the 4-fold (N148L and Q151L) and B-pores (D34F) of *Pseudomonas aeruginosa* bacterioferritin B (BfrB) on BfrB function. The structures show only local structural perturbations and confirm the anticipated hydrophobic interactions. Surprisingly, structures obtained after soaking crystals in Fe²⁺-containing crystallization solution revealed that although iron loads into the ferroxidase centers of the mutants, the side chains of ferroxidase ligands E51 and H130 do not reorganize to bind the iron ions, as is seen in the wt BfrB structures. Similar experiments with a double mutant (C89S/K96C) prepared to introduce changes outside the pores show competent ferroxidase centers that function akin to those in wt BfrB. MD simulations comparing wt BfrB with the D34F and N148L mutants show that the mutants exhibit significantly reduced flexibility and reveal a network of concerted motions linking ferroxidase centers and 4-fold and B-pores, which are important for imparting ferroxidase centers in BfrB with the required flexibility to function efficiently. In agreement, the efficiency of Fe²⁺ oxidation and uptake of the 4-fold and B-pore mutants in solution is significantly compromised relative to wt or C89S/K96C BfrB. Finally, our structures show a large number of previously unknown iron binding sites in the interior cavity and B-pores of BfrB, which reveal in unprecedented detail conduits followed by iron and phosphate ions across the BfrB shell, as well as paths in the interior cavity that may facilitate nucleation of the iron phosphate mineral.



Iron, an essential nutrient for pathogenic bacteria, can also stimulate the formation of reactive oxygen species via the Haber Weiss cycle, in which free iron catalyzes the conversion of hydrogen peroxide and superoxide to the highly toxic hydroxyl radical.^{1,2} Consequently, free levels of iron in bacteria are tightly regulated to ensure sufficiency for metabolic needs, while preventing iron-induced oxidative toxicity. To maintain iron homeostasis, pathogens must balance the need to obtain iron from their host with careful management of intracellular iron levels, which includes storage of iron reserves for subsequent utilization when the nutrient becomes scarce.^{2,3} Bacteria have evolved two types of proteins for storing iron, ferritin (Ftn) and bacterioferritin (Bfr); the latter is unique to bacteria.^{2,4} The significance of bacterial Ftn and Bfr in the life cycle and virulence of pathogens is just beginning to emerge, as reflected in recent findings with Bfr and Ftn mutants of *Mycobacterium tuberculosis*, which are highly susceptible to antibiotics and unable to persist in mouse and guinea pig models of infection,^{5,6} and in recent observations suggesting that immunization of laboratory animals with a *bfrB* mutant strain of *M. tuberculosis* stimulates protective immunity.⁷ In the plant pathogen *Erwinia chrysanthemi* mutation of the *bfr* gene results in impaired iron utilization and growth defects.⁸

The subunit architecture of eukaryotic Ftms, bacterial Ftms, and Bfrs is highly conserved and consists of five α -helices (A–E) arranged in a four-helix bundle (A–D) and a short helix (E) that lies nearly perpendicular to the central axis of the bundle; helices B and C are connected by a long loop that traverses the length of the four-helix bundle. Although the structures of Bfrs are similar to those of Ftms in terms of the overall architecture of the subunits and how they assemble together into 24-mers, their amino acid sequences exhibit little homology (<18%).^{9,10} In addition, Bfrs are unique in possessing intrinsic heme groups, which are bound at 2-fold symmetric intersubunit sites by coordinative interactions with methionine residues from adjacent subunits (Figure 1A). Twenty four subunits and 12 hemes assemble into a spherical and hollow structure (Figure 1B) with an outer diameter of ~120 Å and an inner diameter of ~80 Å, where up to ~3500 iron atoms can be stored in the form of a Fe³⁺ mineral.¹¹ Details pertaining to the self-assembly and stability of the 24-mer bacterioferritin shells are beginning to emerge.^{12,13} The formation of an iron core (iron uptake) requires

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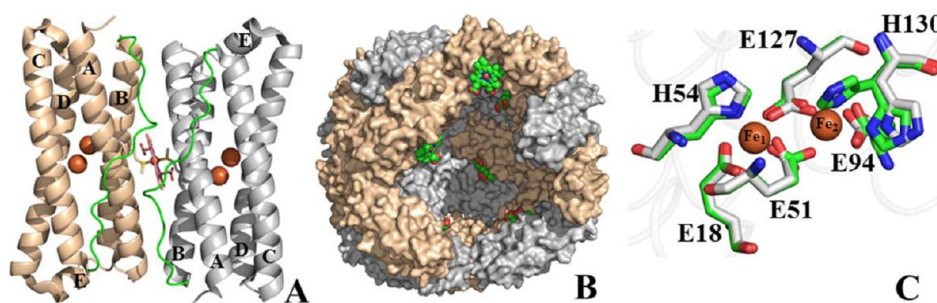


Figure 1. Structural organization of BfrB (PDB: 3IS8). (A) Subunit dimer and associated heme; each subunit is composed of a four-helix bundle (A–D), a perpendicular short helix (E), and a loop connecting the B and C helices (green). The location of the two iron ions in the ferroxidase center of each subunit is shown as orange spheres. (B) Cross sectional view of the 24-mer assembly illustrating the internal cavity where mineral iron is stored; heme molecules are shown in sphere representation (green). (C) Overlay of ferroxidase center residues from the as-isolated (gray) and Fe-soaked (green) wt BfrB structures, illustrating the conformational rearrangements of E51 and H130 upon binding iron.

binding of ferrous iron (Fe^{2+}) to a ferroxidase catalytic site, where it is oxidized to the ferric (Fe^{3+}) state^{11,14,15} and then translocated to the interior cavity. Reincorporation of iron in metabolism requires accepting electrons to reduce Fe^{3+} in the internal cavity and releasing Fe^{2+} to the bacterial cytosol.^{16,17} This dual function (iron uptake and iron release) enables an equilibrium that regulates the range of cytosolic Fe^{2+} concentrations that allow Fur (Fe uptake repressor) to perform a broad range of regulatory functions linking iron homeostasis to broader bacterial metabolism.^{18–20} *Pseudomonas aeruginosa* has two ferritin-like molecules, a bacterial ferritin (FtnA) and a bacterioferritin (BfrB).²¹ The release of iron stored in BfrB involves reduction of the Fe^{3+} mineral, which requires that electrons originating in NADPH are shuttled to BfrB by way of an NADPH ferredoxin reductase (FPR) and a ferredoxin, termed bacterioferritin-associated ferredoxin (Bfd).¹⁷ The X-ray crystal structure of the BfrB–Bfd complex revealed that residues at the complementary BfrB–Bfd interface are highly conserved in Bfr and Bfd sequences from a number of pathogenic bacteria, suggesting that the BfrB/Bfd interaction is important in the regulation of cytosolic iron concentrations in Gram-negative pathogens.¹⁶

In Bfrs the ferroxidase catalytic centers are located in the middle of each subunit (Figure 1A,C). In the structure of as-isolated recombinant wt BfrB, the ferroxidase centers are devoid of iron, and although most of the ferroxidase residues are poised to bind iron, the side chain of H130 is rotated away; we termed this conformation of H130 “gate open” (gray in Figure 1C).²² Soaking crystals of as-isolated wt BfrB in Fe^{2+} solution causes iron to load onto the ferroxidase center, and the H130 side chain is seen in two conformations, one coordinated to Fe_2 (“gate closed”) and the other in the “gate open” state (Figure 1C).²² These structural observations, together with findings obtained from iron incorporation studies in solution, support the idea of a gating mechanism in BfrB, where a ferroxidase channel allows entry of Fe^{2+} from the protein exterior to the ferroxidase center, where a di- Fe^{2+} moiety is coordinated by the ferroxidase ligands, including H130. Oxidation to di- Fe^{3+} is thought to signal H130 to adopt the gate open conformation and allow Fe^{3+} translocation from the ferroxidase center to the internal cavity.²²

The structures of ferritin-like molecules suggest that in addition to the ferroxidase pores, the protein interior communicates with the exterior via 8 3-fold pores, 6 4-fold pores, and 24 B-pores (Figure 2A,B). The 3-fold pores assemble at the intersection of three subunits and in BfrB are formed by layers of alternating

positively and negatively charged residues (Figure 2C).²² The 4-fold pores, which assemble at the intersection of 4-subunits, are lined by the corresponding E-helices; their narrowest portion is composed of two layers made by the side chains of four N148 and four Q151 (Figure 2D). Bacterial Ftns and Bfrs also have B-pores, which are formed at the intersection of three subunits not aligned with an axis of symmetry. Four B-pores surround each 4-fold pore, such that there are 24 B-pores in the 24-mer. In BfrB the narrowest section of a B-pore is defined by D132 and T136 from one subunit, and D34 from a different subunit (Figure 2E). It has been noted that the size of B-pores may allow the traffic of Fe^{2+} across the Bfr shell.^{23,24}

The low amino acid sequence conservation among Ftns, bacterial Ftns and Bfrs^{9,10} profoundly influences packing of the protein shell, charge distribution inside the pores, and possibly pore function: The 3-fold pores of eukaryotic Ftns are lined with negatively charged residues,²⁵ which are thought to function as conduits for iron across the eukaryotic Ftn shell.^{26,27} A similar function, however, is not obvious for the 3-fold pores in Bfrs, which have a layered structure of alternating positive and negative charge. The 4-fold pores of eukaryotic Ftn are lined with hydrophobic residues, but contain hydrophilic amino acids in BfrB (Figure 2D). In reported BfrB structures, a cation has been observed in the pores, which is best modeled as potassium²² and sodium,¹⁶ whereas in two distinct structures of *Azotobacter vinelandii* Bfr the cation was modeled as a barium²⁸ or iron.²⁴ Eukaryotic Ftns lack B-pores, whereas the 24 B-pores in BfrB are lined with hydrophilic and negatively charged residues (Figure 2E). Moreover, in molecular dynamics (MD) simulations carried out with wt BfrB, K^+ ions were seen to traffic in and out of the BfrB shell via B- and ferroxidase pores, but not through the 3-fold or 4-fold pores.²⁹ The MD simulations also suggest that movement of K^+ through B-pores is enabled by periodic kinking/straightening, as well as folding/unfolding of the C-terminal half of helices D, and that these fluctuations are coupled to lateral oscillations of the short E-helices that constitute the 4-fold pores, such that the aggregate of the motions is a periodic expansion and contraction of the B-pores. More recently, elastic network models were used to predict extensive networks of highly correlated residues connecting ferroxidase centers with B- and 4-fold pores in Bfr and bacterial Ftn, leaving the 3-fold pores disconnected from the grid, while in contrast a network of correlated residues in eukaryotic Ftn connects ferroxidase centers and 3-fold pores.¹⁰ These predictions are not only in agreement with the putative distinct function of the different pores in Bfr and eukaryotic

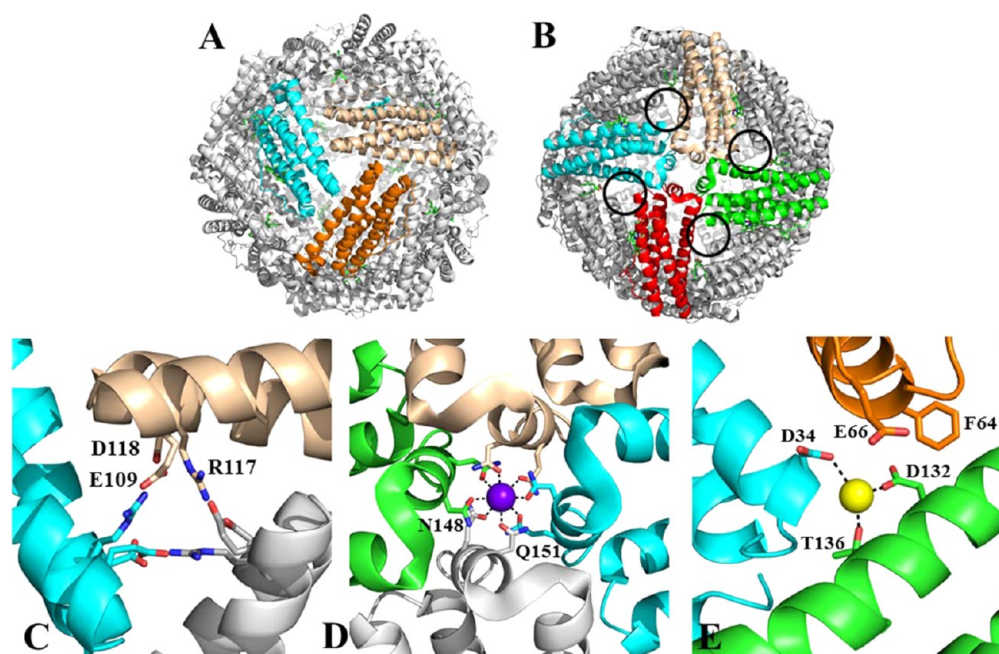


Figure 2. A 24-mer BfrB viewed (A) down one of the eight 3-fold pores, and (B) down one of the six 4-fold pores; four B-pores surrounding a 4-fold pore are highlighted by black circles. (C) Zoomed-in view of a 3-fold pore illustrating the alternating arrangement of positively and negatively charged residues, (D) a 4-fold pore and associated potassium ion (purple sphere) coordinated by the side chains of N148 and Q151, and (E) a B-pore and associated sodium ion (yellow sphere) coordinated by D34, D132, and T136.

Ftn, but also support the idea that cooperative dynamics of the 24-mer architecture are pivotal for iron traffic and iron handling inside the ferritin cavity.

Despite the significant advances made toward understanding Bfr function, many important questions remain unanswered. For example, it is not yet clear how ions (positive and negative) enter or exit the interior cavity through the protein shell, or how these ions are “handled” in the protein interior. It is also not known how the dynamics of individual subunits and 24-mer assemblies contributes to the function of Bfr and the traffic of ions across the protein shell. In an attempt to provide answers to these questions, in this work we investigated the effect that mutations in the 4-fold and B-pores may exert in the structure, iron uptake efficiency and dynamics of BfrB. N148 and Q151, the residues in helices E that line the 4-fold pores and normally coordinate K^+ or Na^+ (Figure 2D), were each replaced by leucine, and D34, one of the negatively charged residues that is part of the narrowest section of the B-pores and coordinates a Na^+ ion within the pore (Figure 2E), was replaced with phenylalanine. Our results show that the site-directed alterations not only exert a local effect on the targeted pores, but also influence the flexibility and reactivity of relatively distant ferroxidase centers. MD simulations carried out with the X-ray crystal structures of select mutants provide a platform for interpretation of the experimental observations in the context of cooperative motions, extending from 4-fold pores, via B-pores, into ferroxidase centers, which are required for efficient oxidation and mineralization of iron by the 24-mer bacterioferritin assembly. Structural studies of the BfrB mutants carried out after exposure of single crystals to Fe^{2+} solution provide experimental support for iron traffic through B-pores, and compilation of iron binding sites in the interior surface of the distinct mutants gives unprecedented insight into the paths followed by iron and phosphate, from entry ports in the Bfr shell toward possible mineralization sites in the interior cavity.

EXPERIMENTAL PROCEDURES

Site-Directed Mutagenesis and Protein Expression.

The pET11a vector containing the *bfrB* gene¹⁷ was mutated to D34F, N148L, Q151L, and C89SK96C using the QuikChange II Site-Directed Mutagenesis kit (Stratagene, La Jolla, CA) using the manufacturer instructions. Primer pair sequences are provided in Supporting Information. PCR products were digested using Dpn I and transformed into XL1-Blue Super-competent cells (Agilent Technologies) for DNA amplification. Plasmid DNA was isolated using the QIAprep Spin Miniprep Kit (QIAGEN), and the sequences verified by ACGT Inc. (Wheeling, IL). Recombinant DNA plasmids with the correct sequence were transformed to *Escherichia coli* ArcticExpress (DE3)RIL competent cells (Agilent Technologies) for subsequent protein expression. The protocols for protein expression, purification, and reconstitution with heme have been described previously.^{17,22}

Crystallization and Data Collection. All crystallization experiments were conducted in Compact 300 or CombiClover 500 (Rigaku Reagents) sitting drop vapor diffusion plates at 20 °C. Equal volumes of BfrB (10 mg/mL in 100 mM potassium phosphate buffer, pH 7.6 and 1 mM TCEP) and crystallization solutions were equilibrated against 75 μ L reservoir volume. Within 1 day, red prismatic crystals were obtained from Wizard 2 (Rigaku Reagents) condition E2 (35% (v/v) 2-methyl-2,4-pentanediol, and 100 mM MES pH 6.5, 200 mM Li_2SO_4). Crystals were transferred to a fresh drop of crystallization solution, which served as the cryoprotectant, and stored in liquid nitrogen for data collection. Fe-soaked crystals were prepared by soaking native crystals for 10 min in 50 mM $FeCl_2$ freshly dissolved in crystallization solution and then frozen in liquid nitrogen. X-ray diffraction data were collected at the Advanced Photon Source beamline 17-ID (IMCA-CAT) using a Dectris Pilatus 6 M pixel array detector.

Structure Solution and Refinement. Intensities were integrated using XDS³⁰ via the XDSAPP³¹ interface or the Autopro³² software package and the Laue class analysis, and data scaling were performed with Aimless.³³ All crystals were isomorphous ($P2_12_12_1$, 24 molecules/asu) with the previously determined BfrB structure (PDB: 3IS7).²² However, BfrB (C89S/K96C) crystals were found to belong to the Laue class 6/*mmm* ($a = 158.57$ Å, $c = 152.38$ Å) with the likely space group $P6_322$ (4 molecules/asu). The Matthews coefficient³⁴ (V_m) suggested that there were likely four ($V_m = 3.2$ Å³/Da, 62% solvent) molecules in the asymmetric unit. This is similar to the Matthew's coefficient observed for the $P2_12_12_1$ crystal form (3.0 Å³/Da, 59% solvent), indicating similar crystal packing in both crystals. Structure solution was conducted by molecular replacement using a single subunit from PDB: 3IS7 as the search model with Phaser.³⁵ All space groups with 622 point symmetry were tested in the molecular replacement searches. The top solution was found in the space group $P6_322$ with four molecules in the asymmetric unit. A second crystal form of as-isolated BfrB (C89S/K96C) was obtained from crystals that had been incubating in the trays for over six months, which resulted in significantly different lattice constants of $a = b = 155.13$ Å, $c = 155.18$ Å. The biological assembly (24-mers) of C89S/K96C BfrB can be generated by application of the space group symmetry operators to the asymmetric units. Structure refinement and manual model building were conducted with Phenix³⁶ and Coot³⁷ respectively. Disordered side chains were truncated to the point where electron density could be observed. Structure validation was conducted with Molprobit.³⁸ Coordinates and structure factors were deposited to the Worldwide Protein Data Bank, and the accession codes are provided in Table 1.

In order to assign the Fe sites for the Fe-soaked BfrB (C89S/K96C) structure, data were collected at a wavelength corresponding to the iron absorption edge (1.7396 Å). This structure was refined using data to 1.8 Å resolution. During subsequent data collection at the synchrotron, diffraction data were collected for another Fe-soaked BfrB (C89S/K96C) crystal, and data were collected at wavelengths of 1.7398 Å and 1.7463 Å, which correspond to the absorption peak and low energy remote respectively. The diffraction data were a bit weaker for this sample and were scaled to 2.1 Å resolution. However, this allowed for the confirmation of the assigned Fe sites, particularly those that displayed weaker electron density and possibly lower occupancy. By computing phased anomalous difference maps the anomalous signal was compared at the peak/remote wavelengths. If Fe is indeed present at a particular site, the anomalous signal should disappear when using the low energy remote wavelength data. Conversely, an atom such as a K⁺ ion, which yields an appreciable anomalous signal at the Fe peak wavelength, would display an increased anomalous signal at the low energy remote wavelength. This second data set (peak and low energy remote) was included with the structure factors deposited to the PDB for the Fe-soaked C89S/K96C structure.

Effect of the Mutations on Iron Oxidation and Uptake in Solution by wt and Mutant BfrB. As-isolated BfrB (wt and mutants) contain only a very small amount of iron in their core.¹⁷ Reconstitution with an iron core was carried out as previously reported.¹⁷ A 10 mM solution of ferrous ammonium sulfate was prepared inside an anaerobic chamber (Coy Laboratories, Grass Lake, MI), placed in a container with a rubber septum, and removed from the chamber. Concentrated

HCl was added to the ferrous ammonium sulfate solution (50 μL/100 mL) through the septum with the aid of a Hamilton microsyringe (final pH ≈ 2.0), and the resultant solution was titrated into a stirred solution of 2 μM BfrB and 1.0 mM TCEP in 100 mM phosphate buffer, pH 7.6, in aliquots delivering 50 Fe²⁺ ions/BfrB. Fifteen minutes were allowed after the addition of each aliquot; upon the addition of the total iron load (500 Fe atoms/BfrB), the solutions were stirred overnight at 4 °C and then passed through a Sephadex G25M size exclusion column (GE Healthcare). The iron content of the samples, prior and after reconstitution with iron, was analyzed using a colorimetric ferrozine-based assay,³⁹ as reported previously.¹⁷ 50 μL concentrated HCl was added to 50 μL of mineralized BfrB and the mixture was incubated for 15 min at room temperature prior to the addition of 50 μL f ascorbic acid (25 mg/mL) and 250 μL of saturated sodium acetate. The concentration of iron was determined using the absorbance at 562 nm ($\epsilon_{562\text{nm}} = 27.9$ mM⁻¹ cm⁻¹) 15 min after the addition of ferrozine (5 mg/mL).

Molecular Dynamics Simulations of BfrB D34F and N148L Mutants. Each mutant BfrB (D34F and N148L) was first immersed in a pre-equilibrated cubic water box of 160 Å in all three dimensions. This box is the same size as that used in the Grand Canonical Monte Carlo/Brownian Dynamics simulations of wt BfrB described previously.^{29,40} Both the box and the BfrB molecule were centered at the origin. The initial ion configurations were the same as those in the E2, E10, and E40 systems described previously;²⁹ the free K⁺ ion numbers are 2840 (E2), 1042 (E10), and 714 (E40) and the numbers of HPO₄²⁻ ions are 1231 (E2), 332 (E10), and 168 (E40). In each of the D34F mutant systems, 24 K⁺ ions were added in the bulk solution to make the total charge of the systems neutral, and they were distributed with 2000 Monte Carlo moves, which were either accepted or rejected based on Metropolis criteria. Water molecules within 2.4 Å from the ions and the BfrB heavy atoms were removed. The systems were then subjected to a 900-ps equilibration cycle with decreasing positional harmonic restraints on heavy atoms not including water oxygen. In the BfrB interior, the removed water molecules that were close to BfrB before the equilibration cycle were quickly replaced by water from the interior cavity of BfrB, creating small vacuum pockets. To overcome this problem, an additional sphere of pre-equilibrated water with 40 Å radius was added to the system on top of the other water molecules (also see ref 29). Any of these newly added water molecules that were within 2.4 Å of either the BfrB heavy atoms or the other water molecules in the system were deleted. Each of the resultant systems was again equilibrated with diminishing positional-harmonic-restraint potentials on the heavy atoms for another 900 ps and then subjected to a 40-ns production.

All the simulations were carried out in NPT (constant particle number, pressure, and temperature) ensembles using the NAMD2.9⁴¹ simulation package. The simulation inputs were obtained from the Quick MD Simulator module in CHARMM-GUI.⁴² The simulations were performed with CHARMM all-atom parameter set PARAM22⁴³ including the dihedral cross-term corrections (CMAP)⁴⁴ and a modified TIP3P water model.⁴⁵ The van der Waals interactions were smoothly switched off at 10–12 Å by a force switching function,⁴⁶ and the electrostatic interactions were calculated using the particle-mesh Ewald method⁴⁷ with a mesh size of ~1 Å for fast Fourier transformation, $\kappa = 0.34$ Å⁻¹, and a sixth-order B-spline interpolation. The temperature (300 K) and pressure

Table 1. Crystallographic Data for BfrB Structures

Data Collection	as-isolated (N148L)	Fe-soaked (N148L)	as-isolated (QJ51L)	Fe-soaked (QJ51L)	as-isolated (D34F)	Fe-soaked (D34F)	as-isolated (C89S/K96C)	as-isolated-2 (C89S/K96C)	Fe-soaked (C89S/K96C)
unit cell parameters (Å, °)	$a = 125.73$ $b = 203.35$ $c = 208.27$ $P_{21}2_12_1$	$a = 125.32$ $b = 203.29$ $c = 207.18$ $P_{21}2_12_1$	$a = 125.67$ $b = 203.15$ $c = 207.79$ $P_{21}2_12_1$	$a = 125.59$ $b = 203.22$ $c = 207.40$ $P_{21}2_12_1$	$a = 125.68$ $b = 203.54$ $c = 206.97$ $P_{21}2_12_1$	$a = 125.49$ $b = 203.40$ $c = 207.25$ $P_{21}2_12_1$	$a = b = 158.57$ $c = 152.38$	$a = b = 155.13$ $c = 155.18$	$a = b = 153.99$ $c = 155.20$
space group	$P_{21}2_12_1$	$P_{21}2_12_1$	$P_{21}2_12_1$	$P_{21}2_12_1$	$P_{21}2_12_1$	$P_{21}2_12_1$	P_{6322}	P_{6322}	P_{6322}
resolution (Å) ^a	49.39–2.00 (2.03–2.00)	49.36–1.95 (1.98–1.95)	49.24–1.95 (1.98–1.95)	49.35–2.25 (2.29–2.25)	49.41–2.05 (2.09–2.05)	49.39–2.20 (2.24–2.20)	49.13–1.65 (1.68–1.65)	48.27–1.80 (1.83–1.80)	48.23–1.80 (1.83–1.80)
wavelength (Å)	1.74013	1.74013	1.00000	1.73964	1.00000	1.73769	1.00000	1.00000	1.73963
temperature (K)	100	100	100	100	100	100	100	100	100
observed reflections	2,270,196	3,760,949	2,639,550	1,749,112	2,240,862	1,490,791	2,688,900	2,015,697	1,825,740
unique reflections	356,578	373,157	384,285	250,626	328,340	257,692	134,788	101,666	99,837
$\langle I/\sigma(I) \rangle^a$	10.8 (1.7)	14.7 (2.2)	10.8 (2.2)	12.2 (2.4)	13.1 (2.2)	18.1 (3.4)	12.3 (2.3)	20.8 (2.3)	22.3 (1.8)
completeness (%) ^a	99.7 (94.1)	97.7 (91.5)	100 (100)	100 (100)	99.5 (98.9)	96.3 (71.8)	100 (99.9)	100 (100)	99.7 (96.5)
multiplicity ^a	6.4 (4.7)	10.1 (8.1)	6.9 (6.4)	7.0 (6.7)	6.8 (6.6)	5.8 (2.4)	19.9 (20.3)	19.8 (19.2)	18.3 (10.2)
$R_{\text{merge}}(\%)^{a,b}$	12.1 (81.5)	14.0 (92.9)	14.4 (95.8)	14.4 (96.7)	11.7 (95.1)	6.2 (25.9)	16.8 (179.1)	10.6 (160.0)	12.1 (131.3)
$R_{\text{meas}}(\%)^{a,d}$	13.2 (91.6)	14.8 (99.1)	15.6 (103.3)	15.2 (104.8)	12.6 (103.1)	7.4 (35.6)	17.7 (188.2)	10.9 (164.3)	12.4 (138.3)
$R_{\text{pim}}(\%)^{a,d}$	5.2 (40.8)	4.6 (33.6)	5.9 (40.9)	5.7 (40.3)	4.8 (39.7)	2.4 (21.4)	3.9 (41.2)	2.4 (37.2)	2.8 (42.7)
$CC_{1/2}^{a,e}$	0.996 (0.609)	0.996 (0.712)	0.996 (0.695)	0.996 (0.680)	0.998 (0.710)	0.998 (0.885)	0.997 (0.783)	1.000 (0.818)	0.999 (0.701)
Refinement									
resolution (Å)	49.39–2.00	49.36–1.95	49.34–1.95	49.35–2.25	41.06–2.05	48.39–2.20	38.10–1.65	37.27–1.80	48.23–1.80
reflections (working/test) ^f	654,121/ 34,746	688,869/ 36,492	364,760/ 19,327	461,135/ 24,449	311,642/ 16,559	485,422/ 24,416	127,956/ 6,770	96,523/ 5,074	180,676/ 9,520
$R_{\text{actof}}/R_{\text{free}}(\%)^c$	16.9/21.1	15.5/19.3	15.6/18.8	15.4/19.2	15.3/19.2	14.9/18.7	14.5/16.2	15.3/17.8	14.7/17.2
no. of atoms (protein/Heme/ K ⁺ /iron/sulfate/water)	30,466/1,032/6/ 0/0/ 2,588	30,665/1,032/6/ 126/40/ 3,005	30,252/5160/ 0/0/ 2,998	30,343/516/0/ 120/40/ 2,110	30,434/516/6/0/ 2,0660/2,066	30,262/516/6/ 144/40/ 2,278	5,151/172/1/0/ 0/762	5,151/172/1/0/ 8/641	5,165/172/1/ 37/ 8/748
Model Quality									
r.m.s deviations									
bond lengths (Å)	0.010	0.956	0.009	0.011	0.016	0.010	0.008	0.010	0.009
bond angles (°)	0.975	0.017	1.008	1.023	1.265	0.985	1.107	0.996	1.013
average B-factor (Å ²)									
all atoms	26.0	22.6	20.9	24.8	28.6	24.8	22.6	27.7	24.8
protein	25.4	19.0	19.9	31.7	28.2	24.1	20.6	26.2	23.1
heme	27.9	21.8	18.0	30.4	25.6	24.4	20.2	24.3	22.9
K ⁺	38.6	28.2			21.5	39.9	15.9	20.6	19.5
iron		45.1		60.0		45.1			46.8
sulfate		48.0		59.8		53.4		52.2	39.3
water	32.9	20.0	29.5	39.3	35.7	31.2	34.0	38.8	35.4
coordinate error (maximum likelihood) (Å)	0.24	0.19	0.17	0.22	0.20	0.20	0.15	0.17	0.18

Table 1. continued

Ramachandran Plot	as-isolated (N148L)	Fe-soaked (N148L)	as-isolated (Q151L)	Fe-soaked (Q151L)	as-isolated (D34F)	Fe-soaked (D34F)	as-isolated (C89S/K96C)	as-isolated-2 (C89S/K96C)	Fe-soaked (C89S/K96C)
	most favored (%)	most favored (%)	most favored (%)	most favored (%)	most favored (%)	most favored (%)	most favored (%)	most favored (%)	most favored (%)
4TO9	99.7	99.6	100	100	99.5	99.8	99.7	99.7	99.5
4TOA	0.3	0.4	0.3	0.3	0.5	0.2	0.3	0.3	0.5
4TOB									
4TOC									
4TOE									
4TOF									
4TOG									
4TOH									

^aValues in parenthesis are for the highest resolution shell. $R_{\text{merge}} = \sum_i |I_i(hkl) - \langle I_i(hkl) \rangle| / \sum_i I_i(hkl)$, where $I_i(hkl)$ is the intensity measured for the i th reflection and $\langle I_i(hkl) \rangle$ is the average intensity of all reflections with indices hkl . $R_{\text{factor}} = \sum_i |F_o(hkl) - F_c(hkl)| / \sum_i F_o(hkl)$; $R_{\text{free}} = \sum_i |F_o(hkl) - F_c(hkl)| / \sum_i F_o(hkl)$; $R_{\text{pim}} = \sum_i |F_o(hkl) - F_c(hkl)| / \sum_i F_o(hkl)$; $R_{\text{merge}}^{60,61}$ is the correlation coefficient of the mean intensities between two random half-sets of data. ^{62,63} For the data sets obtained from iron soaked crystals or native data sets collected at the iron peak wavelength, the number of reflections used during refinement is greater than the number unique reflections reported for data scaling. This is because Friedel pairs were kept separate during refinement and the anomalous scattering factors for the Fe²⁺ atoms were refined.

(1 atm) were kept constant during all the simulations by Langevin dynamics and the hybrid Nosé-Hoover Langevin piston method, respectively. The Langevin damping coefficient was set to 1 ps⁻¹; the decay period and damping time scale were 50 fs, respectively.

RESULTS

X-ray Crystal Structures. The structures of four BfrB mutants (N148L, Q151L, D34F, and C89S/K96C) were determined using crystals grown from purified protein (as-isolated) and using crystals of pure protein after soaking in Fe²⁺ solution (Fe-soaked). Two mutants (N148L and Q151L) were prepared to investigate the effect of replacing the hydrophilic side chains that coordinate K⁺ in the 4-fold pores of BfrB (Figure 2D) with hydrophobic noncoordinative side chains. A third mutant (D34F) was prepared to study the effect of replacing a negatively charged residue, which was shown to bind a Na⁺ ion in the B-pores of BfrB (Figure 2E), with a hydrophobic side chain. As will be shown below, these mutations affect the flexibility of ferroxidase center ligands and the catalytic efficiency of Fe²⁺ oxidation by their respective ferroxidase centers. To show that mutation of residues outside of the pores has no effect on ferroxidase center function, we studied the surface mutant C89S/K96C, which had been prepared for a different investigation. Interestingly, the C89S/K96C mutant not only showed wt-like behavior of the ferroxidase center *in crystallo* and in solution, but also displayed all the iron binding sites observed in the structures of the different mutants reported here and also revealed new ones. The unprecedented observation of six different types of iron binding sites (>200 iron ions not counting heme-iron) in one Bfr structure allows comparison of thermal factors for each type of iron binding site and therefore provides insights into their relative occupancies.

The structures of all the mutants show strong conservation of the subunit architecture and 24-mer assemblies. Superposition⁴⁸ of subunit A from the as-isolated structure of each mutant reported here with subunit A of the previously determined wt BfrB structure (PDB: 3IS7) resulted in the following RMSD between C_α atoms for 153 residues aligned: N148L (0.15 Å), Q151L (0.15 Å), D34F (0.10 Å), and C89S/K96C (0.15 Å). Closer inspection shows that the mutations cause local structural perturbations in the corresponding 4-fold or B-pores relative to the wt structure, as expected. However, comparison of as-isolated and Fe-soaked structures for each of the mutants uncovers important differences: (i) With the exception of the C89S/K96C mutant, iron ions in the ferroxidase centers are not coordinated as in the wt protein, (ii) previously unobserved iron binding sites are present in the interior cavity, near the entrance to 3-fold, 4-fold, and B-pores in all mutants, (iii) sulfate ions are found in the interior cavity at the entrance to 3-fold pores, where they are coordinated by iron ions, (iv) iron ions are observed in B-pores, and (v) the ubiquitous K⁺ ions observed in the 4-fold pores of wt and other mutant BfrB proteins are absent in the Q151L mutant. Pertinent details are presented below.

Four-Fold Pores. In wt BfrB the side chains of N148 and Q151, located in each of the two turns of helices E, point toward the pore interior, where they coordinate a K⁺ ion (Figures 2D and 3A). The structure of 4-fold pores in the as-isolated and Fe-soaked C89S/K96C mutant is nearly identical to that in wt BfrB, including the placement of the K⁺ ion (Figure 3B). As seen previously in the Fe-soaked structure of wt BfrB,²² in the interior cavity of Fe-soaked C89S/K96C four

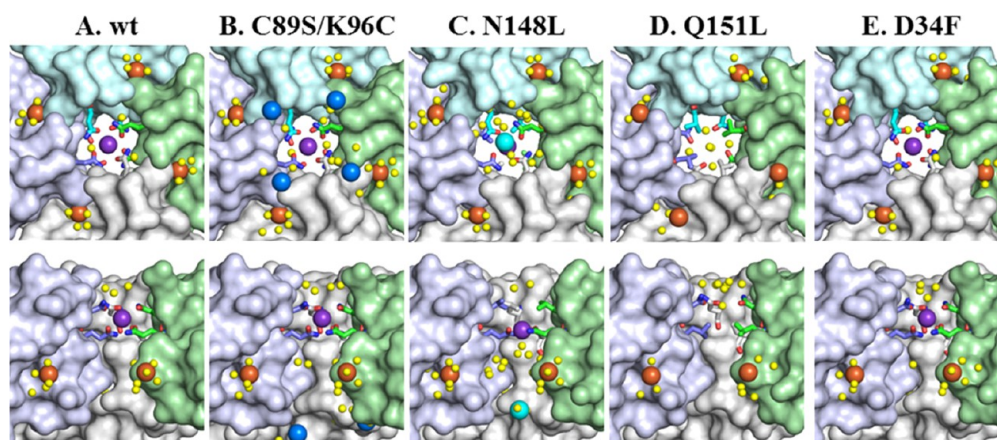


Figure 3. (A–E) Zoomed in view of a 4-fold pore in wt and mutant BfrB molecules. The top row depicts views from the interior cavities, while the bottom row shows cross-sectional views with a subunit removed. Fe_{4f1} iron ions are shown as orange spheres, Fe_{4f2} iron ions as blue spheres, Fe_{4f3} iron ions as cyan spheres, water molecules as yellow spheres, and potassium ions as purple spheres, nitrogen in blue and oxygen in red.

Fe ions (orange spheres) nest in the exterior wall of the 4-fold pore, where they are coordinated by H155 from one subunit (A) and H153 from a different subunit (B) (Figures 3B and S1A of the Supporting Information). These sites were previously termed Fe_{in} to denote their location in the interior cavity. Herein, we will term these sites Fe_{4f1} to indicate that they are the first Fe-sites observed near 4-fold pores. The C89S/K96C structure also revealed previously unobserved Fe sites located deeper in the interior cavity, along the perimeter of the entrance to 4-fold pores from the interior cavity, which we term Fe_{4f2} (blue spheres). Although these Fe ions do not appear to be coordinated by a protein-provided ligand (Figure S1A), interactions with the side chains of one of the terminal residues (E157 or E158), for which there is no observable electron density, cannot be ruled out.

Diffraction data collected at a wavelength of 1.0000 Å did not yield anomalous signal for the K^+ ions in the 4-fold pores. In contrast, diffraction data collected at low energy wavelength (~ 1.74 Å, Fe-edge) resulted in an appreciable anomalous difference density at the K^+ sites, and naturally, at the iron and heme-iron sites. Notably, the anomalous-difference-map peak height increases at the K^+ sites when computed using data collect at a low energy remote wavelength relative to the Fe-edge (Figure S2). This suggests that the assignment of K^+ in the pore interior and Fe ions in the periphery of 4-fold pores in the interior cavity is the correct model.

In the as-isolated N148L structure the conformation of L148 side chains is nearly identical to that of N148 in the wt protein, which results in a tight network of hydrophobic interactions in the pore interior of the mutant. Closer to the interior cavity, the Q151 side chains assume conformations identical to those observed in wt BfrB, and a K^+ is present in the pore interior, but is coordinated only by the side chains of Q151 (Figure 3C). Consequently, K^+ in the 4-fold pores of N148L BfrB is 1.7 Å closer to the interior cavity than equivalent K^+ in the wt protein. As seen previously in the Fe-soaked structure of wt,²² and C89S/K96C BfrB, four Fe_{4f1} ions (orange) nest in the exterior wall of the 4-fold pore inside the BfrB cavity (Figure 3C). The N148L structure also reveals a previously unobserved Fe ion at the entrance to 4-fold pores in the interior cavity, which is termed Fe_{4f3} (cyan spheres). These Fe ions also do not appear to be coordinated by a protein-provided ligand (Figure S1B), although it is likely that the side chains of one of the terminal

residues, for which there is no observable electron density (E157 or E158), provide the necessary stabilizing interactions.

In the structure of as-isolated and Fe-soaked Q151L BfrB the L151 side chains adopt the same conformation as the Q151 side chains in the wt protein, which creates a network of hydrophobic interactions that constrict the interior of 4-fold pores. Interestingly, the K^+ ubiquitously found in *Pa* BfrB structures are absent in the 4-fold pores of the Q151L mutant, in both the as-isolated and Fe-soaked structures (Figure 3D). In the Fe-soaked structure, four Fe_{4f1} ions are present in the exterior perimeter of the 4-fold pore, but unlike N148L, the Fe_{4f3} ion is not observed. The presence of iron is confirmed by strong anomalous difference density observed in diffraction data sets collected at ~ 1.74 Å (Figure S1C). It is also interesting that structured water is absent in the pores below the hydrophobic constriction formed by the L151 side chains in the as-isolated and Fe-soaked structures.

The D34F mutation does not have a structural effect on the 4-fold pores. The side chains of N148 and Q151 coordinate K^+ (Figure 3E) in the as-isolated and Fe-soaked structures. Four Fe_{4f1} ions are observed along the periphery of 4-fold pores in the interior cavity, in positions identical to those observed in wt BfrB. As with the previously discussed structures, anomalous difference density in data sets collected at ~ 1.74 Å corroborate Fe ions at these sites (Figure S1D).

Three-Fold Pores. The 3-fold pores in the structures of the as-isolated N148L, Q151L, D34F, and C89S/K96C mutants are almost identical to their counterparts in wt Bfr (Figure 4 top). The narrowest pore section is lined by side chains from three R117 and three K121, with E109 immediately above (closer to the cytosol) R117 and D118 immediately below (closer to the interior cavity) K121; the K121 side chains cannot be modeled entirely due to disorder in as-isolated BfrB. The innermost layer is wider and demarcated by the side chains of D122 and E125. Soaking crystals of wt BfrB in crystallization solution containing Fe^{2+} led to the observation of a sulfate ion in the 3-fold pores, located between the side chains of R117 and K121 (Figure 4A center).²² Similar iron-soaking experiments carried out with crystals of all mutant proteins allowed observation of a sulfate ion in the innermost layer of each pore, where it is coordinated by three K121 side-chains (Figure 4B–E). Interestingly, in the mutant structures each sulfate ion also interacts with three Fe ions, termed Fe_{3f} (average Fe–O

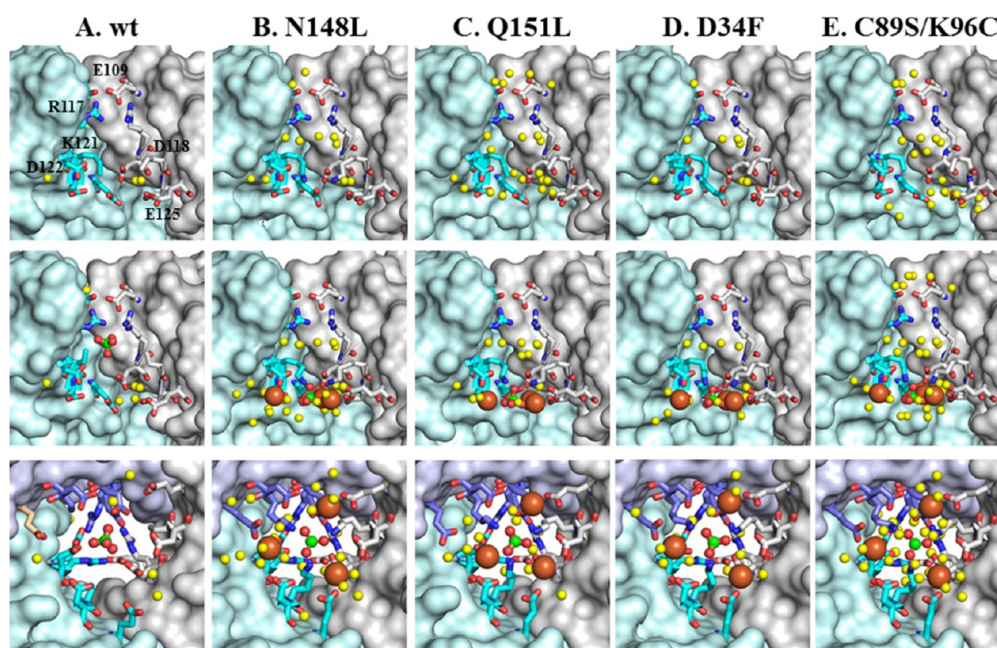


Figure 4. (A–E) View of a 3-fold pore in wt BfrB and mutants. The top row shows cross sectional views of the 3-fold pores in the as-isolated structures. The middle row depicts an identical cross-section view of the 3-fold pores in the Fe-soaked structures, with Fe_{3+} iron ions rendered as orange spheres, sulfate ions as green (S) and red (O) spheres, and structured water as yellow spheres. The bottom row depicts the 3-fold pores in the Fe-soaked structures viewed from the interior cavity, to illustrate the network of interactions involving iron, sulfate, and protein side chains.

distance 4.2 Å), which in turn are coordinated by the side chains of D118, D122, and E125. The side chains of D118 and E125 move from their conformation in the as-isolated structures to coordinate iron in the Fe-soaked structure. The presence of Fe at these sites is supported by anomalous difference density in data sets collected at ~ 1.74 Å (Figure S3).

It is interesting to note that sulfate is absent from the 3-fold pores in all the structures of as-isolated proteins, but is present in the corresponding Fe-soaked structures. An exception is the structure of as-isolated C89S/K96C BfrB obtained from a crystal that had been in the crystallization trays for over 6 months (hereafter as-isolated-2), which showed clear electron density consistent with sulfate ions in the 3-fold pores (Figure S4). In comparison, diffraction data from other as-isolated BfrB crystals was obtained within 20 days of crystal growth. Therefore, it may be reasonable to consider the as-isolated-2 crystal as being soaked for a prolonged period in high sulfate concentration (200 mM Li_2SO_4 from the crystallant), which causes sulfate to slowly occupy the 3-fold pores. As noted in the experimental section, the unit cell parameters differ significantly for the as-isolated and as-isolated-2 C89S/K96C structures. This appears to result in a “tightening up” of the lattice and formation of an intermolecular disulfide bond between C96 of subunit A and its counterpart related by a crystallographic 2-fold axis (Figure S8). It should be noted that soaking of as-isolated C89S/K96C crystals with iron also results in this lattice transformation (see Table 1). Comparison of the 3-fold pores before and after Fe-soak is illustrated in Figure 5: While in the as-isolated-2 structure sulfate nests between R117 and K121 in the middle of the pore, after Fe-soak sulfate is in the most interior pore layer, still coordinated by the side chains of K121, which likely “escort” the anion to its position at the exit of 3-fold pores, where it is seen interacting with iron ions.

A molar ratio of approximately 1.0 phosphate per Fe^{3+} has been observed in the native core of bacterioferritins and bacterial

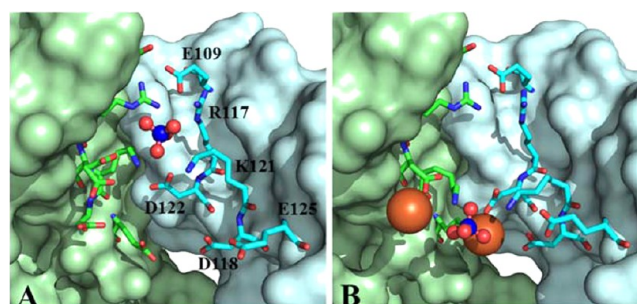


Figure 5. Comparison of 3-fold pores in (A) as-isolated-2 and (B) Fe-soaked C89S/K96C BfrB suggests a possible path for the ingress of sulfate to the interior cavity. Sulfate is shown as blue (sulfur) and red (oxygen) spheres, and Fe_{3+} ions as orange spheres. Key side chains are shown in sticks with nitrogen in blue and oxygen in red.

ferritins, which is distinct from the ratio of approximately 0.1 phosphate per Fe^{3+} observed in the native core of vertebrate ferritins.^{49–51} In this context, comparing the low tendency of sulfate to populate 3-fold pores in as-isolated structures of BfrB (Figure 4, top row) with its ubiquitous presence within the 3-fold pores, or at the exit of these pores into the interior cavity in Fe-soaked BfrB structures (Figure 4, middle row), suggests that phosphate ions utilize 3-fold pores to access the BfrB interior, where they combine with iron ions entering the cavity via other conduits. This idea finds additional support in the highly organized network of iron, sulfate, and acidic side chains observed at the exit of the 3-fold pores into the interior cavity (Figure 4, bottom row) (note: the molecules in 3-fold pores were modeled as sulfate because the crystallization conditions contain 200 mM Li_2SO_4).

B-Pores. The B-pore channels in wt BfrB are formed at the intersection of three subunits not aligned with an axis of symmetry and are demarcated by the side chains of E66, D34,

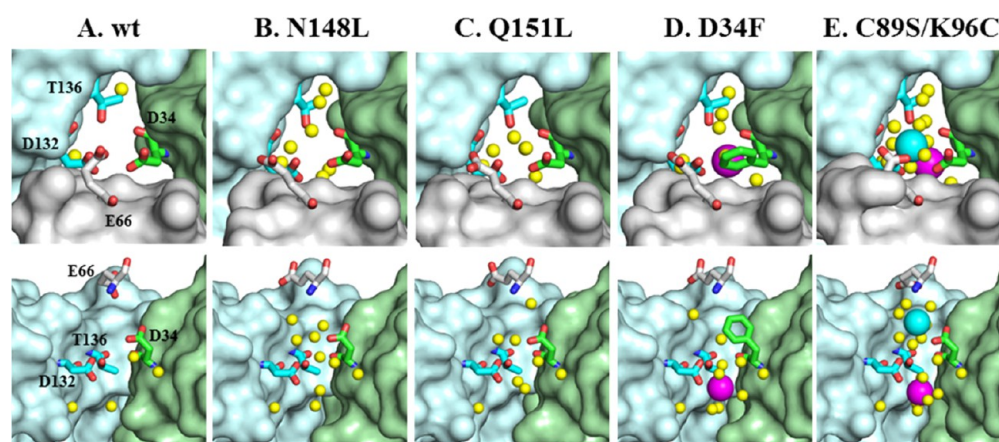


Figure 6. (A–E) Zoomed-in view of B-fold pores in wt BfrB and mutants. The top row depicts the pores viewed from the protein exterior, and the bottom row shows cross sectional views with the gray subunit omitted. $\text{Fe}_{\text{B-1}}$ iron ions are shown as magenta spheres, the $\text{Fe}_{\text{B-2}}$ iron ion is shown as a cyan sphere, water molecules as yellow spheres, nitrogen in blue, and oxygen in red.

T136, and D132, arranged in a corkscrew leading from the outermost (E66) to the innermost (D132) layer (Figure 6A). The structure of B-pores in as-isolated and Fe-soaked N148L and Q151L BfrB are very similar to wt (Figure 6B,C), with the exception of the E66 side chain, which is rotated away and does not participate in the “corkscrew” of hydroxyl and carboxylate groups traversing the channel. In the B-pores of as-isolated D34F BfrB the E66 side chain is also rotated away from the corkscrew, and the side chain of F34 acts as a hydrophobic barrier that occludes the pores and disrupts the hydrophilicity of the corkscrew. Interestingly, the corresponding Fe-soaked structure shows an Fe ion ($\text{Fe}_{\text{B-1}}$) in the innermost layer of the pores, where it is coordinated by D132 (Figure 6D). Anomalous difference density in data sets collected at 1.74 Å support this conclusion (Figure S5A).

The corkscrew of hydroxyl and carboxylate residues seen in wt BfrB is preserved in the B-pores of as-isolated and Fe-soaked C89S/K96C. Remarkably, the Fe-soaked structure of the double mutant reveals two Fe ions aligned along the length of the B-pores (Figure 6E and S5B). One is coordinated by D132 and is located near the interior cavity in a position nearly equivalent to that of the $\text{Fe}_{\text{B-1}}$ ion in the B-pores of Fe-soaked D34F BfrB. The second ($\text{Fe}_{\text{B-2}}$) is near the cytosolic surface, where it is coordinated by D34 and E66. These observations, which provide the first experimental evidence of Fe ions in B-pores, strongly support the idea derived from our MD simulations²⁹ that iron traffics across the Bfr shell using B-pores as conduits.

Ferroxidase Sites. In all the BfrB structures obtained from as-isolated protein, the ferroxidase centers are devoid of iron. Soaking crystals of as-isolated protein in crystallization solution containing iron invariably leads to iron incorporation in the ferroxidase center. In the structures of wt and C89S/K96C BfrB, iron binding at the ferroxidase center is accompanied by a conformation change of the H130 side chain from gate open to gate closed state (Figures 1C and 7A). In contrast, although iron is observed in the ferroxidase centers of D34F, N148L and Q151L BfrB, H130 remains in its gate open conformation, and E51 does not rotate to coordinate and bridge Fe_1 and Fe_2 (Figure 7B–D). The presence of Fe ions in the ferroxidase centers of Fe-soaked structures is supported by anomalous difference density peaks using diffraction data sets collected at ~ 1.74 Å (Figures 7 and S6). The average distance between

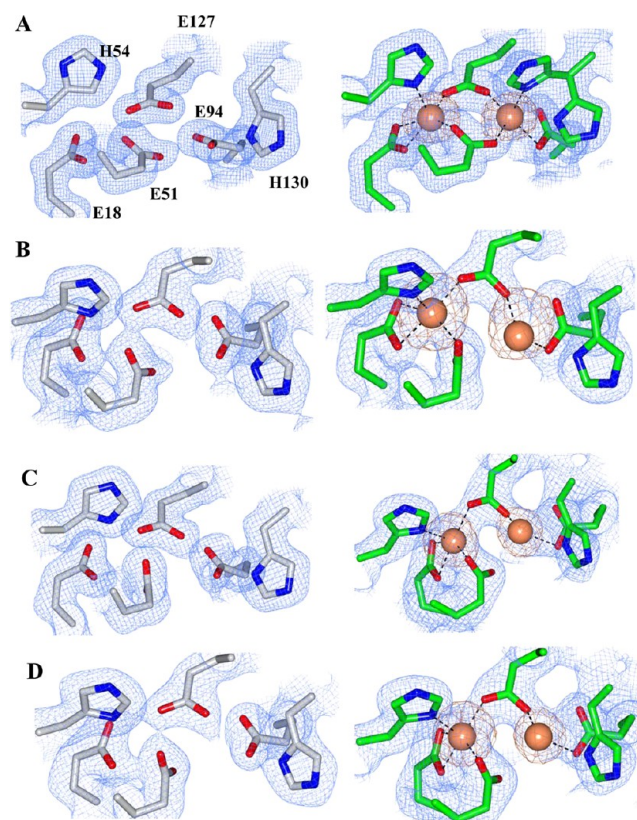


Figure 7. $2F_o - F_c$ electron density map of the ferroxidase ligands (blue mesh) contoured at 1σ and phased anomalous difference map of the ferroxidase center iron atoms (orange mesh) contoured at 3σ for the following BfrB mutant structures: (A) C89S/K96C, (B) N148L, (C) Q151L, and (D) D34F. Ferroxidase centers from as isolated proteins (devoid of iron) are on the left (gray sticks), and ferroxidase centers from iron-soaked structures are on the right (green sticks). Iron atoms are drawn as orange spheres and coordination bonds are represented by dashed lines.

ferroxidase iron ions in all structures is ~ 4.1 Å (Table S3), which suggests di- Fe^{2+} sites.⁵³

The Fe-soaked structure of C89S/K96C BfrB also shows an iron site on the inner surface of the protein, directly below each of the ferroxidase centers ($\text{Fe}_{\text{B-fc}}$), where it is coordinated by H46 (Figures 8 and S7), and close to D50 (~ 4.5 Å). The $\text{Fe}_{\text{B-fc}}$

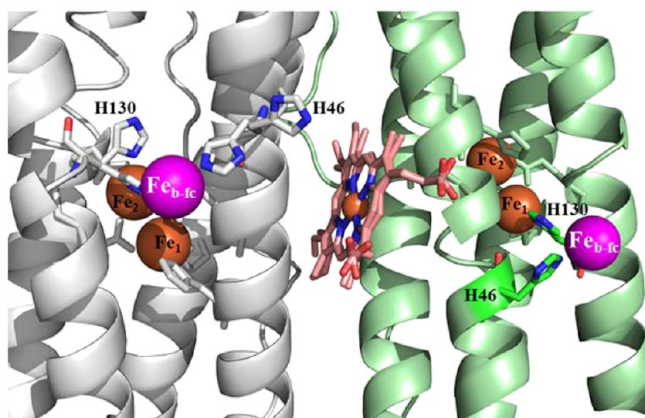


Figure 8. Iron ions below the ferroxidase centers, Fe_{b-fc} (magenta spheres) in the Fe-soaked structure of C89S/K96C BfrB are coordinated by H46. In subunit A (gray) H46 adopts two conformations, one of which coordinates Fe_{b-fc} ; H130 also adopts two conformations, one (gate closed) coordinates Fe_2 (orange), while the second (gate open) is thought to enable its translocation into the interior cavity, aided by H46. In the remainder subunits of the asymmetric unit (B, C, and D), H46 and H130 are observed only coordinated to Fe_{b-fc} and Fe_2 , respectively. The heme is shown in two orientations, which are made evident by the relatively high resolution (1.80 Å) of the structure.

site is equivalent to the iron site observed upon soaking crystals of *E. coli* Bfr in iron-containing solution of crystallant, which was shown to be important in the process of iron mineral growth.⁵⁴ In the asymmetric unit cell of Fe-soaked C89S/K96C BfrB (subunits A, B, C, and D), the side chains of H46 and H130 in subunit A (gray in Figure 8) are observed in two conformations: In one conformation ferroxidase ligand H130 is coordinated to Fe_2 (gate closed) and H46 is coordinated to Fe_{b-fc} , whereas in their alternate conformations both side chains are rotated away from their respective iron ions. In subunits B, C, and D, the H46 and H130 side chains are seen only in their iron bound conformations. The observations in subunit A permit a glimpse into the possible cooperative process of ferroxidase iron translocation, in which the transition of H130 from gate closed to gate open is accompanied by a change in conformation of H46, which serves to coordinate incoming Fe_2 and usher it to the Fe_{b-fc} site. The model is also consistent with the Fe_{b-fc} site serving as an iron depot for the distribution of iron entering via the ferroxidase centers to nucleation and mineralization sites.

Effect of the Mutations on Iron Oxidation and Uptake Function. The X-ray crystal structures show that H130 in the ferroxidase centers of D34F, N148L, and Q151L BfrB does not rotate to coordinate Fe_2 , and that of E51 does not bridge Fe_1 and Fe_2 (see Figures 7 and S6). These observations suggest that mutations in the 4-fold and B-pores adversely affect the function of relatively distant ferroxidase centers. To investigate this idea, we carried out experiments in solution to compare the efficiency of Fe^{2+} oxidation and Fe^{3+} storage exhibited by each mutant relative to wt BfrB. This was carried out by challenging each of the bacterioferritins with 500 Fe^{2+} /BfrB, added in aliquots delivering 50 Fe^{2+} /BfrB, as described in Experimental Methods. Because Fe^{2+} autoxidizes rapidly in aqueous solution at pH above 6.5, and even faster in the presence of phosphate,⁵⁵ the addition of Fe^{2+} to aerobic solutions of BfrB in phosphate buffer, pH 7.6, results in two competing reactions: (i) oxidation of Fe^{2+} at the ferroxidase centers of BfrB and storage of the

resultant Fe^{3+} in the interior cavity (Fe^{3+} is not captured by Bfr or Ftn), and (ii) autoxidation of Fe^{2+} in solution, followed by hydrolysis of Fe^{3+} into an insoluble ferric colloid. The results show that wt and C89S/K96C BfrB oxidize and capture iron with >85% efficiency ($420 \pm 20 \text{ Fe}^{2+}$ /BfrB, and $430 \pm 22 \text{ Fe}^{2+}$ /BfrB, respectively), indicating that the dominant reaction is the oxidation of Fe^{2+} by BfrB. In contrast, the amounts of Fe^{2+} oxidized and captured by the mutants are significantly lower: D34F BfrB oxidized and captured only $110 \pm 30 \text{ Fe}^{2+}$ /BfrB (24%), Q151L captured 320 ± 30 (64%) and the N148L mutant $300 \pm 20 \text{ Fe}^{2+}$ /BfrB (60%). These observations, which show that oxidation of Fe^{2+} by the BfrB mutants does not compete effectively with Fe^{2+} autoxidation, indicate that mutation of important residues at 4-fold pores (N148L, Q151L) and B-pores (D34F) compromises the function of the relatively distant ferroxidase centers. Consequently, the inability of H130 and E51 side chains to rotate and bind iron at their corresponding ferroxidase centers *in crystallo* manifests in solution in the decreased ferroxidase activity exhibited by the 4-fold and B-pore mutants. No attempts were made to determine rates of Fe^{2+} oxidation by BfrB because the only spectroscopic handle to follow the oxidation of Fe^{2+} by BfrB is a broad absorption near 300 nm, which is caused by the formation of Fe^{3+} -O-moieties. Since the ferroxidase activity of the BfrB mutants is compromised, the competing Fe^{2+} autoxidation contributes significantly to the changes in absorbance at 300 nm and therefore precludes meaningful analysis of the progress curves in the context of Fe^{2+} oxidation by BfrB.

Molecular Dynamics Simulations. In an attempt to explain the nature of the disorganization of the ferroxidase centers in the Fe-soaked structures of the D34F and N148L mutants, we investigated their dynamic properties relative to wt BfrB. The per-residue backbone root-mean-square fluctuation (RMSF) plots in the top row of Figure 9 illustrate the relative fluctuations in a BfrB subunit. In wt BfrB (Figure 9A) the flexibility increases with increasing K_2HPO_4 concentrations (from system E40 to E2). In contrast, the flexibility of the mutants is significantly dampened and minimally affected by the same K_2HPO_4 concentrations (Figure 9B,C); the only regions exhibiting a strong tendency to fluctuate are the long loops connecting helices B and C and the N- and C-termini. This is in stark contrast with wt BfrB, where in addition to the BC loops and the N- and C-termini, the C-terminal half of helices D exhibit a strong tendency to fluctuate, followed by the N-terminal half of helix C (Figure 9 middle). A minimal assembly of six subunits containing a 3-fold pore (green star), a 4-fold pore (blue star), and a B-pore (red star), shows that the regions with the highest fluctuations in a subunit of wt BfrB pack to form B- and 4-fold pores (Figure 9 bottom). In contrast, these regions are far less flexible in D34F and N148L BfrB (Figure 9B,C and Supporting Information movies S1-S3).

The structure of D34F BfrB shows that the F34 side chain partially occludes the B-pores and also packs against the L63 side chain. The MD simulations add insight by providing evidence for dampened flexibility in the B pores of the mutant relative to wt BfrB: Figures 10A and 11A depict regions surrounding the B-pores during the MD simulations of wt BfrB, illustrating an unfolded state of the C-terminal section of helix D (cyan), which undergoes frequent folding/unfolding transitions. Such transitions are likely linked to the conformational exchange of D34 and F64 side chains observed in wt BfrB. Figures 10C and 11C, respectively, show the distributions of rotameric states (χ_1, χ_2) for the D34 and F64 side chains in wt BfrB.

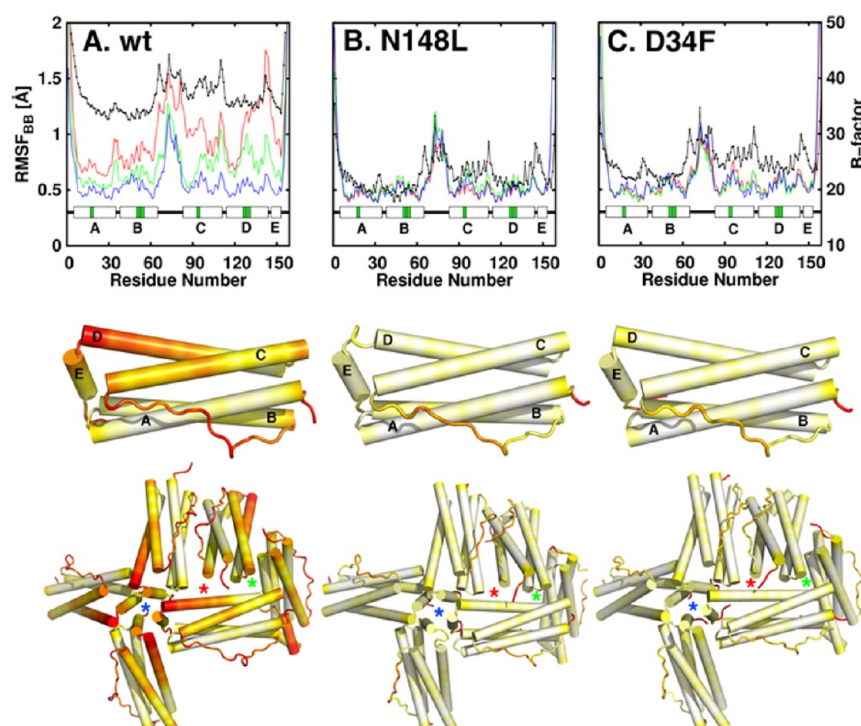


Figure 9. Flexibility in the wt and mutant BfrB molecules. The columns correspond to wt (A), N148L (B), and D34F (C) BfrB. The top row shows plots of per-residue backbone RMSF in system E2 (red), E10 (green), E40 (blue), and per-residue crystallographic B-factors (black); helices A–E are indicated as boxes, and ferroxidase center residues are highlighted in green. The middle row depicts the per-residue backbone RMSF (systems E2) mapped onto a BfrB subunit, and the bottom row shows per-residue backbone RMSF mapped on six subunits of the 24-mer assembly to illustrate relative flexibility at the 4-fold (blue stars), 3-fold (green stars), and B-pores (red stars). Flexibility increases in the color scale from white to red.

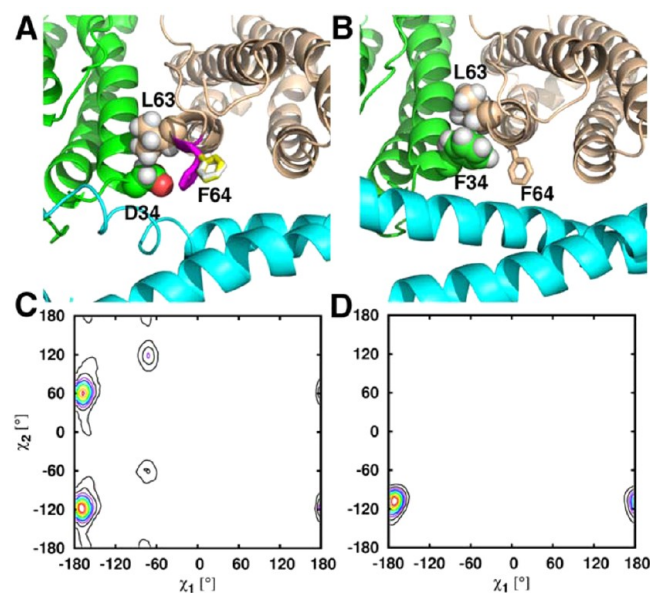


Figure 10. Close up view of a B-pore taken during the MD simulations of (A) wt and (B) D34F BfrB. The three rotameric states of the F64 side chain in wt BfrB are depicted by white, yellow, and magenta sticks in (A), and the three rotameric states of D34 are indicated in the plot shown in (C). The rotameric exchange of D34 and F64 contributes to the breathing motions of the B-pores as well as ion traffic across B-pores in wt BfrB. In the D34F structure packing of F34 against L63 (spheres) likely contributes to the lower flexibility of B-pores in the mutant, which is manifested in only one conformational rotamer of the F34 side chain (D) and only one conformation of the F64 side chain, wheat sticks in (B).

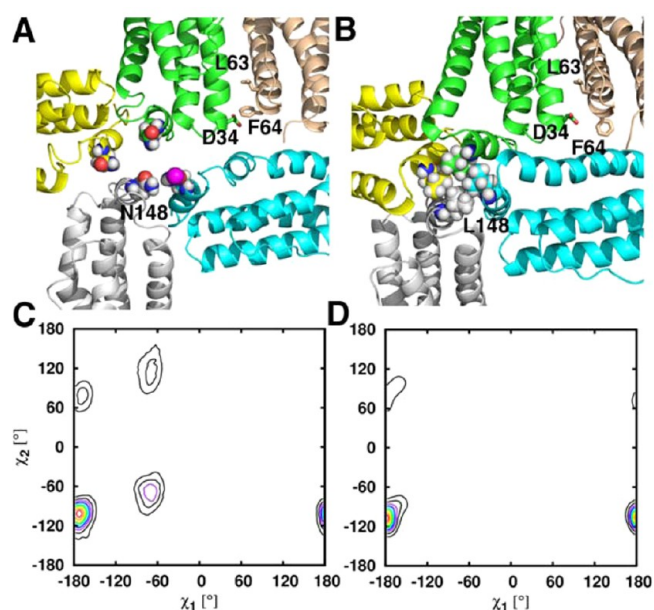


Figure 11. Snapshot obtained from MD simulations illustrating the packing of residues 148 in the 4-fold-pores of (A) wt and (B) N148L BfrB. The comparison of torsion angles sampled by F64 in the wt (C) and N148L (D) simulations are indicative of the dampened B-pore breathing motions observed in the N148L MD simulation relative to those observed in the wt BfrB simulation. Side chains of residue 148 in the 4-fold pores are rendered in spheres, whereas the side chains of D34, L63, and F64 in the B-pores are shown in stick representation.

Table 2. Number of K⁺ Ions Escaped from the BfrB Shell during the Simulations

	E2	E10	E40
WT	63	63	0
N148L	5	14	4
D34F	30	22	8

The dynamic conformational exchange exhibited by these side chains in wt BfrB contributes significantly to the breathing motions of the B-pores and facilitates the translocation of K⁺ across the B-pores (Table 2). In contrast, analyses of B-pores in the MD trajectories of D34F and N148L BfrB show a low-propensity for folding/unfolding transitions in the C-termini of helices D (Figures 10B and 11B), as well as dampened breathing motions of the B-pores (Movies S1–S3, Supporting Information). It is likely that the dampened motions in the B-pore of the D34F mutant stem from the hydrophobic packing between the F34 and L63 side chains (Figure 10B), which limits the conformational freedom of the F34 (Figure 10D) and F64 (Figure 10B) side chains to only one rotamer. In the N148L mutation, hydrophobic packing of L148 side chains inside the 4-fold pores likely stabilizes the interactions of the four E-helices that form a 4-fold pore (Figure 11B). As a result the C-terminal folding/unfolding transitions of helices D are nearly absent, as is the conformational freedom of the F64 side chains (Figure 11D). Thus, the packing between F34 and L63 in the B-pores of D34F, or the hydrophobic packing of L148 side chains in N148L, which stabilizes interactions of the four E-helices that form a 4-fold pore, largely dampens the folding/unfolding transitions of the C-terminal portions of helices D and the breathing motions of the B-pores. This has the effect of reducing the permeability of B-pores to K⁺ ions that cross the BfrB shell using these channels (Table 2).

Structural Perturbations in the Four-Fold and B-Pores Affect the Dynamics of Distant Ferroxidase Centers. In the MD simulations of wt BfrB, the conformational flexibility of the H130 side chains is evident in the average density of H130 side chain conformations sampled during the simulations (Figure 12A), which is located halfway between the gate-open (green) and gate-closed (cyan) conformations observed in the as-isolated and Fe-soaked crystal structures. In contrast, in the MD simulations of the D34F and N148L mutants the flexibility of helix D is lower, and the average (conformation) density of H130 side chains matches the gate-open conformation (Figure 12B,C). These observations, which suggest low conformational freedom of the H130 side chain in the mutants, are in excellent agreement with the findings from the as-isolated and Fe-soaked structures, where only the gate open

conformation is observed (Figure 7). Consequently, our results strongly suggest that flexibility in the C-terminal half of helix D in wt BfrB functions to couple breathing motions in 4-fold and B-pores with ferroxidase center ligands. It is likely that these networked motions facilitate efficient rearrangement of H130 side chains, thus enabling the gating function of this residue in Fe²⁺ capture and oxidation at the ferroxidase centers, and subsequent translocation of Fe³⁺ to the internal cavity for mineralization.

DISCUSSION

A Network of Molecular Fluctuations Connecting Four-fold Pores, B-Pores and Ferroxidase Centers Are Crucial to BfrB function. Bacterioferritins serve as dynamic regulators of cytosolic iron concentrations by performing two functions: (i) iron storage, which is carried out by capturing Fe²⁺ followed by its oxidation to Fe³⁺ and storage as a ferric-phosphate mineral, and (ii) iron release, which requires reduction of the Fe³⁺ mineral, followed by Fe²⁺ mobilization from the interior cavity. The 24-mer spherical assemblies (Ftn and Ftn-like molecules) are typically referred to as protein cages, where iron can be stored. Although a protein cage may suggest a relatively rigid shell that protects the ferric mineral from the reducing potential of the cytosol, recent evidence suggests that the Bfr shell is highly dynamic. Application of a network-weaving algorithm coupled to normal-mode analysis revealed long-range communication between ferroxidase centers and functionally related pores in Ftn-like molecules, which is mediated by a network of highly correlated residues, encompassing approximately 40% of residues in each subunit.¹⁰ In the case of eukaryotic Ftn (M Ftn), the network connects ferroxidase centers with 3-fold pores, an observation that is in agreement with the idea that the 3-fold pores of eukaryotic Ftn are conduits for iron traffic.^{26,27} In comparison, the network of correlated residues in bacterial Ftn and Bfr, connect ferroxidase centers with 4-fold and B-pores. MD simulations carried out with BfrB showed that K⁺ ions move across the protein shell through B-pores, in a process that is largely aided by breathing motions driven by the folding/unfolding of the C-terminal half of the D-helices, which are coupled to lateral displacements of the E-helices that constitute the 4-fold pores.²⁹ The lateral displacements of E-helices typically move two E-helices closer to one another, while the other two are separated further, thus narrowing the 4-fold pores. Although in the MD simulations the 4-fold pores do not permeate K⁺ ions, the lateral displacement of the E-helices, which is driven by the folding/unfolding transitions of the D-helices, enable periodic open/close transitions of the B-pores and facilitate K⁺ traffic. Hence,

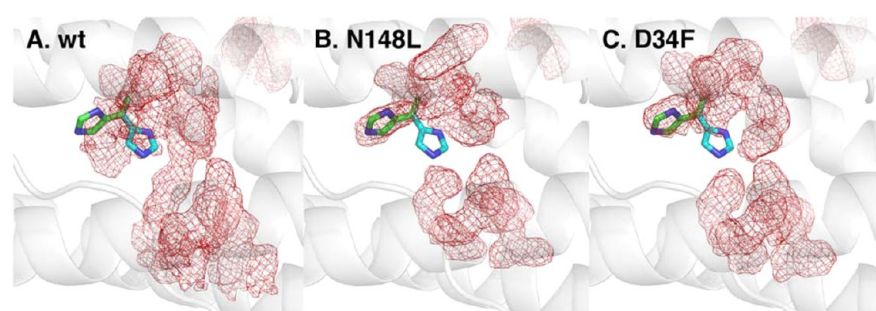


Figure 12. Average heavy atom density of side chain conformations sampled by ferroxidase center residues during the MD simulations is shown as red mesh for wt (A), N148L (B), and D34F BfrB (C). The H130 side chains in the “gate-open” (green) and “gate-closed” (cyan) conformations are shown in sticks. The view is from the exterior surface.

results obtained from coarse grain methods (normal-mode analysis) and atomistic MD simulations suggest a model in which the Bfr function is intimately related to extended networks of coordinated motions that affect a large portion of the protein shell.

In this work we tested the model with mutants designed to decrease pore flexibility: (i) The side chains of N148 and Q151, which coordinate a K^+ ion in the 4-fold pores of wt BfrB (Figure 2), were replaced with Leu to install hydrophobic interactions, which were expected to affect the lateral displacement of the E-helices. (ii) The D34F mutation was carried out with the expectation of occluding the B-pores and installing hydrophobic packing that would reduce B-pore flexibility. Structural determinations reveal local structural perturbations that confirm the installment of hydrophobic packing in the 4-fold and B-pores of the mutants. A surprising finding is the significant loss of Fe^{2+} oxidation (ferroxidase) efficiency exhibited by the mutants, particularly the D34F mutant, which oxidizes and stores only 20% of the iron presented. Remarkably, the Fe-soaked structures of D34F, N148L, and Q151L show that although iron loads onto the ferroxidase center, the diiron center is not coordinated by a full complement of ferroxidase ligands (Figure 7); E51 does not bridge Fe_1 and Fe_2 , and the H130 side chain remains in its gate open conformation. Taken together, the observations indicate that mutations in the 4-fold and B-pores reduce the conformational flexibility of the ferroxidase center, thus reducing its catalytic efficiency. In this context, it is noteworthy that mutations outside the pores (C89S/K96C) do not have an effect on the efficiency of iron oxidation and storage, or in the conformational flexibility of ferroxidase ligands, as evaluated by the conformational competency of E51 and H130 *in crystallo* (Figure 7A), and the unaffected efficiency of iron oxidation and uptake in solution exhibited by the double mutant.

We suspected that the reduced conformational flexibility of the ferroxidase centers in the 4-fold and B-pore mutants may be related to changes in the dynamic properties of the pores resulting from the hydrophobic interactions installed by the mutations. Consequently, in search of additional insight, we turned to MD simulations. As previously revealed by our simulations of wt BfrB, folding/unfolding transitions of the C-terminal half of helices-D drive the dynamic behavior of 4-fold and B-pores. Given that H130 is near the center of helix D, we asked if the folding/unfolding transitions of the helix affect the conformational flexibility of H130 side chains. The MD trajectory shows that in wt BfrB the average position of the H130 side chain is midway between the gate open and gate closed conformations (Figure 12A), a finding that is in agreement with the gating function of the H130 side chain in the binding of Fe^{2+} at ferroxidase centers, oxidation to a di- Fe^{3+} moiety, and subsequent entry of Fe^{3+} to the interior cavity.^{15,22} In contrast, the MD trajectories obtained with the D34F and N148L mutants reveal that helices D exhibit low propensity to undergo folding/unfolding transitions, which results in dampening of motions at the 4-fold and B-pores relative to wt BfrB. Analysis of the conformational freedom of H130 in the mutants showed that the average position of the H130 side chains in the simulations is aligned with the gate-open conformation (Figure 12B,C) observed in the crystal structures. Consequently, it is reasonable to conclude that in BfrB the 4-fold pores, their surrounding B-pores, and the ferroxidase centers are connected by networked fluctuations, and that these

motions enable the gating properties of the ferroxidase center and the ion permeation properties of the pores.

Iron Traffic Paths in BfrB. In previous work we showed that soaking crystals of wt BfrB in crystallization solution containing Fe^{2+} prior to X-ray diffraction reveal iron bound at the ferroxidase center, and iron bound at the periphery of 4-fold pores in the interior cavity, now termed Fe_{4f-1} (Figure 3A). Similar experiments conducted with each of the BfrB mutants allowed us to discover many additional iron binding sites in the interior surface of BfrB, and remarkably, within the B-pores. Visualizing all the iron sites simultaneously provides unprecedented insight into conduits used by iron as it traverses the BfrB shell, as well as the routes it may take on the surface of the interior cavity on its way to mineralization or cavity exit sites. It is noteworthy that all Fe sites except Fe_{4f-3} are observed in the structure C89S/K96C BfrB, which allows comparison of the relative thermal factors for the experimentally observed sites (Table 3), and Figure 13 depicts these iron sites as spheres,

Table 3. Thermal Factors of Iron Ions in the Fe Soak C89S/K96C BfrB Structure

iron site	location	B-factor range (\AA^2)	color in Figure 13
Fe_{heme}	heme	26–28	red
Fe_{4f-1}	4-fold pore	18–25	red
Fe_{4f-2}	4-fold pore	81–105	yellow
Fe_{3f}	3-fold pore	37–40	orange
Fe_{B1} , Fe_{B2}	B pore	45–63	orange
Fe_1 , Fe_2	fc ^a	18–30	red
Fe_{b-fc}	below fc	56–81	yellow

^afc: ferroxidase center.

color-coded by B-factor range. Since atomic occupancies and B-factors are highly correlated parameters,⁵⁶ Fe ions were refined with occupancy factors of 1.0 and the isotropic B-factors were examined in an effort to distinguish fully occupied and partially occupied/transient ions. Note that iron sites ubiquitously found in all Fe-soaked structures (Fe_1 , Fe_2 , and Fe_{4f-1}) have the lowest B-factors (red), which are comparable in magnitude to the protein and heme thermal factors. Together the observations suggest that iron at these sites is relatively stable and nearly 100% occupied. Iron in the B-pores (Fe_{B-1}/Fe_{B-2}) and iron near the entrance to 3-fold pores (Fe_{3f}) exhibit B-factors nearly twice as large (orange), which suggests these sites as more transient. Finally, the Fe_{b-fc} and Fe_{4f-2} sites (yellow) show thermal factors 4–5-fold larger than the protein B-factors, which suggests that these sites are only transiently occupied. To aid visualization of the full-complement of iron sites observed in this study, in Figure 13 we also include Fe_{4f-3} (orange), the iron site observed immediately above the entrance to 4-fold pores in Fe-soaked N148L BfrB (see Figure 3-C). The iron ions in Figure 13 are superposed on the structure of the Fe-soaked D34F mutant, which was selected because it is one of the structures where the H130 side chains remain in the gate open position even after iron loads onto the ferroxidase center. With the H130 side chain in the gate open position, it is possible to observe the ferroxidase iron ions Fe_1 and Fe_2 (red) within ferroxidase centers, and how the gate open conformation of H130 may enable translocation of Fe_2 into the interior cavity, where it binds to H46 at site Fe_{b-fc} (yellow). This is most evident in the ferroxidase centers located at the extreme left or top sections of Figure 13. Residues lining the B-pores are colored in cyan, and their inspection illustrates how B-pores

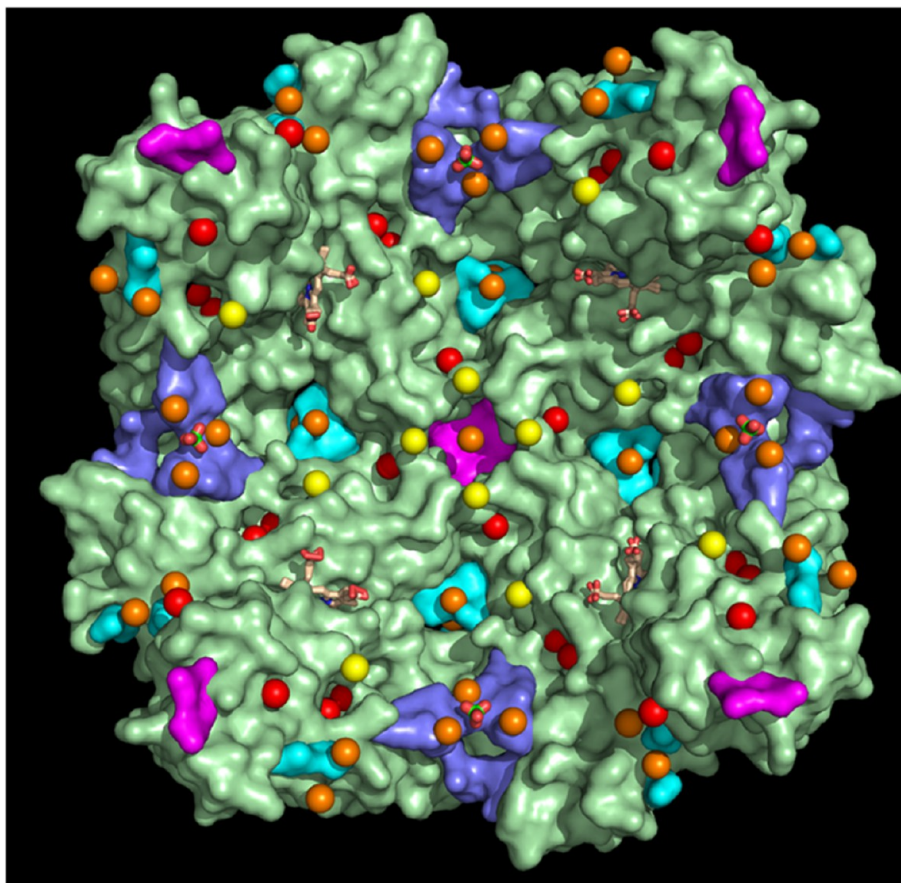


Figure 13. Transverse view of the BfrB interior cavity depicting all iron sites observed in the crystal structure of Fe-soaked C89S/K96C BfrB, color coded according to the corresponding *B*-factor ranges listed in Table 3; red (Fe_1 , Fe_2 , Fe_{4f1} , Fe_{heme}), orange (Fe_{3f} , Fe_{B-1} , Fe_{B-2}), and yellow (Fe_{4f2} , Fe_{B-fc}). The Fe_{4f3} iron, shown immediately on top of the 4-fold pore in the center of the figure is from the Fe-soaked N148L structure, rendered in orange to reflect the range of thermal factors exhibited by these iron ions ($31\text{--}54\text{ \AA}^2$) relative to those in Table 3. Residues lining 4-fold pores are in magenta, residues lining 3-fold pores are in blue and those lining B-pores are in cyan. Sulfate ions are in green and red sticks, and heme molecules are rendered as wheat sticks.

may function as conduits for iron traffic across the Bfr shell, with two iron ions (Fe_{B-1} , Fe_{B-2} , orange) aligned along the length of the channels (see also Figure 6E). Residues delineating the 4-fold pores are colored in magenta. The 4-fold pore at the center of the figure illustrates the three layers of iron binding sites located around the external perimeter of the pore inside the BfrB cavity (see also Figure 3): (1) Four Fe_{4f1} ions (red) surround the largest diameter of the pore, where the pore wall meets the interior surface. (2) Four Fe_{4f2} ions (yellow) surround the narrowest perimeter of the pore, where the pore wall meets the interior cavity. (3) One Fe_{4f3} ion (orange) is located immediately above the entrance to the pore from the interior cavity. Remarkably, iron ions are not observed inside 4-fold pores, which in our structures always bind K^+ , with the exception of the pores in the Q151L structure. Residues lining the 3-fold pores are blue. Note how each of the sulfate ions at the entrance to each of the pores from the interior cavity (green and red sticks) is surrounded by three Fe_{3f} ions (orange; see also Figure 4).

The picture that emerges from this analysis suggests that iron ions can enter the BfrB cavity via ferroxidase centers or through B-pores. Iron processed through ferroxidase centers will likely enter the cavity as Fe^{3+} , which may then be transported by coordinating residues on the interior surface to mineralization sites, or toward 3-fold pores where it binds phosphate ions

accessing the interior cavity via 3-fold pores, prior to mineralization. Iron entering through B-pores is likely in the ferrous oxidation state. Given the large number of B-pores (24 in total) on the Bfr structure, Fe^{2+} accessing the interior cavity via B-pores would be in relative close proximity to a ferroxidase center, where it can be oxidized by electron exchange with Fe^{3+} bound to the ferroxidase sites, or, at the growing Fe^{3+} mineral, as has been proposed for eukaryotic Ftn⁵⁷ and *E. coli* Bfr.⁵⁸ It is noteworthy that in spite of the cumulative large number of iron binding sites we have been able to observe, we have not yet found evidence of iron ions inside the 4-fold pores of BfrB. Instead, the structures reveal several iron ions bound to the external perimeter of these pores inside the BfrB cavity, which could have reached those sites after accessing the interior cavity via B-pores. Hence, although we cannot rule out the possibility that iron ions traverse BfrB using 4-fold pores, it is tempting to speculate that the 4-fold pores in BfrB function mainly as binding sites for Fe^{2+} in the interior cavity, prior to oxidation at ferroxidase centers, or as nucleation sites for incorporation of iron ions into a growing Fe^{3+} mineral. The data also suggest that iron and phosphate combine near the 3-fold pores, which seem to be the channels used by the anion to traverse the BfrB shell.

In conclusion, our approach, which combined biochemistry with X-ray crystallography and MD simulations, provides

unprecedented insight into Bfr function. Our findings demonstrate that a vast network of concerted motions connecting 4-fold and B-pores with ferroxidase centers is crucial for efficient ferroxidase activity and for permeation of ions across the Bfr cavity. Our results also reveal a large number of iron binding sites in the interior cavity, which aid in the traffic, oxidation, and mineralization of the iron, and that the 3-fold pores are the likely conduits of phosphate, which is necessary to form the ferric phosphate mineral core in Bfr.

■ ASSOCIATED CONTENT

■ Supporting Information

Phased anomalous difference maps of the different iron sites in observed in the structures of Fe-soaked D34F, N148L, Q151L, and C89S/K96C BfrB are shown in Figures S1–S3 and S5–S7. $2F_o - F_c$ maps for sulfate ions in the 3-fold pores of Fe-soaked N148L, Q151L, and C89S/K96C BfrB are shown in Figure S3, and $2F_o - F_c$ maps of sulfate ions in the 3-fold pores of as-isolated-2 C89S/K96C BfrB are depicted in Figure S4. Anomalous peak heights and B-factors for iron atoms in the Fe-soaked structure of C89S/K96C structure are listed in Table S1, anomalous peak heights and B-factor ranges for iron atoms in the structures of Fe-soaked D34F, N148L, and Q151L BfrB are listed in Table S2, and distances between ferroxidase center ligands and ferroxidase iron ions in the structures of Fe-soaked WT, C89S/K96C, D34F, N148L, and Q151L BfrB are listed in Table S3. Sequence of primer pairs used to introduce the site directed mutations. This material is available free of charge via the Internet at <http://pubs.acs.org>.

Accession Codes

Coordinates and crystallographic structure factors for the distinct BfrB structures have been deposited in the Protein Data Bank under accession codes listed in Table 1.

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Notes

The authors declare no competing financial interest.

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