



Structure of the Large Extracellular Loop of FtsX and Its Interaction with the Essential Peptidoglycan Hydrolase PcsB in Streptococcus pneumoniae

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ABSTRACT Streptococcus pneumoniae is a leading killer of infants and immunocompromised adults and has become increasingly resistant to major antibiotics. Therefore, the development of new antibiotic strategies is desperately needed. Targeting bacterial cell division is one such strategy, specifically by targeting proteins that are essential for the synthesis and breakdown of peptidoglycan. One complex important to this process is FtsEX. FtsEX comprises a cell division-regulating integral membrane protein (FtsX) and a cytoplasmic ATPase (FtsE) that resembles an ATP-binding cassette (ABC) transporter. Here, we present nuclear magnetic resonance (NMR) solution structural and crystallographic models of the large extracellular domain of FtsX, denoted extracellular loop 1 (ECL1). The structure of ECL1 reveals an upper extended β -hairpin and a lower α -helical lobe, each extending from a mixed α - β core. The helical lobe mediates a physical interaction with the peptidoglycan hydrolase PcsB via the coiled-coil domain of PcsB (PscB_{CC}). Characterization of S. pneumoniae strain D39-derived strains harboring mutations in the α -helical lobe shows that this subdomain is essential for cell viability and required for proper cell division of S. pneumoniae.

IMPORTANCE FtsX is a ubiquitous bacterial integral membrane protein involved in cell division that regulates the activity of peptidoglycan (PG) hydrolases. FtsX is representative of a large group of ABC3 superfamily proteins that function as "mechanotransmitters," proteins that relay signals from the inside to the outside of the cell. Here, we present a structural characterization of the large extracellular loop, ECL1, of FtsX from the opportunistic human pathogen *S. pneumoniae*. We show the molecular nature of the direct interaction between the peptidoglycan hydrolase PcsB and FtsX and demonstrate that this interaction is essential for cell viability. As such, FtsX represents an attractive, conserved target for the development of new classes of antibiotics.

KEYWORDS NMR structure, *Streptococcus pneumoniae*, cell division, peptidoglycan hydrolases, protein-protein interactions

Streptococcus pneumoniae is a Gram-positive, opportunistic respiratory pathogen (1-3) that has acquired antibiotic resistance worldwide (4-6). This ovococcal bacterium relies on highly conserved cell wall machinery to divide and grow (7, 8). The cell wall is primarily composed of peptidoglycan (PG), a macromolecule composed of repeating subunits of N-acetylglucosamine and N-acetylmuramic acid linked by PG peptide side chains (9, 10). Regulation of the synthesis and remodeling of PG is essential

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for bacterial growth and viability, due to the turgor pressure bacterial cells must withstand (10-12). One vital process for the synthesis of PG is the controlled insertion of new strands of PG. This process requires timed cleavage of the old PG matrix to allow incorporation of new nascent strands (13). PG hydrolases are the primary enzymes that carry out PG cleavage and remodeling (14, 15). Thus, regulation of these hydrolases and activation at specific times during the cell cycle is required for proper cell growth. Specific protein complexes are utilized by bacterial cells to regulate these enzymes. This work focuses on understanding the structure and function of one of these protein complexes.

From Mycobacterium tuberculosis to Caulobacter crescentus, the ATP-binding cassette (ABC) transporter-like protein complex FtsEX acts as a key regulator of PG hydrolysis and divisome assembly (16-19). The proposed mechanism of FtsEX activation of PG hydrolases is as follows. FtsE, upon sensing an unknown signal from inside the cell, hydrolyzes ATP to ADP. Hydrolysis causes a conformational change that is transmitted through the membrane via FtsX, an integral membrane protein with two extracellular loops (ECLs), denoted the large (ECL1) and small (ECL2) loops. These extracellular loops interact with either cell wall hydrolases or effector proteins, which results in activation of PG hydrolysis via an unknown mechanism (16, 18, 20-25). In Escherichia coli, it has been demonstrated that FtsX interacts with the effector protein EnvC to activate the PG amidases AmiA and AmiB (24, 25). In addition, FtsX interacts with other division proteins, such as FtsA, where it regulates the polymerization of FtsA and recruitment of downstream division proteins (26). In other organisms, including Bacillus subtilis and M. tuberculosis, FtsEX also activates PG hydrolases (16, 23). Interestingly, FtsEX is nonessential in rod-shaped bacteria like E. coli and B. subtilis (23, 24, 26-28). However, in S. pneumoniae, FtsEX is absolutely essential (21) and depletion of FtsEX results in cell rounding and cessation of growth (20, 21).

In the case of S. pneumoniae, genetic experiments suggest that both outward-facing domains of FtsX, ECL1 and ECL2, interact with the essential PG hydrolase PcsB via its long coiled-coil (CC) domain (20, 21). However, there is little direct biochemical evidence for this interaction. ECL1 and ECL2 are thought to allosterically activate the catalytic activity of the cysteine, histidine-dependent amidohydrolase/peptidase (CHAP) domain of PcsB (20). The crystal structure of full-length PcsB, including the CC domain, an alanine-rich linker region, and the CHAP domain, provides insight into the mechanism of how this may occur (22). While the PcsB structure implies that FtsEX activates PcsB by displacing the catalytic domain from the CC domain, the exact nature of the FtsX-PcsB interaction remains unknown.

In order to understand how FtsX activates PcsB, we determined the structure of the large extracellular loop of FtsX (FtsX_{ECL1}) by both multidimensional nuclear magnetic resonance (NMR) spectroscopy and X-ray crystallography. FtsX_{ECL1} harbors a conserved mixed α - β core and a lower α -helical lobe extending from the core, identified previously in M. tuberculosis FtsX (16), and S. pneumoniae FtsX_{ECL1} has a unique extended β -hairpin. The N-terminal β 1 and C-terminal β 6 strands are adjacent in the core and connect ECL1 to the transmembrane 1 (TM1) and TM2 helices, respectively, in the membrane. PscB_{CC}-mediated chemical shift perturbations of spectra obtained by ¹H-¹⁵N heteronuclear single quantum coherence (HSQC) spectroscopy of FtsX_{ECL1} reveal that the helical lobe consisting of the $\alpha 2$ helix and the α 2- β 5 linker (residues 107 to 134) of FtsX_{ECL1} interacts with PscB_{CC}. To determine if this interaction is required for FtsX function in bacterial cells, we constructed a merodiploid strain that allows for conditional expression of mutant ftsX. We demonstrate that specific amino acid substitutions in the FtsX-PcsB interface are lethal or cause pronounced morphological defects despite the fact that these FtsX_{ECL1} mutant proteins are expressed at nearly wild-type levels. These findings support the model that a direct physical interaction between FtsX and PcsB is required for activation of PcsB PG hydrolytic activity.



RESULTS

The three-dimensional structure of $FtsX_{ECL1}$. The three-dimensional structure of FtsX_{FCI.1} (residues 46 to 168) was solved by both NMR spectroscopy (Fig. 1A) and X-ray crystallography (Fig. 1B). The folded structure (residues 57 to 166) reveals a central core composed of a four-stranded antiparallel β -sheet (β 1, β 6, β 4, and β 5) and two helices (α 1 and α 3), an α -helical lobe (residues 107 to 135) harboring the α 2 helix, and an extended β -hairpin (β 2 and β 3). The β -hairpin and helical lobes are connected to the central core by hinges. Details for structural determination of FtsX_{FCI,1} by NMR in the absence of detergent are presented in Materials and Methods, and structure statistics are summarized in Table 1. The solution structure shows that while the central mixed α - β core adopts a well-defined conformation, the two appended lobes are highly dynamic on multiple timescales (see below), presenting a range of conformations among the 20 members of the $FtsX_{ECL1}$ NMR structural ensemble (Fig. 1A).

Three different structures, with resolutions ranging from 2.0 to 2.3 Å, were solved by X-ray crystallography, each in the presence of a different detergent. The detergents used were dodecyltrimethylammonium chloride (detergent 1 [D1]), *n*-undecyl-β-Dmaltoside (detergent 2 [D2]), and *n*-decyl-β-D-maltoside (detergent 3 [D3]). The presence of detergents was critical, as in their absence, crystals diffracted at very low resolution (≤4 Å), suggesting significant mobility in some protein regions. Details from the crystallographic determination are provided in Materials and Methods, and structure statistics are summarized in Table 2. In all cases, two independent protein molecules are present in the asymmetric unit (Fig. S1 in the supplemental material), arranged in such a way as to keep hydrophobic residues from their α -helical lobes buried. As explained below, the α -helical-lobe region is relevant for PcsB interaction. In total, six independent structures were determined for both the protein structure and associated detergent molecules (Fig. S2). Different conformations were observed for the β -hairpin and the helical lobe among the six structures, depending on the presence and identity of the detergent molecule bound to FtsX_{ECL1} (Fig. S1B). The structural variations observed in these crystallographic structures, however, are less dramatic than those observed in the NMR conformer bundle obtained in the absence of ligand (Fig. 1E). As expected, regions presenting structural variations in crystal structures correspond to those showing a highly dynamic behavior by NMR.

In any case, the crystal structures suggest that changes in the protein backbone and side chains of the helical lobe occur when a detergent ligand is bound (Fig. 1C). These changes create a cavity in which the detergent molecules insert (Fig. 1D), with a large part of the helical lobe (from Q111 to E127) affected by the interaction with detergent (Fig. 1C). Residues Y112, W123, E127, and F126 are strongly perturbed upon detergent ligand binding (Fig. 1C), with the hydroxyl moiety of Y112 interacting with carboxylate of E127. Changes in W123 and F126 stabilize the hydrophobic region of detergent 1. A similar interaction pattern is observed for detergents 2 and 3; additional interactions are observed in N131 with detergents 2 and 3, which are characterized by a larger hydrophilic head (maltose) (Fig. S1E and F). Full stabilization of the hydrophobic tail of the detergents is completed by the same hydrophobic residues (W123 and F126) but from a symmetry-related molecule (Fig. S1G). Although a physiological role for detergent binding to the helical lobe is unknown, many of these same residues are important for the interaction with PcsB (see below).

A structural comparison with the M. tuberculosis FtsX_{ECL1} (PDB code 4N8N) (16) reveals differences in both the overall structure (root mean square deviation [RMSD] of 2.2 Å) and the appended lobes of the core domain (Fig. 1F). The main differences between the mycobacterial and pneumococcal FtsX_{ECL1} domains are the presence of an extra helix (α 1) and a disulfide bond in the *M. tuberculosis* ECL1 that are absent in the pneumococcal ECL1 and an extended β -hairpin (residues 71 to 87) that is unique to the pneumococcal ECL1 domain. Of note, this β -hairpin was conserved among aligned streptococcal species sequences (see Fig. S7A for species list), with β 3 being more conserved than β 2. It is also worth noting that in the M. tuberculosis ECL1, the upper

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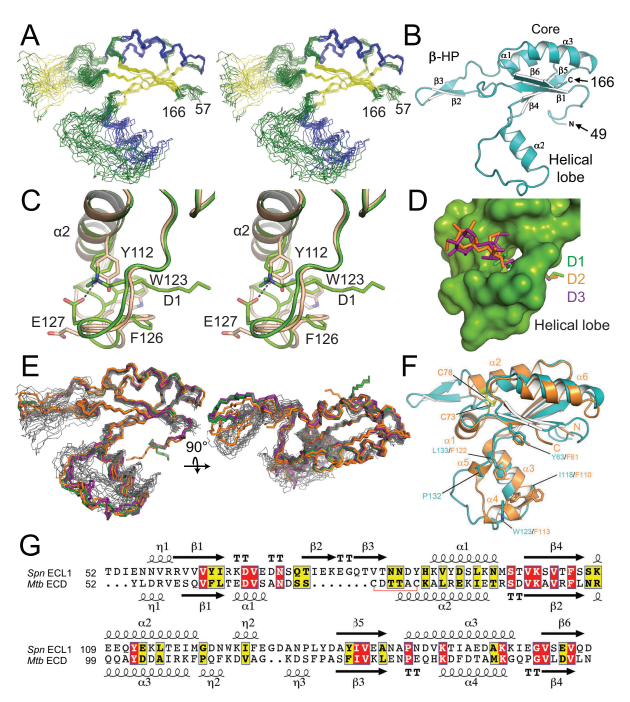


FIG 1 The structure of FtsX_{ECL1} from Streptococcus pneumoniae. (A) Stereoview of the 20 conformers of the FtsX_{ECL1} NMR structure as backbone traces, with helices shown in blue and β -strands shown in yellow. N and C termini of the domain are indicated by the residue numbers 57 and 166. (B) Cartoon representation of the $FtsX_{ECL1}$ structure obtained by X-ray crystallography (chain B in $FtsX_{ECL1}$ -D1) in which the different secondary structure elements have been numbered and labeled. N and C termini of the domain are indicated by the residue numbers 49 and 166. β-HP, β-hairpin. (C) Stereoview showing changes in helical lobe upon interaction with dodecyltrimethylammonium chloride. Apo form (chain B) is colored in pale brown, and holo form (chain A) is colored in green. Relevant residues affected by the presence of the detergent are depicted as sticks. The polar interaction is represented by a dashed line. (D) Surface representation of the FtsX_{ECL1} crystal structure in which the three different detergent molecules are superimposed and shown in sticks, as follows: dodecyltrimethylammonium chloride, n-undecyl- β -p-maltoside, and n-decyl- β -p-maltoside are colored in green, orange, and purple, respectively. (E) Overlay of the backbone traces for the six FtsX_{ECL1} crystal structures (colored in green, orange, and purple in the presence of dodecyltrimethylammonium chloride, *n*-undecyl-β-Dmaltoside, and n-decyl- β -p-maltoside, respectively) with the backbone traces of the 20 FtsX $_{ECL1}$ NMR conformers (in gray). (F) Cartoon representation of the FtsX_{ECL1} crystallographic structure (in cyan) overlaid on the M. tuberculosis FtsX_{ECD} structure (in orange) (PDB code 4N8N). Cysteine residues involved in disulfide bond formation are shown as yellow sticks and indicated with an arrow. (G) Structure-based sequence alignment (54) of extracellular domains from S. pneumoniae (Spn ECL1) (PDB code 6HFX, chain A) and M. tuberculosis (Mtb ECD) (PDB code 4N8N, chain B). Secondary structure elements from each protein are indicated and numbered.



TABLE 1 Structural statistics for NMR solution structure of FtsX_{ECL1}^a

Parameter	Value
NMR distance and angle restraints	
Total NOE-based distance constraints	1,711
Intraresidue $(i = j)$	232
Sequential $(i - j = 1)$	499
Medium range $(1 < i - j < 5)$	365
Long range $(i-j \ge 4)$	615
Maximum distance violation (Å)	0.49
Dihedral angle restraints	110
Maximum dihedral angle violation (°)	7
Total number of RDCs used (measured)	82 (112)
Q factor	0.11 (0.39)
Correlation (experimental to calculated)	0.99 (0.89)
RMS deviations from idealized covalent geometry	
Covalent bond lengths (Å)	0.011
Covalent angle values (°)	1.3
Ramachandran analysis (%) ^{b,c}	
Most favored	86.5
Additionally allowed	8.4
Generously allowed	1.8
Disallowed	3.3
RMSD values (Å)	
Backbone atoms (all)	1.5
All heavy atoms (all)	1.9
Backbone atoms (core ^b)	0.4
All heavy atoms (core ^b)	0.9

^aFrom the ensemble of the 20 best NMR structures. PDB code 6MK7; BMRB code 30523.

and lower lobes form a large hydrophobic cleft with four exposed Phe residues (F61, F110, F113, and F122), and this region was suggested as a strong candidate for the interaction surface between FtsX and PG hydrolase RipC (16). These phenylalanine residues are not conserved in the pneumococcal ECL1 (Fig. 1F and G), but the hydrophobic nature of this region is preserved (Y63, I118, W123, and L133).

FtsX_{ECL1} is dynamic in solution. We also performed additional NMR experiments to explore the mobility of the FtsX_{FCL1} domain, both to validate the heterogeneity of the structural ensemble in solution and to elucidate function. The ${}^{1}D_{HN}$ residual dipolar couplings (RDCs) obtained by weak alignment in Pf1 filamentous phage correspond well to previously determined secondary structure elements (29), with uniform values for the entire length of $\alpha 1$ and $\alpha 2$, as expected for straight helices. In contrast, ${}^{1}D_{HN}$ values near 0 for the N-terminal tail, the very C terminus, and the nonhelical part of the helical lobe between $\alpha 2$ and $\beta 5$ are suggestive of significant conformational disorder in solution (Fig. S3A). These regions of small or zero RDCs are regions of very high RMSD in the NMR structure bundle (Fig. S3B). As anticipated, the correlation between experimentally measured RDCs and predicted RDCs back-calculated from the structures (30) is high, but only in the core subdomain, and the experimental and predicted RDCs match poorly in the β -hairpin for most of the crystal structures (Fig. S3C and D).

We previously reported that the ¹⁵N-{¹H} heteronuclear nuclear Overhauser effect (hNOE) is low or negative at the termini, indicating that they are highly flexible in solution (29). The hNOE is also smaller in the β 2- β 3 hairpin region, as well as in the C-terminal end of the $\alpha 2$ helix and the subsequent coiled region leading into $\beta 5$ (Fig. S3E) (29). Mapping these dynamics data onto a representative structure from the solution NMR ensemble shows that these regions with fast-timescale dynamics correspond to the β -hairpin and the helical lobe (Fig. S3F). Information on picosecond-to nanosecond (ps-ns) fast-timescale motions extracted from the R_2/R_1 ratio also reveals

 $[^]b$ Calculated over the core of the structure, excluding the helical lobe and β -hairpin, residues 59 to 65, 89 to 108, and 134 to 165.

^cAs computed by Procheck.

TABLE 2 Crystallographic data collection and refinement statistics

	Values for ^a :			
Parameter	FtsX _{ECL1} -D1	FtsX _{ECL1} -D2	FtsX _{ECL1} -D3	
Data collection statistics				
Wavelength (Å)	0.97934	0.97934	0.97934	
Space group	P 4 ₃ 2 ₁ 2	P 4 ₃ 2 ₁ 2	P 4 ₃ 2 ₁ 2	
Unit cell dimensions				
а, b, с (Å)	75.87, 75.87, 97.93	84.61, 84.61, 106.19	75.49, 75.49, 95.31	
α, β, γ (°)	90, 90, 90	90, 90, 90	90, 90, 90	
Temp (K)	100	100	100	
X-ray source	Synchrotron	Synchrotron	Synchrotron	
Resolution range (Å)	47.05-(2.05-2.0)	44.97-(2.38-2.3)	46.58-(2.23-2.16)	
No. of unique reflections	19,992 (1,436)	17,781 (1,700)	15,360 (1,496)	
Completeness (%)	100 (100)	100 (100)	99.85 (99.93)	
Multiplicity	24.4 (25.9)	22.1 (22.5)	16.6 (21.8)	
$R_{pim}{}^{b}$	0.013 (0.535)	0.007 (0.123)	0.010 (0.241)	
/i	21.9 (1.5)	41.7 (4.6)	24.6 (2.55)	
CC1/2 ^c	1.00 (0.67)	1.00 (0.98)	1.00 (0.89)	
Refinement statistics				
Resolution range (Å)	41.14-2.0	42.31-2.3	46.58-2.16	
$R_{\text{work}}/R_{\text{free}}^{d}$	0.24/0.28	0.26/0.30	0.25/0.31	
No. of atoms				
Protein	1,888	1,732	1,856	
Water	25	21	23	
Ligand	16	47	33	
RMS deviations				
Bond length (Å)	0.008	0.009	0.010	
Bond angles (°)	1.27	1.22	1.29	
Ramachandran plot				
Favored/outlier regions (%)	95.65/0.43	93.20/2.43	92.41/0.00	
Monomers per AU	2	2	2	
PDB code	6HE6	6HEE	6HFX	

aValues between parentheses correspond to the highest-resolution shells.

that the $\alpha 2-\beta 5$ linker is highly flexible, while the $\beta 1-\beta 2$ linker and β -hairpin show elevated R_2/R_1 ratios, specifically indicative of slow, millisecond (ms) timescale conformational exchange (Fig. S3G and H). These findings were directly confirmed by Carr-Purcell-Meiboom-Gill (CPMG) relaxation dispersion NMR spectroscopy (Fig. S3I and J) (31). We conclude that the β -hairpin exhibits flexibility on both the sub-ns and ms timescales. Interestingly, the β 1- β 2 linker also shows increased B factors that are qualitatively consistent with the ms timescale conformational exchange observed by NMR.

These complex motions observed in the solution dynamics experiments are reflected in the heterogeneity of the NMR structure, with high $C\alpha$ RMSDs particularly in the $\alpha 2-\beta 5$ linker but also in the β -hairpin and $\beta 1-\beta 2$ linker, thus validating the conformational spread in the ECL1 structure in solution (Fig. S3B). The dynamic nature of the helical lobe is also reflected in the heterogeneity and B factors of the crystal structures (Fig. S3K and L). Full-length FtsX itself is likely a dimer in vivo (32), and one can speculate that the flexible helical-lobe and β -hairpin regions may contribute to dimerization or to interactions with a binding partner like PcsB.

The PscB_{CC} domain interacts with FtsX_{ECL1}. The ¹H-¹⁵N HSQC spectrum for ECL1 has excellent chemical shift dispersion and lends itself readily to studies of proteinprotein interactions (Fig. S4A). In contrast, full-length PcsB is 42 kDa and forms a dimer

 $[^]bR_{
m pim}$ measures the precision of averaged intensities. $R_{
m pim} = \Sigma_{hkl} [1/(N-1)] \ 1/2 \ \Sigma_i \ | \ I_i(hkl) - I_i(hkl) \ | \ I_i(hkl) \ |$

 $[[]l(hkl)] \mid /\Sigma_{hkl}\Sigma_{l}(hkl)$, where $\Sigma_{l}(hkl)$ is the ith measurement of reflection hkl, [l(hkl)] is the weighted mean of all measurements, and N is the redundancy for the hkl reflection.

cCC1/2 is the correlation coefficient between intensity estimates from half data sets.

 $^{{}^}dR_{\mathrm{work}}/R_{\mathrm{free}} = \sum_{hkl} |F_o - F_c| / \sum_{hkl} |F_o|$, where F_c is the calculated and F_o is the observed structure factor amplitude of reflection hkl for the working/free (5%) set, respectively.



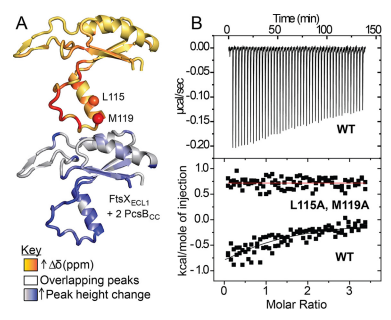


FIG 2 FtsX_{FCI 1} binds PscB_{CC}. (A) Significant chemical shifts and peak height changes upon ¹H-¹⁵N HSQC titration of 50 μ M 15 N FtsX $_{ECL1}$ with 100 μ M unlabeled PscB $_{CC}$ map to the lower lobe of FtsX $_{ECL1}$. Chemical shifts ($\Delta\delta$, ppm) and peak height changes are mapped as color gradients on the FtsX_{ECL1} structure, orange to red and light gray to blue, respectively. L115 and M119 lpha carbons are shown as spheres on the upper image. Peaks that overlap in the ¹H-¹⁵N HSQC spectra are colored white. Chemical shifts and peak height changes upon the addition of 2 molar equivalents of $PscB_{CC}$ to $FtsX_{ECL1}$ are mapped to the structure. (B) Representative titration of $PscB_{CC}$ with wild-type (WT) $FtsX_{ECL1}$ or L115A, M119A $FtsX_{ECL1}$ as monitored by ITC. Conditions used were 50 mM potassium phosphate, 50 mM NaCl, 0.5 mM EDTA, pH 7.0, at 25.0°C. Top, corrected ITC data; bottom, kcal/mole of injection versus time. The black line overlapping the WT data indicates the best fit to a one-site binding model. Fitting parameters are summarized in Table 3. The red line drawn through the L115A M119A $FtsX_{ECL1}$ data is for reference.

and thus is challenging to study by NMR due to its size. We therefore constructed truncation mutants of PcsB, focusing on the coiled-coil domain (PscB_{CC}), thereby limiting the molecular weight to 23 to 24 kDa. ¹⁵N-labeled PscB_{CC}(47–267) (comprising residues 47 to 267) and $PscB_{CC}(47-254)$ (comprising residues 47 to 254) are both characterized by ¹H-¹⁵N HSQC spectra with limited ¹H signal dispersion (Fig. S6A), consistent with the high helical content of this domain. Circular dichroism spectroscopy confirms that these proteins are primarily α -helical, in agreement with the crystal structure (22), indicating that they are properly folded (Fig. S6A) and can be used for ECL1 binding studies.

To determine if PcsB physically interacts directly with FtsX_{ECL1}, we titrated unlabeled $PscB_{CC}(47-267)$ into ¹⁵N-labeled FtsX_{ECL1} at molar ratios of 1:1, 2:1, 4:1, and 6:1 and recorded the ¹H-¹⁵N HSQC spectra (Fig. S4A and B). Numerous crosspeaks move in response to the addition of $PscB_{CC}$ to ^{15}N -labeled $FtsX_{ECL1}$. The largest changes occur in the helical lobe of FtsX_{ECL1} (Fig. 2A, Fig. S4 and S5). In particular, residues M119, W123, I125, F126, and G128 exhibit the greatest changes in crosspeak position and intensity in the sample with a 1:2 molar ratio of FtsX_{ECL1} to PscB_{CC} (Fig. 2A, Fig. S4C); when additional PscB_{CC} is added, these crosspeaks broaden beyond detection (Fig. S5B and C). A reciprocal ¹H-¹⁵N HSQC spectroscopy experiment with ¹⁵N-labeled PscB_{CC}(47-254) further confirms an interaction with unlabeled FtsX_{ECL1} (Fig. S6B), as multiple crosspeaks shift upon the addition of increasing amounts of FtsX_{ECL1} (Fig. S6B). We measured the binding affinity of the $FtsX_{ECL1}$ - $PscB_{CC}$ complex using isothermal titration calorimetry (ITC), which reveals an association equilibrium constant (K_a) of 3.0 \times 10⁴ ${\rm M}^{-1}$ (${\rm K}_d \sim 34~\mu{\rm M}$) (Table 3, Fig. 2B). These data provide the first biochemical evidence for a direct physical interaction between $PscB_{CC}$ and $FtsX_{ECL1}$ (Fig. 2, Fig. S4 to S6).

The interaction region of FtsX_{ECL1} with PscB_{CC} is essential for cell growth and **proper morphology.** Having identified the interaction region between FtsX_{FCI.1} and

TABLE 3 Thermodynamic parameters of wild-type and mutant FtsX_{FCL1} from direct analysis of isothermal titration calorimetry^a

Protein	$K_a (\times 10^4 \text{ M}^{-1})$	$K_d (\mu M)^b$	ΔH (kcal mol $^{-1}$)	ΔS (kcal mol ⁻¹ K ⁻¹)
FtsX _{ECL1}	3.0 ± 0.5	34 ± 6	-2.1 ± 0.2	13.4
L115A M119A FtsX _{FCL1}	NA^c	NA^c	NAc	NAc

^aSolution conditions were 50 mM sodium phosphate, 50 mM NaCl, 0.5 mM EDTA, pH 7.0, at 25°C. The chemical model used to fit the data is indicated in supplemental materials and methods in Text S1.

PscB_{CC}, we next sought to determine the degree to which this interaction interface mapped by NMR spectroscopy contributes to pneumococcal viability. A multiplesequence alignment of this region (residues 102 to 155) among bacterial species in which FtsX has been studied and in related streptococcal species (Fig. S7A) reveals that amino acids in this region are either partially or completely conserved (Fig. S7A). We therefore decided to target E109, Q111, L115, M119, W123, F126, and N131 for substitution with alanine, singly or in combination (Fig. S7A). Given the essentiality of the interaction of FtsX_{ECL1} and PscB_{CC}, we predicted that mutating these residues might be lethal (20). To allow for the cross-in of potentially lethal point mutations, we employed the Janus cassette method to insert point mutations at the native site of ftsX into a strain containing an ectopic copy of $ftsX^+$ under a zinc-inducible promoter (Fig. S7B) (33). We then transformed markerless mutant alleles of ftsX in the presence of zinc. This allows for expression of the wild-type $ftsX^+$ and mutant ftsX simultaneously. As long as the mutant ftsX was not dominant negative, we could obtain a strain that expresses the wild-type copy of ftsX+ under zinc induction and mutant ftsX only in the absence of zinc (Fig. S7B).

Zinc toxicity has been observed to cause aberrant cell morphology and growth inhibition in S. pneumoniae when cells are not supplemented with manganese (34, 35). To rule out any deleterious effects of the zinc-and-manganese (Zn-Mn) addition used to induce ftsX expression, we measured the growth of the parent and the FtsX merodiploid strain in the presence of these metals. To verify that the addition of 0.45 mM ZnCl₂ and 0.045 mM MnSO₄ (Zn-Mn) did not cause growth or morphological defects, cells were grown in the presence and absence of Zn-Mn and imaged at 3 h and 6 h into the growth curve (Fig. S7C and D). Wild-type cells (strain D39 Δcps rpsL1) had no morphological or growth defects at these time points with or without the addition of Zn-Mn

In contrast, the FtsX merodiploid strain (P_{Zn} -ftsX Δ ftsX) had significant morphological and growth defects at 3 or 6 h in the absence of Zn-Mn (Fig. S7C to E). Cessation of growth and aberrant cell morphology were observed in 90% of cells at 3 h and 95% of cells at 6 h growth (Fig. S7D). These cells were significantly shorter and rounder than wild-type cells (Fig. S7E), and a large variability in their volumes was observed (Fig. S7E), as previously found for a strain expressing ftsX+ under a fucose-inducible promoter (21). If the strain was grown in the presence of Zn-Mn, FtsX was expressed and the strain had no growth or morphological defects (Fig. S7C to E). This indicates the defects observed were solely due to the absence of FtsX.

We next constructed three classes of amino acid substitution or insertion mutants (Table 4) in an effort to disrupt the FtsX_{FCL1}-PscB_{CC} interaction defined by NMR spectroscopy. These are designated class I (single amino acid changes), class II (multiple amino acid changes), and class III [insertion of a (Gly₃Ser)₂ linker] mutants. Class I strains were made by introducing single-amino-acid substitutions in the merodiploid strain and measuring growth or morphology defects (Table 4, Fig. S8). Class I mutants targeted both the $\alpha 2$ helix and the loop region (residues 107 to 120 and 121 to 130, respectively) of the FtsX_{ECL1} helical lobe (Table 4, Fig. 3A, Fig. S8A). Single-amino-acidsubstitution mutations of FtsX (including E109A [a change of E to A at position 109], L115A, M119A, W123A, F126A, N131A, and N131D) resulted in morphological defects without Zn-Mn (Table 4, Fig. S8C and D) but did not induce a measurable growth

bK_D, equilibrium dissociation constant.

^cNA, not applicable since no heat was observed.



TABLE 4 Amino acid changes made in vivo to disrupt the FtsX_{ECL1}-PcsB interaction^a

Category of change(s) made ^b	Amino acid change(s) or location of insertion	Defect(s) in shape and/or growth	Location(s) in FtsX _{ECL1}
Class I	E109A	Shape only	α2 helix
	E109Q	No	α 2 helix
	Q111A	Noc	α 2 helix
	L115A	Shape only	α 2 helix
	M119A	Shape only	α 2 helix
	W123A	Shape only	Loop ^d
	F126A	Shape only	$Loop^d$
	N131A	Shape only	$Loop^d$
	N131D	Shape only	Loop ^d
Class II	E109A N131A	Shape only	α 2 helix, loop ^d
	Q111A L115C	Shape only	α 2 helix
	L115A M119A	Yes	α 2 helix
	E109A Q111A L115A M119A	Yes	α 2 helix
	F126A W123A N131A	Yes	$Loop^d$
Class IIIe	Residue 51	Yes	Post-TM1
	Residue 78	Shape only	β -Hairpin
	Residue 173	NA^f	Pre-TM2

^aSee the supplemental material for the corresponding strain number and strain construction for each amino acid change or insertion

phenotype (Table 4, Fig. S8B). The expression of FtsX(L115A) resulted in cell shape defects (aspect ratio, length, width, and volume) (Fig. S8C and D), while the expression of FtsX(M119A) only resulted in a change in cell volume (Fig. S8D). These differences were not due to misexpression of FtsX, as Western blot analysis indicates that all mutant proteins were expressed at or near wild-type levels (Fig. 4C). Two other single-amino-acid substitutions (E109Q and Q111A) did not strongly affect growth, morphology, or expression (Table 4, Fig. 4D).

In contrast to the somewhat modest physiological impact of class I substitutions, selected class II mutants (Table 4) exhibited severe morphological and growth defects (Fig. 3, Table 4, Fig. S9). In strains with mutations targeting the α 2 helix [FtsX(E109A Q111A L115A M119A)] or the coil [FtsX(F126A W123A N131A)], ≥98% of cells had severe growth and morphology defects. The growth and morphology of these strains were similar to those of cells in which ftsX was depleted (Fig. 3B to D). Cells expressing these ftsX alleles became significantly rounder and shorter (Fig. 3D) and showed growth inhibition in the absence of zinc (Fig. 3B). Importantly, in the presence of zinc, the cells were indistinguishable from wild-type cells at 3 h (Fig. 3D). At 6 h with zinc, these cells exhibited changes in width and volume, which could be due to overexpression of FtsX at this time point or the expression of wild-type and mutant FtsX simultaneously (Fig. 3D). Western blotting confirmed that FtsX(E109A Q111A L115A M119A) and FtsX(F126A W123A N131A) were expressed in the absence of zinc at 6 h postdepletion (Fig. 4A). The triple FtsX(F126A W123A N131A) mutant expressed in the absence of zinc migrated slightly higher than wild-type FtsX on an SDS-PAGE gel, but it was expressed (Fig. 4A). Taken together, these results reveal that both the α 2 helix and the loop in the helical lobe of FtsX_{FCL1} are important for FtsX function in vivo and confirm the functional importance of the physical interaction of FtsX and PcsB mapped by NMR spectroscopy.

Some of the class II mutants we characterized had just two amino acid changes in the α 2 helix or the extended loop of the helical lobe (Table 4, Fig. S9). We observed that

bClass I, single amino acid changes; class II, multiple amino acid changes; class III, insertion of a (Gly₃Ser)₂ linker.

In the absence of zinc, this strain had no morphology defects. Morphology defects were observed at 3 h when the strain was expressing both the wild-type copy of FtsX and FtsX(Q111A).

^dRefers to the unstructured loop after the α 2 helix in the helical lobe of the FtsX_{ECL1} structure.

eThese strains have an insertion of the amino acid sequence GGGSGGGS after the indicated residue.

^fNA, not applicable, since the strain did not express FtsX with the (Gly₃Ser)₂ insertion. See Fig. 4E for Western blot data.

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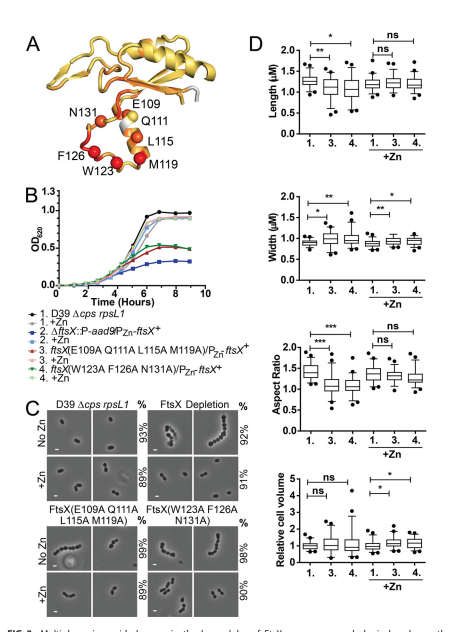


FIG 3 Multiple amino acid changes in the lower lobe of $FtsX_{ECL1}$ cause morphological and growth defects. (A) Amino acid changes made are mapped to the structure of FtsX $_{\rm ECL1}$. The lpha carbon of each residue is shown as a colored sphere on the structure, and the orange-to-red coloring represents the peak height changes in ${}^{1}H^{-15}N$ HSQC spectra upon the addition of 2 molar equivalents of PscB $_{CC}$ to FtsX_{ECL1}. (B) Representative growth curves of strains with amino acid changes in the lower lobe of $\mathsf{FtsX}_{\mathsf{ECL1}}$ compared to the growth of an FtsX depletion strain. These strains were grown without or with 0.45 mM ZnCl₂ with 0.045 mM MnSO₄ (Zn-Mn; indicated as +Zn). Strains shown are as follows: black circle, strain D39 rpsL1 Δcps wild-type parent (1, strain IU1824); gray circle, IU1824 + Zn; blue square, D39 rpsL1 Δcps $\Delta ftsX::P-aad9/bgaA::tet-P_{zn}-ftsX^+$ (2, strain IU12376); light blue square, IU12376 +Zn; red triangle, D39 rpsL1 \(\Delta cps \) ftsX(E109A \(\Q111A \) L115A \(\M119A)/bgaA::tet-P_{zn}-ftsX^+ \) (3, strain \(\IU12861); \(\pink \) triangle, IU12861 +Zn; green inverted triangle, D39 rpsL1 Δcps ftsX(F126A W123A N131A)/bgaA::tet-P_{zn}ftsX+ (4, strain IU12864); light green inverted triangle, IU12864 +Zn. This growth curve experiment was repeated three times with similar results. (C) Representative images of strains at 6 h of growth. The genotype or phenotype of the strain is indicated above each panel. No Zn, Zn-Mn was not added; +Zn, Zn-Mn was added; %, percentage of cells in the population that are morphologically similar to the images shown. More than 50 cells per strain, condition, and experimental repeat were analyzed. These experiments were performed three times independently with similar results. Scale bar shown is equal to 1 μ m. (D) Lengths, widths, aspect ratios, and relative cell volumes of strains at 6 h of growth. Strains are indicated according to numbering in panel B. More than 50 cells were measured per strain and condition over two experimental replicates. For statistical analysis, the Kruskal-Wallis test (one-way ANOVA) with Dunn's multiple-comparison posttest was used to determine if lengths, widths, aspect ratios, and relative cell volumes were significantly different between strains and conditions. ns, not significant; *, P < 0.05; **, *P* < 0.005; ***, *P* < 0.0005.



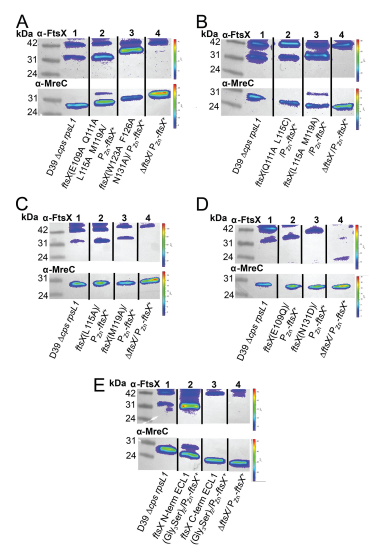


FIG 4 FtsX mutants with amino acid changes are expressed at nearly wild-type levels. Representative blots of anti-FtsX and anti-MreC (Western blotting control for loading) antibodies are shown, with the genotype indicated under each lane. Expected molecular weight (MW) for FtsX is 34.2 kDa, and expected MW for MreC is 29.7 kDa. Samples were grown without Zn and harvested at 6 h of growth. Western blots were imaged as described in supplemental materials and methods in Text S1. (A) FtsX(E109A Q111A L115A M119A) and FtsX(W123A F126A N131A) are expressed at or above wild-type levels without zinc. Lane 1, D39 rpsL1 Δcps (strain IU1824); lane 2, D39 rpsL1 Δcps ftsX(E109A Q111A L115A M119A)/bgaA:: $tet-P_{Zn}-ftsX^+$ (strain IU12861); lane 3, D39 rpsL1 Δcps ftsX(W123A F126A N131A)/bgaA:: $tet-P_{Zn}-ftsX^+$ (strain IU12864); lane 4, D39 rpsL1 Δcps $\Delta ftsX/bgaA$::tet-P_{Zn}- $ftsX^+$ (strain IU13461). (B) FtsX(Q111A L115C) and FtsX(L115A M119A) are expressed at nearly wild-type levels without zinc. Lane 1, D39 rpsL1 \(\Delta cps\) (IU1824); lane 2, D39 rpsL1 Δcps ftsX(Q111A L115C)/bgaA:: $tet-P_{Zn}$ - $ftsX^+$ (IU13064); lane 3, D39 rpsL1 Δcps ftsX(L115A)M119A)/bgaA:: $tet-P_{Zn}$ - $ftsX^+$ (IU13066); lane 4, D39 rpsL1 Δcps $\Delta ftsX/bgaA$:: $tet-P_{Zn}$ - $ftsX^+$. (C) FtsX(L115A)and FtsX(M119A) are expressed at nearly wild-type levels without zinc. Lane 1, D39 rpsL1 \(\Delta cps\) (IU1824); lane 2, D39 rpsL1 Δcps ftsX(L115A)/bgaA::tet-P_{Zn}-ftsX⁺ (IU12521); lane 3, D39 rpsL1 Δcps ftsX(M119A)/ bgaA::tet-P_{Zn}-ftsX+ (IU12637); lane 4, D39 rpsL1 Δcps ΔftsX/bgaA::tet-P_{Zn}-ftsX+ (IU13461). (D) FtsX(E109Q) and FtsX(N131D) are expressed at nearly wild-type levels without zinc. Lane 1, D39 rpsL1 \(\Delta cps\) (IU1824); lane 2, D39 rpsL1 Δcps $ftsX(E109Q)/bgaA::tet-P_{Zn}-ftsX^+$ (IU13088); lane 3, D39 rpsL1 Δcps ftsX(N131D)/res $bgaA::tet-P_{Zn}-ftsX^+$ (IU13089); lane 4, D39 rpsL1 Δcps $\Delta ftsX/bgaA::tet-P_{Zn}-ftsX^+$ (IU13461). (E) FtsX with (Gly₃Ser)₂ after residue 51 is expressed, whereas FtsX with (Gly₃Ser)₂ after residue 173 is not expressed. These are referred to as ftsX N-term ECL1(Gly₃Ser)₂ and ftsX C-term ECL1(Gly₃Ser)₂, respectively. Lane 1, D39 rpsL1 Δcps (IU1824); lane 2, D39 rpsL1 Δcps ftsX N-term ECL1(Gly₃Ser)₂/ $bgaA::tet-P_{Zn}-ftsX^+$ (IU12629); lane 3, D39 rpsL1 Δcps ftsX C-term ECL1(Gly₃Ser)₂/ $bgaA::tet-P_{Zn}-ftsX^+$ (IU12869); lane 4, D39 rpsL1 Δcps ΔftsX/bgaA::tet-P_{7n}-ftsX⁺ (IU13461). These experiments were performed two to three times independently.

99% of cells expressing FtsX(L115A M119A) exhibited strong growth and morphology defects at 6 h in the absence of zinc (Table 4, Fig. S9B and C), and these cells displayed decreases in length, width, and volume relative to those of wild-type cells (Fig. S9D). This mutant was expressed at nearly wild-type levels in the absence of zinc (Fig. 4B). These data confirm that the tandem L115A M119A substitution disrupts FtsX function, even though these substitions result in only slight morphological defects as individual single mutations (Table 4, Fig. S8B to D, Fig. S9B to D). Another double mutant, FtsX(Q111A L115C), induced the formation of long chains and a "boxy" cell morphology (Table 4, Fig. S9C). This mutant resulted in shorter cells with a significantly different aspect ratio than the wild type (Fig. S9D), but this strain had no growth phenotype (Fig. S9B). In contrast, another double substitution class II mutant, FtsX(E109A N131A) (Table 4), exhibited no strong morphology or growth defects.

Finally, class III insertion mutants (Table 4) were constructed and used to evaluate whether other regions of $FtsX_{ECL1}$ were important for the $FtsX_{ECL1}$ -PcsB interaction or for FtsX function. We inserted a $(Gly_3Ser)_2$ flexible linker either approximately where $FtsX_{ECL1}$ is predicted to enter (residue 51) or exit (residue 173) the membrane bilayer or in the β -hairpin, which exhibits significant conformational disorder over a range of timescales (Fig. 1A). An insertion after residue 51 in $FtsX_{ECL1}$ was detrimental to both growth and morphology (Table 4), and this insertion did not disrupt FtsX expression (Fig. 4E). The insertion after residue 173 in $FtsX_{ECL1}$ also caused growth and morphology defects, but this FtsX variant was not expressed in cells (Table 4, Fig. 4E). The β -hairpin ($Gly_3Ser)_2$ insertion (Fig. 1B, see appendix in Text S1) was introduced after amino acid 78 of $FtsX_{ECL1}$, which corresponds to the tip of the β -turn in the β -hairpin (Fig. 1B). This strain also exhibited no growth defect, but these cells were significantly smaller, although only at the 3 hr time point (see appendix in Text S1). We conclude that the β -hairpin does not play a major role in FtsX-PcsB interaction, consistent with the NMR mapping experiments.

FtsX_{ECL1}(L115A M119A) is stably folded and unable to bind PscB_{CC}. We reasoned that if the defects observed in class I and class II mutants were due to the disruption of the FtsX_{ECL1}-PscB_{CC} interaction, this should affect the affinity of this interaction as measured by ITC. We first characterized the L115A M119A double mutant by 1 H- 15 N NMR and CD spectroscopy to confirm its structural integrity. The CD spectrum resembled that of the wild type, as did the 1 H- 15 N HSQC spectrum, with clear chemical shift perturbations only among those resonances in the immediate vicinity of the double substitution (Fig. S10B). Both pieces of data suggest a local rather than global perturbation of the α 2-loop lobe in the FtsX_{ECL1} structure upon introduction of the L115A M119A double substitution. As anticipated, titration of PscB_{CC}(47–267) with FtsX_{ECL1}(L115A M119A) reveals no detectable binding (no observable heat) (Fig. 2B, Table 3) compared to the results using wild-type FtsX_{ECL1}. These data confirm that the helical lobe of FtsX_{ECL1} interacts with PscB_{CC} and that this interaction is required for viability and proper cell shape.

In contrast to the L115A M119A double mutant, the severely functionally compromised representative triple mutant $FtsX_{ECL1}$ (W123A F126A N131A) and quadruple mutant $FtsX_{ECL1}$ (E109A Q111A L115A M119A) exhibited more pronounced structural perturbations that nonetheless map only to the helical lobe. The triple mutant was indistinguishable from the L115A M119A and wild-type $FtsX_{ECL1}$ derivatives by CD spectroscopy, while the quadruple mutant exhibited less molar ellipticity, or secondary structure (Fig. S10D). Inspection of their 1H - ^{15}N HSQC spectra reveals that although the core and β -hairpin domains essentially resemble those of the wild type, each of these mutants exhibits considerable perturbation of resonances throughout the helical lobe (Fig. S10A and C). Since these mutants are functionally compromised, these structural findings strongly support the conclusion that the structural integrity of the lower lobe is essential for the physical interaction with PcsB and the function of FtsX in pneumococcal cells.



DISCUSSION

This study presents a comprehensive analysis of the solution and X-ray structures of the outward-facing large extracellular loop of FtsX (FtsX_{FCI 1}) from *S. pneumoniae* and defines a physical interaction site with the coiled-coil domain of peptidoglycan hydrolase PcsB (PscB_{CC}). Our FtsX_{ECL1} structures reveal a globular fold that, while similar to the large extracellular loop of FtsX from M. tuberculosis (16), is characterized by unique features. The upper β -hairpin distinguishes S. pneumoniae FtsX_{ECL1} from that of M. tuberculosis, and despite being characterized by significant conformational dynamics on a range of timescales, it is not required for the interaction of S. pneumoniae FtsX_{FCI.1} with PcsB. The function of this domain is not well defined by our data, but it could play a role in another process, e.g., FtsX dimerization, interaction with the small extracellular loop (ECL2), or interaction with another domain of PcsB. Alternatively, it could be a result of the difference in cell wall architecture between S. pneumoniae and M. tuberculosis. On the other hand, the helical lobe of FtsX_{ECL1}, common to the FtsX structures of both S. pneumoniae and M. tuberculosis, is vital for the interaction with PcsB in vitro, and this interface is functionally important in vivo. Increasing numbers of Ala substitutions tested here increasingly disrupt this interaction and ultimately cause dramatic growth and morphology defects, indicating that the helical lobe of FtsX_{FCL1} is essential for regulation of PcsB during cell division. Interestingly, this region of FtsX_{FCI 1} corresponds with the region shown to be important for the interaction of M. tuberculosis FtsX_{ECD} with its PG hydrolase, RipC (16). This suggests that the helical lobe could be a conserved functional determinant for the interaction of FtsX with cognate hydrolases or adaptor proteins across many species of bacteria.

We propose that the helical lobe of FtsX_{ECL1} is important for the activation of cognate hydrolase activity either directly or indirectly through adaptors (Fig. 5). The exact role of the second extracellular loop of FtsX (FtsX_{ECL2}) is unknown, but it may also regulate this process, as temperature-sensitive mutations in *pcsB* were found to be suppressed by mutations in the coding region for FtsX_{ECL2} (20). Previous work suggests that FtsEX forms a dimer (32), as dimerization of the FtsE ATPase domain is likely a necessary condition for ATP hydrolysis (Fig. 5A) (36, 37). After formation of the complex, ATP hydrolysis by FtsE results in a conformational change in FtsX, releasing PcsB from what we anticipate is an inhibited state (Fig. 5B and C) (22). This interaction is mediated by the helical lobe of FtsX_{ECL1}, although the membrane and, possibly, lipid binding by FtsX_{ECL1} and FtsX_{ECL2} itself may also play a role. We propose that the interaction of FtsX_{ECL1} with the PcsB coiled-coil domain communicates release of the PcsB CHAP domain from an inhibited state and, thus, is important for modulating PG hydrolysis by PcsB (Fig. 5B and C).

Recently, the structure of *Aggregatibacter actinomycetemcomitans* MacB was reported and suggested to be a structural paradigm for the ABC3 transporter superfamily that includes FtsX (38). MacB was proposed to function as a mechanical pump to drive enterotoxin transport through TolC in *E. coli* (38). Crow et. al found that MacB itself did not transport enterotoxin but drove TolC to transport it instead, due to the lack of a central cavity in the MacB structure (38). As such, they proposed MacB as a model for other so-called "mechanotransmitters" belonging to this same ABC3 transporter superfamily. While this proposed function of mechanotransmission may well characterize MacB and FtsX, the overall structure of FtsX_{ECL1} from *S. pneumoniae* is clearly distinct from the periplasmic domain of MacB, thus revealing that MacB does not readily provide a structural basis for understanding FtsX-dependent peptidoglycan hydrolases. Future work using reconstituted FtsEX and PcsB complexes will allow for understanding how this common mechanotransmission principle extends throughout the ABC3 superfamily.

MATERIALS AND METHODS

NMR spectroscopy. Spectra of ¹⁵N- or ¹⁵N¹³C-labeled FtsX_{ECL1} were recorded at 298 K on Varian (Agilent) DDR 600- or 800-MHz spectrometers equipped with cryogenic probes in the METACyt Biomolecular NMR Laboratory at Indiana University Bloomington. NMR samples contained 50 mM potassium phosphate, pH 7.0, 50 mM NaCl, and 10% (vol/vol) D₂O, with 0.2 mM 4,4-dimethyl-4-silapentane-1-

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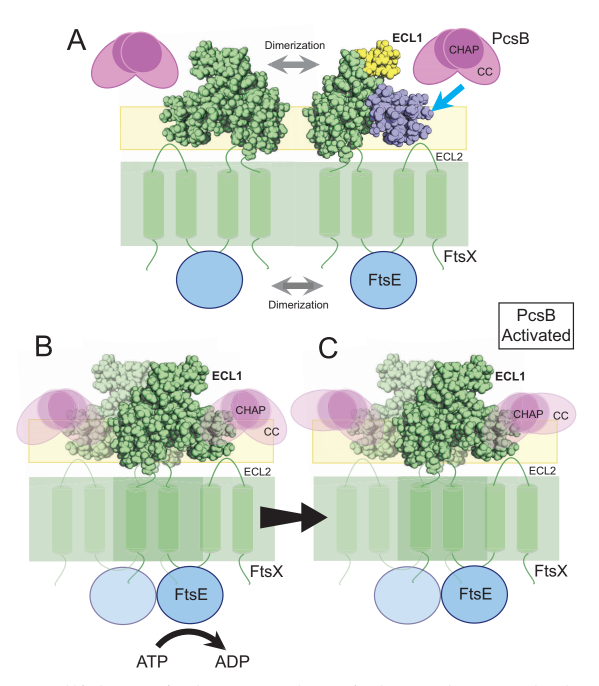


FIG 5 Model for the activation of PcsB by FtsX_{ECL1}. (A) FtsEX dimerizes to form the active complex. PcsB is secreted into the extracellular milieu. Attraction of PcsB to the area of active FtsX complexes might be mediated by its propensity to interact with membranes (32). The ECL1 and ECL2 loops are indicated on FtsX. FtsX_{ECL1} is shown in green, with the β-hairpin and α-helical lobe shaded in yellow and blue, respectively. (B) After formation of the active complex, ATP hydrolysis by FtsE causes a conformational change in FtsX. (C) PcsB interacts with FtsX_{ECL1} via its coiled-coil domain, and this interaction causes activation of the peptidoglycan hydrolytic activity of PcsB. PcsB, along with other factors in the cell, allows cell division to proceed normally. Functional FtsX, FtsE, and PcsB are all required for efficient cell division.

sulfonic acid (DSS) for chemical shift referencing. The typical concentrations of FtsX_{ECL1} were 50 μ M for ¹⁵N HSQC spectra and 400 μ M for triple-resonance and dynamics experiments. ¹J_{HN} splittings for residual dipolar couplings (RDCs) were measured using two-dimensional in-phase/anti-phase (2D IPAP) [¹⁵N, ¹H]-HSQC spectra (39), recorded on an isotropic sample and on a sample aligned with 20 mg/ml phage *Pf*1 (ASLA Biotech). ¹D_{HN} was calculated from ¹D_{HN} = ¹J_{HN} (anisotropic) – ¹J_{HN} (isotropic). Aromatic sidechains were assigned using the HBCBCGCDHD and HBCBCGCDCEHE experiments (40). For experiments detecting PscB_{CC} binding, ¹⁵N FtsX_{ECL1} was kept at 50 μ M, and ¹H-¹⁵N HSQC spectra were recorded with the following concentrations of PscB_{CC}(47–267): 0, 50 μ M, 100 μ M, 200 μ M, and 400 μ M. nmrPipe, Sparky, CARA (http://cara.nmr.ch), CCPNMR, and NMRbox (41–44) were used for data processing and



analysis. Resonance assignments and dynamics data are available in the BMRB under accession code 30523. These NMR data were used to calculate and refine the solution structure of ECL1 (see supplemental materials and methods in Text S1), with the ensemble of the 20 lowest-energy structures (see Table 1 for structure statistics) deposited in the Protein Data Bank (PDB code 6MK7).

X-ray crystallography. Crystallization screenings were performed by using high-throughput crystallization techniques in a NanoDrop robot with Innovadyne SD-2 microplates (Innovadyne Technologies, Inc.) and screening using PACT Suite and JCSG Suite (Qiagen), JBScreen Classic 1 to 4 and 6 (Jena Bioscience), and Crystal Screen, Crystal Screen 2, and Index HT (Hampton Research). Positive conditions in which crystals grew were optimized by the sitting-drop vapor-diffusion method at 291 K by mixing 1 μ l of protein solution and 1 μ l of precipitant solution, equilibrated against 150 μ l of precipitant solution in the reservoir chamber. The best crystals were obtained under a crystallization condition of a solution containing 0.1 M sodium citrate, pH 5.6, 0.2 M potassium-sodium tartrate, and 2 M ammonium sulfate. These crystals were further optimized by cocrystallization in the presence of detergents dodecyltrimethylammonium chloride (detergent 1 [D1]; 46 μM), n-undecyl-β-p-maltoside (detergent 2 [D2]; 0.59 mM), and n-decyl- β -D-maltoside (detergent 3 [D3]; 1.8 mM). Crystals were cryoprotected in the precipitant solution supplemented with 25% (vol/vol) glycerol prior to flash cooling at 100 K. Diffraction data were collected in the XALOC beamline at the ALBA synchrotron (CELLS-ALBA, Spain), using a Pilatus 6 M detector and a wavelength of 0.979257 Å. Crystals diffracted up to 2.0- to 2.3-Å resolution and belonged to the P 4_3 2_1 2 space group. The collected data sets were processed with XDS (45) and Aimless (46). Two FtsX_{FCI 1} molecules were found in the asymmetric unit for all three structures, yielding $Matthews \ coefficients \ of \ 2.59 \ \mathring{A}^3/Da \ (FtsX_{ECL}-D1), \ 3.49 \ \mathring{A}^3/Da \ (FtsX_{ECL}-D2), \ and \ 2.49 \ \mathring{A}^3/Da \ (FtsX_{ECL}-D1) \ (47)$ and solvent contents of 52.5% (Fts X_{ECL} -D1), 64.7% (Fts X_{ECL} -D2), and 50.7% (Fts X_{ECL} -D3). Structure determination was performed by molecular replacement using the online server Morda (http://www .biomexsolutions.co.uk/morda/). Refinement and manual model building were performed with Phenix (48) and Coot (49), respectively. The data for $FtsX_{ECL}$ -D1 and $FtsX_{ECL}$ -D3 presented translational noncrystallographic symmetry (fractional coordinates of [-0.498, -0.498, 0.50] and height relative to origin of 79.60% for FtsX_{ECL}-D1, and fractional coordinates of [-0.498, -0.498, 0.50] and height relative to origin of 78.70% for $FtsX_{ECL}$ -D3) that were treated with Phenix (48). The stereochemistry of the final model was checked by MolProbity (50). Data collection and processing statistics are shown in Table 2.

Bacterial strains, plasmids, and growth conditions. Bacterial strains and plasmids used in this study are listed in the supplemental material. *S. pneumoniae* strains were derived from IU1945, an unencapsulated derivative of serotype 2 *S. pneumoniae* strain D39 (51). Strains were grown on Trypticase soy agar II with 5% (vol/vol) defibrinated sheep blood (TSAII-BA) plates or in Becton, Dickinson brain heart infusion (BHI) broth at 37°C in an atmosphere of 5% CO₂. *E. coli* strains for protein expression were derived from strain BL21(DE3) (catalog number C2527H; NEB). *E. coli* strains were grown in Luria-Bertani (LB) broth or in M9 minimal medium supplemented with 15 NH₄Cl at 37°C with shaking at 150 rpm. When required, tetracycline (0.25 to 2.5 μg/ml), kanamycin (250 μg/ml), spectinomycin (150 μg/ml), streptomycin (250 μg/ml), ampicillin (100 μg/ml), and/or isopropyl β-D-1-thiogalactopyranoside (IPTG) (1 mM) was added to *S. pneumoniae* or *E. coli* culture medium. *S. pneumoniae* strains requiring zinc for expression of essential genes were grown with 0.45 mM ZnCl₂ and 0.045 MnSO₄.

Growth curve experiments and phase-contrast microscopy of strains. For physiological and morphological analyses of strains, cells were inoculated from frozen glycerol stocks into BHI broth, serially diluted, and incubated for 10 to 12 h statically at 37°C in 5% CO₂ overnight. If zinc was required for growth of cultures, 0.45 mM ZnCl₂ and 0.045 MnSO₄ were added to overnight tubes. The next day, cultures ranging from an optical density at 620 nm (OD₆₂₀) of ≈0.05 to 0.4 were diluted into fresh BHI to an OD_{620} of ≈ 0.003 in 4-ml volumes, and two identical cultures for each strain were prepared, one with 0.45 mM ZnCl₂-0.045 MnSO₄ and one without. These cultures were grown under the same growth conditions as described above. Growth was monitored turbidimetrically every 45 min to 1 h with a Genesys 2 spectrophotometer (Thermo Scientific). For microscopic analyses, samples (1 to 2 μ l) were taken at 3 h and 6 h and examined using a Nikon E-400 epifluorescence phase-contrast microscope with a 100× Nikon Plan Apo oil-immersion objective (numerical aperture, 1.40) connected to a CoolSNAP HQ2 charge-coupled device (CCD) camera (Photometrics). Images were processed using NIS-Elements AR software (Nikon), and measurements and calculation of cell width, length, volume, and aspect ratio were performed as described previously (52, 53). Statistical significance was determined using GraphPad Prism (GraphPad Software, Inc.) by comparing values for cell width, length, volume, and aspect ratio measured for at least 50 cells over two experimental replicates. To determine if values were significantly different between strains and conditions, the Kruskal-Wallis test (one-way analysis of variance [ANOVA]) with Dunn's multiple comparison posttest was used.

For additional materials and methods, please see Text S1.

Data availability. The ensemble of the 20 lowest-energy structures from using NMR data to calculate and refine the solution structure of ECL1 has been deposited in the Protein Data Bank (PDB code 6MK7). The atomic coordinates of FtsX_{ECL1} determined by cocrystallization in the presence of detergents 1, 2, and 3 have been deposited in the Protein Data Bank with PDB codes 6HE6, 6HEE, and 6HFX, respectively. NMR spectroscopy resonance assignments and dynamics data are available in the BMRB under accession code 30523.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/mBio .02622-18.

TEXT \$1, PDF file, 2.6 MB.

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FIG S1, EPS file, 13.2 MB.

FIG S2, EPS file, 2.9 MB.

FIG S3, EPS file, 8.8 MB.

FIG S4, EPS file, 4.3 MB.

FIG S5, EPS file, 8.5 MB.

FIG S6, EPS file, 8.1 MB.

FIG S7, EPS file, 6.2 MB.

FIG S8, EPS file, 5.8 MB.

FIG S9, EPS file, 5.9 MB.

FIG S10, EPS file, 19.2 MB.

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