CXCL10 Acts as a Bifunctional Antimicrobial Molecule against *Bacillus anthracis*

Katie R. Margulieux, a Jay W. Fox, b Robert K. Nakamoto, c Molly A. Hughes a

Division of Infectious Diseases, Department of Medicine, University of Virginia, Charlottesville, Virginia, USA; Department of Microbiology, Immunology, and Cancer Biology, University of Virginia, Charlottesville, Virginia, USA; Department of Molecular Physiology and Biological Physics, University of Virginia, Charlottesville, Virginia, USA

ABSTRACT *Bacillus anthracis* is killed by the interferon-inducible, ELR(−) CXC chemokine CXCL10. Previous studies showed that disruption of the gene encoding FtsX, a conserved membrane component of the ATP-binding cassette transporter-like complex FtsE/X, resulted in resistance to CXCL10. FtsX exhibits some sequence similarity to the mammalian CXCL10 receptor, CXCR3, suggesting that the CXCL10 N-terminal region that interacts with CXCR3 may also interact with FtsX. A C-terminal truncated CXCL10 was tested to determine if the FtsX-dependent antimicrobial activity is associated with the CXCR3-interacting N terminus. The truncated CXCL10 exhibited antimicrobial activity against the *B. anthracis* parent strain but not the ΔftsX mutant, which supports a key role for the CXCL10 N terminus. Mutations in FtsE, the conserved ATP-binding protein of the FtsE/X complex, resulted in resistance to both CXCL10 and truncated CXCL10, indicating that both FtsX and FtsE are important. Higher concentrations of CXCL10 overcame the resistance of the ΔftsX mutant to CXCL10, suggesting an FtsX-independent killing mechanism, likely involving its C-terminal α-helix, which resembles a cationic antimicrobial peptide. Membrane depolarization studies revealed that CXCL10 disrupted membranes of the *B. anthracis* parent strain and the ΔftsX mutant, but only the parent strain underwent depolarization with truncated CXCL10. These findings suggest that CXCL10 is a bifunctional molecule that kills *B. anthracis* by two mechanisms. FtsE/X-dependent killing is mediated through an N-terminal portion of CXCL10 and is not reliant upon the C-terminal α-helix. The FtsE/X-independent mechanism involves membrane depolarization by CXCL10, likely because of its α-helix. These findings present a new paradigm for understanding mechanisms by which CXCL10 and related chemokines kill bacteria.

IMPORTANCE Chemokines are a class of molecules known for their chemoattractant properties but more recently have been shown to possess antimicrobial activity against a wide range of Gram-positive and Gram-negative bacterial pathogens. The mechanism(s) by which these chemokines kill bacteria is not well understood, but it is generally thought to be due to the conserved amphipathic C-terminal α-helix that resembles cationic antimicrobial peptides in charge and secondary structure. Our present study indicates that the interferon-inducible, ELR(−) chemokine CXCL10 kills the Gram-positive pathogen *Bacillus anthracis* through multiple molecular mechanisms. One mechanism is mediated by interaction of CXCL10 with the bacterial FtsE/X complex and does not require the presence of the CXCL10 C-terminal α-helix. The second mechanism is FtsE/X receptor independent and kills through membrane disruption due to the C-terminal α-helix. This study represents a new paradigm for understanding how chemokines exert an antimicrobial effect that may prove applicable to other bacterial species.
against a wide variety of Gram-negative and Gram-positive bacteria such as *Escherichia coli*, *Staphylococcus aureus*, *Listeria monocytogenes*, and *Bacillus anthracis* (14−17). Additionally, CXCL10 exerts an antimicrobial effect against both laboratory strains and clinical isolates of *E. coli*, including a multidrug-resistant strain obtained from a clinical sample (21). Like the structure of other chemokines, that of CXCL10 consists of a relatively unstructured N terminus, three antiparallel β-sheets, and a C-terminal amphipathic α-helix (22−24). The interferon-inducible, ELR(−) CXC chemokines interact with CXCR3, a receptor that is found on host immune cells such as T-cells, B-cells and natural killer cells (25, 26). These chemokines bind to the receptor via a series of interactions that are dependent upon the interaction of the chemokine N terminus with the extracellular domain of CXCR3 (amino acids [aa] 1 to 35) (8, 27). Further interaction of the chemokine N terminus and β-sheets with other regions of CXCR3 then occurs in order to activate the signaling cascade necessary to initiate chemotaxis of host immune cells (8, 26, 27).

*B. anthracis* is a Gram-positive, spore-forming bacterium that causes the disease anthrax following the exposure of susceptible mammals to spores and their subsequent germination into vegetative bacilli (28−31). Our laboratory has previously shown that CXCL10 exhibits *in vitro* antimicrobial activity against both *B. anthracis* spores and vegetative cells (17, 32). *In vivo* neutralization of CXCL10 resulted in a significantly higher mortality rate in a mouse lung infection model with C57BL/6 mice, which are inherently resistant to intranasal infection by the *B. anthracis* Sterne strain (32). Blocking of CXCR3 resulted in no difference in survival from control (vehicle-treated) mice (32). These *in vivo* data support a direct antimicrobial role for CXCL10 that is unrelated to its interaction with CXCR3 and suggest that CXCL10 is produced locally at a high enough concentration to actively kill the bacteria at sites of infection. Given the charge and structural similarity of the CXCL10 C-terminal α-helix to cationic antimicrobial peptides, it was initially hypothesized that the α-helix was responsible for its antimicrobial activity, likely via membrane disruption based on characterized mechanisms utilized by cationic antimicrobial peptides like LL37 (19, 33, 34).

Our laboratory previously screened a *B. anthracis* Sterne strain transposon mutant library to identify bacterial gene products associated with the antimicrobial activity of recombinant human CXCL10 (17, 32, 35). Of particular interest was the finding that disruption of the *ftsX* gene, which encodes FtsX, resulted in the resistance of *B. anthracis* vegetative cells to CXCL10 (35). FtsX is a conserved membrane component of the FtsE/X complex that appears to be a prokaryotic ATP-binding cassette transporter (35−38). The FtsE/X complex in other bacterial species, such as *Bacillus subtilis*, *E. coli*, and *Mycobacterium tuberculosis*, has been shown to play a key role in cell wall peptidoglycan (PG) processing through the activation of PG hydrolytic enzymes by FtsX (37−41). FtsE possesses conserved Walker motifs that are responsible for ATP binding and hydrolysis (40, 42, 43). Work conducted by others using *B. subtilis* and *M. tuberculosis* suggests that ATP hydrolysis by FtsE induces a conformational change in FtsX resulting in the activation of PG hydrolases for cell wall processing, a critical function for maintenance of bacterial cell integrity (39−41). Major classes of antimicrobials (e.g., beta-lactam antibiotics) kill bacteria by inhibiting specific steps in PG synthesis, which also affects bacterial cell integrity (44).

In *silico* analysis of the *B. anthracis* FtsX protein revealed that a 27-aa region (aa 54 to 80) in the large external loop has 45% similarity to the N-terminal amino acid sequence of CXCR3 (aa 9 to 35), the mammalian receptor for CXCL10 (35). Importantly, this N-terminal amino acid sequence is involved in the binding of the N-terminal region of CXCL10 (8, 45, 46). The similarity between CXCR3 and FtsX suggested that the N-terminal region of CXCL10 may also interact with the CXCR3-similar region of FtsX. Thus, the antimicrobial activity of CXCL10 may be primarily the result of a receptor-mediated effect that leads to cell lysis and not necessarily to depolarization resulting from the insertion of the C-terminal α-helix into the cytoplasmic membrane. In the present study, we investigated two potential ways in which CXCL10 may kill bacteria. Our data suggest that the antimicrobial activity of CXCL10 is mediated by its interaction with FtsE/X, which utilizes a mechanism separate from that of the C-terminal α-helix. In the absence of functional FtsE/X, higher concentrations of CXCL10 exhibit a bactericidal effect that appears to be dependent on the presence of its C-terminal α-helix and general membrane depolarization. However, the result of each pathway (i.e., cell lysis) is the same, with killing of *B. anthracis* being the most effective when both mechanisms are present and active.

**RESULTS**

The CXCR3-similar region of FtsX is involved in CXCL10 interaction with *B. anthracis*. FtsE/X is widely conserved among Gram-positive and Gram-negative bacterial species, with a similarity ranging between 46 and 75%, as determined by BLASTP analysis and amino acid sequence alignment (see Text S1 in the supplemental material) (47, 48). In general, bacterial FtsX protein structures consist of four transmembrane domains and two extracellular loops, although functional differences between the FtsX proteins of Gram-positive and Gram-negative bacterial species have been observed (35, 42). In Gram-positive bacteria, FtsX, along with FtsE, exists in an FtsE/X complex that is involved in PG processing during cellular elongation and spore formation. In Gram-negative bacteria, FtsX and the FtsE/X complex have been shown to direct PG processing at the bacterial septum during cell division (37−41). Our initial investigation into *B. anthracis* FtsX identified two regions of similarity between FtsX and the known mammalian CXCL10 receptor, CXCR3 (50) (see Text S1). While those specific regions are not conserved between Gram-positive and Gram-negative species, there are other regions of similarity between CXCR3 and FtsX, notably located in the exposed outer loops. In this study, the focus of investigation involved the region of *B. anthracis* FtsX (aa 54 to 80) with sequence similarity to the region of human CXCR3 (aa 9 to 35) that is primarily responsible for chemokine binding, since that similarity between FtsX and CXCR3 suggested a region for possible interaction with CXCL10.

A peptide competition assay was performed to determine if the CXCR3-similar region of FtsX is involved in facilitating the antimicrobial activity of CXCL10. A 27-aa peptide representing the CXCR3-like region of FtsX was synthesized, as well as a scrambled peptide to control for sequence specificity (Fig. 1A). The antimicrobial activity of CXCL10 alone or the peptide-CXCL10 mixture against the *B. anthracis* Sterne strain, hereby designated as parent strain, was examined after 2 h of incubation (35). There was a statistically significant increase in the survival of bacteria exposed to the FtsX peptide-CXCL10 mixture at molar ratios of both 10:1 and 20:1 (Fig. 1B). No significant increase in survival was observed.
in the scrambled peptide-CXCL10 competition assay (Fig. 1B). These results suggested that the CXCR3-similar region of FtsX was important in mediating the antimicrobial effect of CXCL10 and that the interaction required a specific sequence and was not just due to peptide charge interactions.

C-terminally truncated CXCL10 (CTTC) synthesis and physical characteristics. The importance of the CXCR3-similar region of FtsX indicated a possible site for the interaction of CXCL10 with \textit{B. anthracis}. It was previously determined through the work of others that the relatively unstructured N-terminal region of CXCL10 initially interacts with CXCR3, resulting in immune cell activation (8, 27). We hypothesized that the N-terminal region of CXCL10 may also interact with bacterial FtsX, resulting in an antimicrobial effect. The amphipathic \( \alpha \)-helix of various chemokines has been considered to be the portion of the proteins responsible for antimicrobial activity because of its positive charge and secondary structural resemblance to cationic antimicrobial peptides and defensins (19, 33, 34). However, since our findings unexpectedly implicated the N-terminal portion of CXCL10 in eliciting an antimicrobial effect, we conducted the following series of experiments.

To determine if it is the N-terminal region or the C-terminal \( \alpha \)-helix of CXCL10 that is responsible for antimicrobial activity against \textit{B. anthracis}, a truncated CXCL10 protein was synthesized that lacked the C-terminal 23 aa (Fig. 2A). The sequence of CTTC was confirmed by mass spectrometry (data not shown). Circular dichroism (CD) spectra confirmed that CTTC lacked the \( \alpha \)-helical content, as evidenced by the absence of the spectral peaks at 190 and 210 nm characteristic of intact CXCL10 (Fig. 2B and C). Importantly, analysis of the CTTC spectrum revealed that the content of \( \beta \)-strands (39%) was similar to that of the properly folded chemokine on the basis of the published crystal structure (Protein Data Bank code 1080), thus confirming that the secondary structures of CTTC were intact (23, 45, 51).

CTTC retains antimicrobial activity against the \textit{B. anthracis} parent strain. Various concentrations of CXCL10 and CTTC were compared for antimicrobial activity against the \textit{B. anthracis} parent strain. Micromolar concentrations were used instead of the previously published micrograms per milliliter because of the mass difference between CXCL10 and CTTC (35). Concentration-dependent killing of the \textit{B. anthracis} parent strain was observed, with no viable bacteria detected at concentrations \( \geq 1.2 \) \( \mu \text{M} \)

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure1}
\caption{The CXCR3-similar region of FtsX is important in mediating CXCL10 antimicrobial activity against the \textit{B. anthracis} parent strain. (A) A synthetic peptide consisting of the 27 aa (aa 54 to 80) of FtsX with similarity to the CXCR3-binding region (aa 9 to 35) of CXCL10 (35) and a scrambled control. (B) A peptide competition assay with the FtsX peptide in a 10:1 or 20:1 molar ratio with 0.46 \( \mu \text{M} \) CXCL10 resulted in statistically significant less killing of the \textit{B. anthracis} parent strain than by CXCL10 alone. Incubation with the control peptide had no effect on killing. Bacterial viability was measured by alamarBlue reduction, and fluorescence is expressed as a percentage of that of the strain-specific untreated control. Data points represent mean values \( \pm \) the standard errors of the means; \( n = 3 \) separate experiments using triplicate wells. **, \( P \leq 0.001 \); ***, \( P \leq 0.0001 \).}
\end{figure}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure2}
\caption{Complete amino acid sequence and secondary-structure comparison of CXCL10 and CTTC. (A) Amino acid sequences of intact CXCL10 and synthetic protein CTTC (aa 1 to 54) lacking the C-terminal \( \alpha \)-helix. The amino acids removed from intact CXCL10 are in bold. (B, C) CD showed a distinct \( \alpha \)-helical pattern in the intact CXCL10 spectrum, as indicated by peaks at wavelengths of 190 and 210 nm; these peaks were absent from the CTTC spectrum. The calculated \( \beta \)-sheet contents of both the CXCL10 and CTTC spectra were consistent with the estimated percentage of \( \beta \)-sheets in the published CXCL10 crystal structure (45).}
\end{figure}
CXCL10. CTTC also exhibited concentration-dependent killing but was less potent than intact CXCL10 (Fig. 3A). A statistically significant difference in antimicrobial activity between CXCL10 and CTTC was observed at all of concentrations tested. Thus, in the absence of the amphipathic α-helix, CTTC retained some antimicrobial activity, suggesting that the N terminus and other portions of CXCL10 could kill *B. anthracis* by a mechanism that was independent of the presence of the C-terminal α-helix.

The *B. anthracis* Δ*ftsX* mutant is not susceptible to CTTC. Based on previous studies in our laboratory (35), *B. anthracis* Δ*ftsX* was relatively resistant to CXCL10 when compared with the parent strain. However, this resistance could be overcome by higher concentrations of CXCL10 (Fig. 3B). In a previously published study, we showed that complementation of the *B. anthracis* Δ*ftsX* mutant with plasmid pUVA113, encoding an isopropyl-β-D-thiogalactopyranoside (IPTG)-inducible wild-type *ftsX* gene, restored susceptibility to CXCL10 compared to that of the Δ*ftsX* mutant complemented with the empty control vector pUTE973 (35). Therefore, to investigate whether the CXCL10 C-terminal α-helix plays a role in this killing, the antimicrobial activity of CTTC against the *B. anthracis* Δ*ftsX* mutant was assessed. In the absence of both FtsX and the CXCL10 α-helix, no antimicrobial activity was observed (Fig. 3B). A statistically significant difference between the activity of CXCL10 and that of CTTC against the Δ*ftsX* mutant was observed at all concentrations above 0.70 μM. The Δ*ftsX*-complemented strain *B. anthracis* Δ*ftsX*/pUVA113 was susceptible to CTTC, whereas the Δ*ftsX* mutant complemented with empty vector plasmid pUTE973 remained resistant (see Fig. S1 in the supplemental material). These results suggested at least two possible mechanisms by which CXCL10 kills *B. anthracis*. One mechanism is mediated through the interaction between the N-terminal region of CXCL10 and bacterial FtsX and does not require the C-terminal α-helix (Fig. 3A); and the other, observed at higher concentrations of intact CXCL10 and in the absence of bacterial FtsX, requires the presence of the C-terminal α-helix (Fig. 3B).

**B. anthracis** FtsE Walker motif mutant strain. To test whether an active FtsE/X complex is necessary for CXCL10-mediated antimicrobial activity, an FtsE mutant with point mutations in the Walker A and B motifs was generated (52). FtsE/X is encoded within a single operon, and point mutations within the ftsE gene Walker motifs disrupt the function of FtsE without interfering with the expression of FtsX (40). Walker motifs are highly conserved between bacterial species and are involved in ATP binding and hydrolysis. Point mutations within Walker motifs are well documented to result in loss of the protein energy providing function (40, 42, 43, 49, 53, 54). Point mutations in ftsE resulted in two amino acid substitutions, K123A and D481N. The resulting mutant strain was designated *B. anthracis* ftsE(K123A/D481N). A genetically complemented *B. anthracis* ftsE(K123A/D481N) strain was generated. The wild-type *B. anthracis* parent strain *ftsE* gene was inserted into plasmid pUTE973 (55) to generate plasmid pUVA424, in which *ftsE* was under the control of an IPTG-inducible promoter. Because the *B. anthracis* ftsE(K123A/D481N) mutant was generated by chromosomal mutations, *B. anthracis* ftsE(K123A/D481N)/pUVA424 carried both mutated and wild-type copies of *ftsE*.

Phenotypically, the *B. anthracis* parent strain grew as long, smooth chains of bacilli that were not affected by the presence and/or expression of pUTE973 or pUVA424 (see Fig. S2A to F in the supplemental material). In comparison, the *B. anthracis* ftsE(K123A/D481N) mutant bacilli resembled the *B. anthracis* Δ*ftsX* mutant (35), with a “kinked” appearance (see Fig. S2G). Addition of IPTG alone in the absence of any plasmids had no effect on the mutant phenotype of *B. anthracis* ftsE(K123A/D481N) (see Fig. S2I). The similarity in morphology between the ftsE(K123A/D481N) and Δ*ftsX* strains was as expected for mutations within the same functional complex. This morphological phenotype was also observed in similarly constructed *B. subtilis* mutants (40). The *B. anthracis* ftsE(K123A/D481N) mutant carrying the empty control vector (pUTE973) or an *ftsE* complementation plasmid (pUVA424) with no IPTG induction had the morphological phenotype of the ftsE(K123A/D481N) mutant strain (see Fig. S2H, I, and K). Induction of *ftsE* gene expression with IPTG generated a morphological phenotype similar to that of the *B. anthracis* parent strain (see Fig. S2L). A 24-h growth curve revealed that the ftsE(K123A/D481N) mutant strain had a slightly longer lag phase than the parent strain prior to initiating log-phase.
The concentrations tested (Fig. 4B), similar to the observed with the ftsE mutant (designated as “ftsE” in this figure) had a slightly longer lag phase prior to initiating log-phase growth than the B. anthracis parent strain, a growth characteristic similar to that observed in the B. anthracis ΔftsX mutant (35). (B) The B. anthracis ftsE(K123A/D481N) mutant was relatively resistant to CXCL10 and resistant to CTTC at all of the concentrations tested, from 0 to 2.8 μM. Results were similar to those observed with the B. anthracis ΔftsX mutant. A statistically significant difference between CXCL10 and CTTC was observed at concentrations of ≥0.9 μM. (C) The B. anthracis ftsE(K123A/D481N) mutant transformed with IPTG-inducible plasmid pUTE973 (empty vector control) exhibited resistance to 0.9 μM CXCL10 in the absence or presence of 25 μM IPTG. Transformation of the B. anthracis ftsE(K123A/D481N) mutant with IPTG-inducible plasmid pUVA424 (ftsE complementation vector) resulted in restoration of susceptibility to 0.9 μM CXCL10 only upon treatment with 25 μM IPTG to induce gene expression. (D) The B. anthracis ftsE(K123A/D481N) mutant transformed with plasmid pUTE973 exhibited resistance to 2.3 μM CTTC in the absence or presence of 25 μM IPTG. The B. anthracis ftsE(K123A/D481N) mutant transformed with pUVA424 exhibited susceptibility to 2.3 μM CTTC only upon the addition of 25 μM control. Data points represent mean values ± the standard errors of the means; n = 3 separate experiments using IPTG to induce gene expression. Bacterial viability was measured by alamarBlue reduction, and fluorescence is expressed as a percentage of that of the strain-specific untreated triplicate wells in each experiment. *, P ≤ 0.01; **, P ≤ 0.0001; ****, P < 0.0001.

FIG 4 The B. anthracis ftsE(K123A/D481N) mutant was resistant to CXCL10 and CTTC, in contrast to the parent strain. (A) The B. anthracis ftsE(K123A/D481N) mutant exhibited greater resistance to CXCL10 and CTTC as compared to the parent strain (Fig. 4B). The kinked morphology of ftsE(K123A/D481N) and that previously reported for ΔftsX (35) suggested that FtsE and FtsX play a key role in cell wall synthesis and/or cell division, as has been observed in other bacterial species (39, 56, 57).

The B. anthracis ftsE(K123A/D481N) mutant is resistant to CXCL10 and CTTC. To test the potential role of FtsE, as part of an active FtsE/X complex, in CXCL10-mediated antimicrobial activity, the B. anthracis ftsE(K123A/D481N) mutant was exposed to CXCL10 or CTTC at various concentrations. The B. anthracis ftsE(K123A/D481N) mutant exhibited greater resistance to CXCL10-mediated killing compared to the parent strain (Fig. 4B). This resistance of the ftsE(K123A/D481N) mutant was overcome by higher concentrations of CXCL10 (Fig. 4B), similar to what was observed with the ΔftsX mutant (Fig. 3B). The B. anthracis ftsE(K123A/D481N) mutant strain was resistant to CTTC at all of the concentrations tested (Fig. 4B), similar to the ΔftsX mutant (Fig. 3B). IPTG-induced expression of the ftsE-containing vector (pUVA424) in the ftsE(K123A/D481N) mutant strain restored susceptibility to CXCL10 at 0.9 μM (Fig. 4C) and to CTTC at 2.3 μM (Fig. 4D). In contrast, IPTG induction of the empty vector control (pUTE973) in the ftsE(K123A/D481N) mutant strain did not restore susceptibility to CXCL10 or CTTC (Fig. 4C and D). These data suggest that a functional FtsE/X complex is important for the antimicrobial effect of CXCL10. Importantly, the ftsE gene was intact (i.e., wild type) in the ftsE(K123A/D481N) mutant strain in these experiments. Therefore, the data indicate that interaction of CXCL10 or CTTC with B. anthracis FtsX was not, in and of itself, sufficient for killing. Instead, an active FtsE/X complex was necessary for FtsX-mediated killing. Table 1 summarizes the calculated 50% effective concentration (EC50) values of CXCL10 and CTTC against the B. anthracis parent, ΔftsX, and ftsE(K123A/D481N) strains.

B. anthracis parent strain membrane depolarization in response to CXCL10 or CTTC exposure. The CXCL10 C-terminal α-helix does not appear to be required for the FtsE/X-mediated antimicrobial activity of CXCL10 in B. anthracis. However, it is likely that it plays a role in the non-FtsE/X-dependent mechanism of killing. Therefore, membrane depolarization studies were con-
TABLE 1 Calculated EC_{50} values of CXCL10 and CTTC for the B. anthracis parent, ΔftsX, and ftsE(K123A/D481N) strains\textsuperscript{a}

| Strain                | CXCL10 EC_{50} (\mu M) | CTTC EC_{50} (\mu M) | Fold change |
|-----------------------|------------------------|----------------------|-------------|
| Parent                | 0.68 ± 0.03            | 1.83 ± 0.07          | 2.7         |
| ΔftsX mutant          | 1.59 ± 0.05            | NA\textsuperscript{b} | NA          |
| ftsE(K123A/D481N) mutant | 1.39 ± 0.07           | NA                   | NA          |

\textsuperscript{a} The fold difference between the EC\textsubscript{50} values of CXCL10 and CTTC for the B. anthracis parent strain is shown. The mutant strains showed no susceptibility to CTTC; thus, there is no EC\textsubscript{50} or fold change compared to CXCL10.

\textsuperscript{b} NA, not applicable because no killing was observed.

The membrane of the B. anthracis parent strain underwent depolarization in the presence of CXCL10. The extent of depolarization increased with increasing concentrations of CXCL10 (Fig. 5A). Depolarization occurred to a lesser but statistically significant degree in the presence of CTTC (Fig. 5A). The degree of depolarization also increased with increasing concentrations of CTTC (Fig. 5A). Statistical analysis was conducted by comparing untreated and treated groups through a two-way analysis of variance (ANOVA). The effect of CTTC on the parent strain may indicate that some other portion of CXCL10, not just the α-helix, can produce depolarization or that cell lysis mediated through the FtsE/X-mediated pathway also leads to depolarization. The corresponding viability from CFU determination was plotted on a log scale and showed that the cells died upon exposure to either CXCL10 or CTTC (Fig. 5C). Because of an inoculum effect resulting from the higher bacterial concentration used in this particular assay relative to the concentration of CXCL10 or CTTC, the degree of bacterial killing was less than that observed in the standard antimicrobial assay shown in Fig. 3.

Membrane depolarization of the B. anthracis ΔftsX mutant occurs only in the presence of the CXCL10 α-helix. CXCL10 caused significant membrane depolarization of the B. anthracis ΔftsX mutant, in contrast to the untreated control (Fig. 5B). However, no significant membrane depolarization was observed when the B. anthracis ΔftsX mutant was treated with CTTC (Fig. 5B). These data suggested that the membrane depolarization induced by CTTC in the parent strain was dependent upon the presence of FtsX. Thus, an FtsE/X-dependent pathway is likely to lead to a lack of cell integrity with concomitant membrane disruption. A decrease in bacterial viability was correlated with membrane depolarization upon incubation with CXCL10; a less pronounced decrease in viability was observed with CTTC (Fig. 5D). These results indicated that the C-terminal α-helix of CXCL10 was capable of causing membrane depolarization and that its activity was independent of FtsX.

**DISCUSSION**

The antimicrobial activity of the interferon-inducible ELR(−) CXC chemokine CXCL10 has been demonstrated against a broad range of Gram-positive (including B. anthracis, L. monocytogenes, and S. aureus) and Gram-negative (E. coli and Pseudomonas aeruginosa) pathogens, but the exact mechanism(s) of action has remained unclear (14–16, 18, 19). In the present study, we showed that CXCL10 can kill B. anthracis through multiple mechanisms. One mechanism involves the bacterial FtsE/X complex and is mediated through interaction with the N-terminal region(s) of CXCL10. It is independent of the amphipathic C-terminal α-helix. Another mechanism is FtsE/X independent and is mediated by the depolarization of the cytoplasmic membrane by the CXCL10 C-terminal α-helix. Such findings are novel and unexpected in the chemokine field and open up a number of mechanistic possibilities for how chemokines target microbes beyond nonspecific membrane disruption, while also supporting defensin-like antibacterial targeting as a second mechanism.

Previous electron microscopy studies have shown that CXCL10 localizes to the bacterial membrane in vegetative cells and to the internal coat-cortex interface in spores (17). That an extracellular region of FtsX, a membrane protein, had limited similarity to CXCR3 suggested a possible interaction with CXCL10 through receptor mimicry. This observation led to the hypothesis that the N terminus of CXCL10 is important for interaction with the target microorganism through that region of FtsX, especially since it is the portion of the chemokine that facilitates interaction with CXCR3. A peptide competition assay showed the CXCR3-like region of FtsX was indeed important in interacting with CXCL10 and mediating antimicrobial activity. We then tested CTTC to determine if the N-terminal region of CXCL10 promotes antimicrobial activity or if the C-terminal α-helix of CXCL10 is necessary for killing. CTTC was less active against the B. anthracis parent strain and did not kill the ΔftsX mutant strain. The EC_{50} values indicated that CTTC is 2.7-fold less effective than CXCL10 at killing the parent strain and lacks the ability to kill the ΔftsX mutant strain altogether (Table 1). These results indicated that an FtsX-dependent mechanism of killing was independent of the CXCL10 C-terminal α-helix, highlighting a novel interaction.
between the CXCR3 receptor-binding region(s) of CXCL10 with bacterial FtsX (Fig. 6A), as chemokine antimicrobial activity has previously been hypothesized to be mediated through the C-terminal α-helix (19, 33, 34). The finding that other portions of CXCL10 may mediate killing represents a paradigm shift and opens avenues for the study and development of new therapies based on chemokine architecture.

FtsE/X is a unique bacterial complex that warrants further study as an antimicrobial target since its pivotal role in PG processing through hydrolase activation is widely conserved among bacterial species (38–41, 43, 60). Interestingly, there are differences between Gram-positive and Gram-negative bacteria in the roles of FtsE/X (37–41, 49). In Gram-positive bacteria, namely, *Bacillus* spp., there appear to be redundancies in the enzymatic activities involved in PG processing during cellular elongation, such that a disruption in one pathway could result in cell wall instability and bacterial cell lysis (44, 61, 62). Notably, the *B. anthracis* ftsE(K123A/D481N) mutant in our study exhibited a morphology similar to that previously reported for the ΔftsX mutant, in which bacilli appeared “kinked,” with frequent curves and sharp angles among the growing chains, consistent with an alteration in cell wall processing (35, 39, 56, 57). The interaction of CXCL10 with FtsE/X may provoke or disrupt PG processing from which the bacteria cannot recover, resulting in bacterial cell death.

Our results determined that CXCL10 FtsX-dependent antimicrobial activity requires both FtsX and a functional FtsE ATP-binding protein, thus implicating an energy requirement for the CXCL10-mediated mechanism. This report is the first description of an active FtsE/X complex contributing to the bactericidal effect of an antimicrobial molecule. The membrane disruption studies described here show that depolarization also occurred through the FtsX-mediated mechanism. Given the role of FtsE/X in PG processing, it is possible that CXCL10 provokes PG turnover, which leads to cell lysis through PG degradation and the loss of the cell’s

![Membrane depolarization occurs after exposure of *B. anthracis* to CXCL10.](image)

**FIG 5** Membrane depolarization occurs after exposure of *B. anthracis* to CXCL10. (A) *B. anthracis* parent strain membrane depolarization occurs after exposure to CXCL10 or CTTC. (B) *B. anthracis* ΔftsX mutant membrane depolarization occurs after exposure to CXCL10 but not upon exposure to CTTC. The positive control (LL37 at 25 μM) induced membrane depolarization. Exposure to the negative control (CCL5 at 2.8 μM) did not cause membrane depolarization (n = 4 or 5 separate experiments). (C) Killing of the bacteria by CXCL10 or CTTC was correlated with the membrane depolarization assay results obtained under the same conditions (n = 4 or 5 separate experiments). Numbers of CFU per milliliter are plotted on a log scale. (D) Treatment of the *B. anthracis* ΔftsX mutant with CXCL10 or CTTC at the concentrations indicated results in less killing than treatment of the parent strain shown in panel A (n = 4 or 5 separate experiments). Incubation with the positive control (LL37) resulted in less survival of both strains than of the untreated control strain. The negative control (CCL5) was not bactericidal (n = 4 or 5 separate experiments). Statistical analysis was performed by two-way ANOVA. ns, not significant; *, P ≤ 0.01; **, P ≤ 0.001; ***, P ≤ 0.0001; ****, P < 0.0001.
ability to withstand environmental conditions. Studies are under way in our laboratory to investigate the nature of this interaction and how bacterial cell death may occur in response to CXCL10, potentially from provoking or disrupting the PG processing pathway. Understanding the function of the unique and widely conserved FtsE/X complex may lead to the utilization of this novel bacterial target for new antimicrobial therapies.

The FtsE/X-independent mechanism of killing by CXCL10 is likely mediated through the nonspecific interaction of its C-terminal α-helix with the bacterial membrane, similar to what has been described for cationic antimicrobial peptides such as LL37 or defensins (19, 33, 34) (Fig. 6B). The B. anthracis ΔftsX mutant only undergoes membrane depolarization in the presence of intact CXCL10, suggesting that the amphipathic structural characteristics of the C-terminal region are responsible for membrane disruption and lysis. It appears that the CXCL10 C-terminal α-helix is not necessary for killing of the parent strain. However, in the absence of FtsX and at higher concentrations of CXCL10, death can occur through membrane disruption. When both mechanisms are present/active, CXCL10-mediated bacterial killing appears to be the most effective. Further study of the structural characteristics of CXCL10 and its related chemokines should lead to a better understanding of their biological activities, as well as a broader application of therapeutic potential for chemokine α-helices against a wide range of bacterial species.

Our findings represent an exciting advance in the understanding of how chemokines function as antimicrobial molecules, as well as in the study of host-pathogen interactions within the innate immune system. The primary significance is the new information involving the bifunctional activity of CXCL10, with mechanisms involving a previously unrecognized portion of CXCL10 that exhibits antimicrobial activity through interaction with the novel bacterial target FtsE/X and also the CXCL10 C-terminal α-helix-mediated antimicrobial killing of B. anthracis. Our research suggests that as chemokines are further studied as antimicrobial molecules, there will be heightened recognition of more specific mechanisms by which they kill microorganisms beyond nonspecific membrane targeting and disruption. Studies of the antimicrobial effects of chemokines are likely to reveal additional mechanisms of killing and lead to the development of novel small molecules as innovative therapies to treat a wide range of bacterial pathogens, including already established and emerging antibiotic-resistant organisms.

MATERIALS AND METHODS

Bacterial strains and culture conditions. B. anthracis Sterne strain ATCC 7702 (pXO1+ pXO2−; American Type Culture Collection, Manassas, VA) was designated the parent strain and used in these experiments. The B. anthracis ΔftsX, ΔftsX/pUTE973, ΔftsX/pUA113, and ΔftsX(K123A/D481N) mutant strains were derived from the Sterne strain (35). Vegetative cells were prepared by inoculating B. anthracis parent strain spore stocks or B. anthracis ΔftsX and ΔftsX(K123A/D481N) mutant vegetative cell frozen stocks into 10 ml of brain heart infusion (BHI) broth (Difco, Franklin Lakes, NJ) and incubating them at 37°C with shaking (250 rpm). Mid-log-phase cultures were prepared on the next day by diluting the overnight cultures 1:20 in fresh BHI broth and incubating them at 37°C with shaking for ~2 h until an optical density at 600 nm between 0.6 and 0.65 was achieved, at which time the cultures were used for experiments. All laboratory work involving B. anthracis Sterne or Sterne-derived strains was approved by the University of Virginia Institutional Biosafety Committee. Biosafety level 2 practices were used for all work involving B. anthracis Sterne or Sterne-derived strains.

Amino acid sequence alignment. The National Library of Medicine Basic Local Alignment Search Tool (http://blast.ncbi.nlm.nih.gov/ Blast.cgi) sequence alignment program was used to align Gram-positive and Gram-negative FtsX protein sequences with B. anthracis Sterne (parent) strain FtsX (see Text S1 in the supplemental material). The following species and FtsX identification codes were used to perform individual BLASTP analyses: B. anthracis Sterne (AAT57322.1), B. subtilis 168 (NP_391405.1), M. tuberculosis H37Rv (CAAA49620.1), S. pneumoniae (AJD71681.1), E. coli F11 (EDV68789.1), and K. pneumoniae (KLA37751.1). Alignment of the full sequences of all six strains was conducted with the T-Coffee multiple-sequence alignment software (see Text S1) (http://tcoffee.crg.cat/apps/tcoffee/do:regular) and formatted with the Boxshade program (http://www.ch.embnet.org/software/BOX_form.html) (47, 48).
Generation of *B. anthracis* *ftsE*(K123A/D481N) and plasmid vector pUV424 for gene complementation studies with *B. anthracis* *ftsE*(K123A/D481N). Point mutations of the *B. anthracis* gene *ftsE* were performed as previously described (35, 52). Briefly, *B. anthracis* *ftsE* was cloned into plasmid vector pGEM in α-select silver efficiency *E. coli* (Bio-line, Taunton, MA) where Walker A and B mutations were performed (see Text S2A in the supplemental material). The mutated *ftsE* gene was amplified by PCR and sequenced to verify the mutations in the Walker A and B motifs. The bacterial strain with the two point mutations (K123A and D481N) in the *ftsE* Walker A and B motifs was designated *ftsE*(K123A/D481N).

Complementation of the *B. anthracis* *ftsE*(K123A/D481N) mutant strain was performed as previously described in detail (35). The generated *ftsE*(K123A/D481N) complementation plasmid (pUV424) or an empty control plasmid vector (pUTE973) was electroporated into *B. anthracis* *ftsE*(K123A/D481N) bacilli. All isolates were tested and verified by PCR and sequencing.

**CXCL10 and CTTC.** Recombinant human CXCL10 was purchased from PeproTech (Rocky Hill, NJ). CXCL10 was dissolved in sterile H$_2$O containing 0.3% (wt/vol) human serum albumin (HSA; Grifols, Los Angeles, CA) at a concentration of 1 mg ml$^{-1}$ and stored at −80°C in 10-μl aliquots to limit freeze-thaw cycles to no more than one. The CXCL10 peptide was commercially synthesized and capped by amidation on the C terminus by United Biosystems (Herndon, VA) (see Text S2B in the supplemental material). Lyophilized CTTC was dissolved in sterile H$_2$O at a concentration of 5 mg ml$^{-1}$ and stored in 20-μl aliquots as described above. Sequence and structure verification was carried out as described in Text S2B.

**Antimicrobial assays and light microscopy studies.** Vegetative cells were diluted to approximately 2.5 × 10$^5$ CFU ml$^{-1}$ in fresh Dulbecco's modified Eagle's medium ( Gibco-Invitrogen, Carlsbad, CA) containing 10% (vol/vol) fetal bovine serum (HyClone, Logan, UT) and the various concentrations of CXCL10 or CTTC were added. Aliquots of 100 μl were placed in triplicate wells of a 96-well plate and incubated at 37°C with 5% CO$_2$ for approximately 4 h, unless otherwise noted, at which point alamarBlue dye (ABD Serotech, Oxford, United Kingdom) was added at a 1:10 dilution and allowed to reach visual saturation in an untreated control under protection from light. Active metabolism of the vegetative cells was quantified by the generation of a fluorescent signal from the reduction of the alamarBlue oxidation-reduction dye by measuring the fluorescence at 530-nm excitation and 590-nm emission wavelengths with a fluorescence plate reader (Bio-tek Instruments, Winooski, VT) (see Text S2C). The antimicrobial activity was quantified via the generation of a fluorescent signal from the reduction of the alamarBlue oxidation-reduction dye by measuring the fluorescence at 530-nm excitation and 590-nm emission wavelengths with a fluorescence plate reader (Bio-tek Instruments, Winooski, VT) (see Text S2C).

**Membrane depolarization measurement.** After the exposure of vegetative cells to various micromolar concentrations of CXCL10 or CTTC, membrane depolarization was measured by a modification of a previously published protocol (59). Controls are discussed in Text S2C in the supplemental material. Briefly, *B. anthracis* cells were grown to mid-log phase as described above. Cells were collected and resuspended in respiration buffer, and 0.1 mM diS-C$_3$-3 (AnaSpec, Fremont, CA) was added to cell aliquots. Initial baseline fluorescence ($F_0$) readings, test antimicrobial fluorescence (*F*) readings, and maximum depolarization (*F$_{max}$) readings were collected every 5 s for 150 to 300 s. Using the initial (*F$_0$), test (*F*), and final, maximum fluorescence readings (*F$_{max}$), the values of each group were averaged and the following calculation was used to determine percent depolarization by the test molecule compared to complete depolarization by melittin: % Depolarization = [(*F$_{max}$ − *F*)/(*F$_{max}$ − *F$_0$)] × 100 (59). In parallel with the above-described depolarization experiments, the numbers of CFU per milliliter were determined to measure bacterial survival (see Text S2C).

**Statistical analyses.** Statistical analyses and graphing were performed with the GraphPad Prism 4.0 and 6.0 software. Experimental groups were analyzed with an unpaired, two-tailed Student t test or two-way ANOVA, as noted in the figure legends. Differences were considered significant at a *P* value of <0.05.

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at http://mbio.asm.org/lookup/suppl/doi:10.1128/mBio.00334-16/-/DCSupplemental.

Figure S1, TIF file, 0.1 MB.

Figure S2, TIF file, 1.4 MB.

Text S1, DOCX file, 0.02 MB.

Text S2, DOCX file, 0.02 MB.

**ACKNOWLEDGMENTS**

We thank Debra J. Fisher for her technical assistance and Borina Mehrad, Barbara J. Mann, Girija Ramakrishnan, and members of their laboratories at the University of Virginia School of Medicine for their advice and helpful discussions. We thank Stephen A. Morse (Centers for Disease Control and Prevention, Atlanta, GA) for providing helpful comments on the manuscript.

**FUNDING INFORMATION**

This work was supported by National Institutes of Health/National Institute of Allergy and Infectious Diseases grants R01 AI099097 (M.A.H.) and R21 AI072469 (M.A.H.) and National Institutes of Health grant T32 AI055432 (K.R.M.). The funders had no role in the study design, data collection and interpretation, or the decision to submit the work for publication. This work used the Aviv 410 CD spectropolarimeter in the Shared Instrumentation Core, which is supported by the University of Virginia School of Medicine.

**REFERENCES**

1. Mortality and Causes of Death Collaborators. 2015. Global, regional, and national age-sex specific all-cause and cause-specific mortality for 240 causes of death, 1990–2013: a systematic analysis for the Global Burden of Disease Study 2013. Lancet 385:117–171. http://dx.doi.org/10.1016/S0140-6736(14)61682-2.

2. Luster AD. 1998. Chemokines-chemotactic cytokines that mediate inflammation. N Engl J Med 338:436–445. http://dx.doi.org/10.1056/NEJM199802123380706.

3. Bonecchi R, Galliera E, Borroni EM, Corsi MM, Locati M, Mantovani A. 2009. Chemokines and chemokine receptors: an overview. Front Biosci (Landmark Ed) 14:540–551. http://dx.doi.org/10.2741/3261.

4. Griffith JW, Sokol CL, Luster AD. 2014. Chemokines and chemokine receptors: positioning cells for host defense and immunity. Annu Rev Immunol 32:659–702. http://dx.doi.org/10.1146/annurev-immunol-032713-120145.

5. Zlotnik A, Yoshie O. 2000. Chemokines: a new classification system and their role in immunity. Immunity 12:121–127. http://dx.doi.org/10.1016/S1074-7613(00)01065-X.

6. Zlotnik A, Yoshie O. 2012. The chemokine superfamily revisited. Immunity 36:705–716. http://dx.doi.org/10.1016/j.immuni.2012.05.008.

7. Allen SJ, Crown SE, Handel TM. 2007. Chemokine: receptor structure,
24. Mehrad B, Keane MP, Strieter RM. 2007. Chemokines as mediators of angiogenesis. Thromb Haemost 97:755–762. http://dx.doi.org/10.1160/TH07-01-0040.

25. Lacotte S, Brun S, Muller S, Dumortier H. 2009. CXCR3, inflammation, and autoimmune diseases. Ann N Y Acad Sci 1173:310–317. http://dx.doi.org/10.1111/j.1546-1672.2009.04013.x.

26. Dur M, Peschel A. 2002. Chemokines meet defenses: the merging concepts of chemotacticants and antimicrobial peptides in host defense. Infect Immun 70:6515–6517. http://dx.doi.org/10.1128/IAI.70.12.6515-6517.2002.

27. Yang D, Chen Q, Hoover DM, Staley P, Tucker KD, Lubkowski J, Crawford MA, Zhu Y, Green CS, Burdick MD, Sanz P, Alem F, O’Brien Oppenheim JJ, Biragyn A, Kwak LW, Yang D. 2009. Review. The CXCR3 binding chemokine IP-10/CXCL10: structure and receptor antagonistic functions. Immunol Cell Biol 87:259–266. http://dx.doi.org/10.1038/icb.2009.15.

28. Oppenheim JJ, Biragyn A, Kwak LW, Yang D. 2003. Roles of antimicrobial peptides such as defensins in innate and adaptive immunity. Ann Rheum Dis 62(Suppl 2):ii17–ii21. http://dx.doi.org/10.1136/ard.supp1_2.i17.

29. Yung SC, Murphy PM. 2012. Antimicrobial chemokines. Front Immunol 3:276. http://dx.doi.org/10.3389/fimmu.2012.00276.

30. Vaidehi N, Bhattacharya S, Larsen AB and antagonistic functions. Immunol Cell Biol 87:259–266. http://dx.doi.org/10.1038/icb.2009.15.

31. Booth V, Keizer DW, Kamphuis MB, Clark-Lewis I, Sykes BD. 2002. The CXCR3 binding chemokine IP-10/CXCL10: structure and receptor interactions. Biochimie 41:10418–10425. http://dx.doi.org/10.1016/j.bita.2002.06.020.

32. Trott A, Costantini S, Colonna G. 2009. Modelling of the membrane receptor CXCR3 and its complexes with CXCL9, CXCL10 and CXCL11 chemokines: putative target for new drug design. Mol Immunol 47:332–339. http://dx.doi.org/10.1016/j.molimm.2009.09.013.

33. Loetscher M, Loetscher P, Brass N, Meese E, Moser B. 1998. Lymphocyte-specific chemokine receptor CXCR3: regulation, chemokine binding and gene localization. Eur J Immunol 28:3696–3705. http://dx.doi.org/10.1002/(SICI)1521-4141(199811)28:11<3696::AID-IMMU3696>3.0.CO;2-W.

34. Groom JR, Luster AD. 2011. CXCR3 ligands: redundant, collaborative and antagonistic functions. Immunol Cell Biol 89:207–215. http://dx.doi.org/10.1038/icb.2010.138.
organization of immunity. Immunol Rev 195:58–71. http://dx.doi.org/10.1034/j.1600-065X.2003.00067.x.

47. Notredame C, Higgins DG, Heringa J. 2000. T-Coffee: a novel method for fast and accurate multiple sequence alignment. J Mol Biol 302:205–217. http://dx.doi.org/10.1006/jmbi.2000.4042.

48. Di Tommaso P, Moretti S, Xenarios I, Orosbitg M, Montanyoila A, Chang J-M, Taly J-F, Notredame C. 2011. T-Coffee: a web server for the multiple sequence alignment of protein and RNA sequences using structural information and homology extension. Nucleic Acids Res 39:W13–W17. http://dx.doi.org/10.1093/nar/gkr245.

49. Yang DC, Peters NT, Parzych KR, Uehara T, Markovski M, Bernhardt TG. 2011. An ATP-binding cassette transporter-like complex governs cell-wall hydrolysis at the bacterial cytokinetic ring. Proc Natl Acad Sci U S A 108:e1052-60. http://dx.doi.org/10.1073/pnas.1107780108.

50. Shahabuddin S, Ji R, Wang P, Brailoiu E, Dun N, Yang Y, Aksoy MO, Kelsen SG, Aksoy MO, Dun N. 2006. CXCR3 chemokine receptor-induced chemotaxis in human airway epithelial cells: role of p38 MAPK and PI3K signaling pathways. Am J Physiol Cell Physiol 291:C34–C39. http://dx.doi.org/10.1152/ajpcell.00441.2005.

51. Greenfield NJ. 2006. Using circular dichroism spectra to estimate protein secondary structure. Nat Protoc 1:2876–2890. http://dx.doi.org/10.1038/nprot.2006.202.

52. Janes BK, Stibitz S. 2006. Routine markerless gene replacement in Bacillus anthracis. Infect Immun 74:1949–1953. http://dx.doi.org/10.1128/IAI.74.3.1949-1953.2006.

53. Sham LT, Jensen KR, Bruce KE, Winkler ME. 2013. Involvement of FtsE ATPase and FtsX extracellular loops 1 and 2 in FtsEX-PcsB complex function in cell division of Streptococcus pneumoniae D39. mBio 4.e00431–13. http://dx.doi.org/10.1128/mBio.00431-13.

54. Mir MA, Rajeswari HS, Veeraraghavan U, Ajitkumar P. 2006. Molecular characterisation of ABC transporter type FtsE and FtsX proteins of Mycobacterium tuberculosis. Arch Microbiol 185:147–158. http://dx.doi.org/10.1007/s00203-005-0079-z.

55. Pfughoefl KJ, Sumby P, Koehler TM. 2011. Bacillus anthracis sin locus and regulation of secreted proteases. J Bacteriol 193:631–639. http://dx.doi.org/10.1128/JB.01083-10.

56. Rueff A-S, Chastanet A, Dominguez-Escobar J, Yao Z, Yates J, Prejean M-V, Delumeau O, Noirot P, Wedlich-Soldner R, Filipe SR, Carballido-López R. 2014. An early cytoplasmic step of peptidoglycan synthesis is associated to MreB in Bacillus subtilis. Mol Microbiol 91:348–362. http://dx.doi.org/10.1111/mmi.12467.

57. Formstone A, Errington J. 2005. A magnesium-dependent mreB null mutant: implications for the role of mreB in Bacillus subtilis. Mol Microbiol 55:1646–1657. http://dx.doi.org/10.1111/j.1365-2958.2005.04506.x.

58. Su M, He C, West CA, Mentzer SJ. 2001. Cytolytic peptides induce biphasic permeability changes in mammalian cell membranes. J Immunol Methods 252:63–71. http://dx.doi.org/10.1016/S0022-1759(01)00334-9.

59. Chongsrirawatana NP, Barron AE. 2010. Comparing bacterial membrane interactions of antimicrobial peptides and their mimics. Methods Mol Biol 618:171–182. http://dx.doi.org/10.1007/978-1-60761-594-1_12.

60. Sham LT, Barendt SM, Kopecky KE, Winkler ME. 2011. Essential PcsB putative peptidoglycan hydrolase interacts with the essential FtsXSpn cell division protein in Streptococcus pneumoniae D39. Proc Natl Acad Sci U S A 108:E1061–E1069. http://dx.doi.org/10.1073/pnas.1108323108.

61. Novak R, Charpentier E, Braun JS, Tuomanen E. 2000. Signal transduction by a death signal peptide: uncovering the mechanism of bacterial killing by penicillin. Mol Cell 5:49–57. http://dx.doi.org/10.1016/S1097-2765(00)80402-5.

62. Yount NY, Yeaman MR. 2013. Peptide antimicrobials: cell wall as a bacterial target. Ann N Y Acad Sci 1277:127–138. http://dx.doi.org/10.1111/nyas.12005.