Structural Basis of Ca\textsuperscript{2+}-Dependent Self-Processing Activity of Repeat-in-Toxin Proteins

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\textbf{ABSTRACT} The posttranslational Ca\textsuperscript{2+}-dependent “clip-and-link” activity of large repeat-in-toxin (RTX) proteins starts by Ca\textsuperscript{2+}-dependent structural rearrangement of a highly conserved self-processing module (SPM). Subsequently, an internal aspartate-proline (Asp-Pro) peptide bond at the N-terminal end of SPM breaks, and the liberated C-terminal aspartyl residue can react with a free ε-amino group of an adjacent lysine residue to form a new isopeptide bond. Here, we report a solution structure of the calcium-loaded SPM (Ca-SPM) derived from the FrpC protein of \textit{Neisseria meningitidis}. The Ca-SPM structure defines a unique protein architecture and provides structural insight into the autocatalytic cleavage of the Asp-Pro peptide bond through a “twisted-amide” activation. Furthermore, in-frame deletion of the SPM domain from the ApxIVA protein of \textit{Actinobacillus pleuropneumoniae} attenuated the virulence of this porcine pathogen in a pig respiratory challenge model. We hypothesize that the Ca\textsuperscript{2+}-dependent clip-and-link activity represents an unconventional strategy for Gram-negative pathogens to adhere to the host target cell surface.

\textbf{IMPORTANCE} The Ca\textsuperscript{2+}-dependent clip-and-link activity of large repeat-in-toxin (RTX) proteins is an exceptional posttranslational process in which an internal domain called a self-processing module (SPM) mediates Ca\textsuperscript{2+}-dependent processing of a highly specific aspartate-proline (Asp-Pro) peptide bond and covalent linkage of the released aspartyl to an adjacent lysine residue through an isopeptide bond. Here, we report the solution structures of the Ca\textsuperscript{2+}-loaded SPM (Ca-SPM) defining the mechanism of the autocatalytic cleavage of the Asp414-Pro415 peptide bond of the \textit{Neisseria meningitidis} FrpC exoprotein. Moreover, deletion of the SPM domain in the ApxIVA protein, the FrpC homolog of \textit{Actinobacillus pleuropneumoniae} attenuated the virulence of this porcine pathogen in a pig infection model, indicating that the Ca\textsuperscript{2+}-dependent clip-and-link activity plays a role in the virulence of Gram-negative pathogens.

\textbf{KEYWORDS} RTX toxins, cell adhesion, clip-and-link, host-pathogen interactions, nuclear magnetic resonance

The Ca\textsuperscript{2+}-dependent “clip-and-link” activity represents a distinct type of posttranslational autoprocessing of proteins. It consists of Ca\textsuperscript{2+}-induced autocatalytic cleavage of a specific Asp-Pro peptide bond within a polypeptide precursor that generates a reactive C-terminal Asp anhydride, which can then react with an ε-amino group of an adjacent lysine residue to form a new covalent isopeptide bond (1). This process is mediated by a self-processing module (SPM), a 177-residue-long polypeptide domain...
that is exclusively present in a specific subset of repeat-in-toxin (RTX) proteins of Gram-negative bacteria (2). These very large proteins (>1,000 residues) are secreted by a dedicated type I secretion system (T1SS) that recognizes a C-terminal noncleavable secretion signal preceded by a variable number of tandem repeats of a consensus motif GGXGDXXXX (3). These RTX repeats form a calcium-binding parallel β-roll structure that folds vectorially, from the C to N terminus in the course of secretion, as the RTX substrates exit from the calcium-depleted bacterial cytosol into the calcium-rich extracellular milieu (4, 5). Even though many RTX proteins serve as major virulence factors of Gram-negative pathogens, the RTX proteins carrying the SPM domain were not found to exert any cytotoxic activity on host cells. Possibly, the Ca²⁺-dependent clip-and-link activity of FrpC, an iron-regulated RTX protein of Neisseria meningitidis, might be involved in the interaction of the bacterium with the host cell surface (6). Recent biochemical and biophysical characterization of the SPM domain of FrpC revealed that the Ca²⁺-dependent clip-and-link activity results from the Ca²⁺-induced folding of the SPM polypeptide (2). Three EF-hand-like Ca²⁺-binding loops were predicted to be a key feature of an SPM structure, which appears to be stabilized by a noncovalent π-π interaction between two tryptophan residues (W₄₅₁ and W₅₁₉) arranged in a T-shaped orientation. Its formation involves a cooperative structural transition from an unfolded conformation to a well-folded state that enables a highly specific processing of the peptide bond between the Asp₄₁₄ residue and the N-terminal Pro₄₁₅ residue of the SPM amino acid sequence (residues 415 to 591 of FrpC). As a result, the D₄₁₄-P₄₁₅ bond breaks, and a reactive anhydride is formed at the newly released D₄₁₄ residue, which upon nucleophilic attack by an ε-amino group of an adjacent polypeptide chain can form a new covalent isopeptide bond (Fig. 1A). Since the cross-linking activity of SPM can be inhibited by nucleophile scavengers, such as the reducing agents with a free thiol group (e.g., 1,4-dithiothreitol), the self-processing activity of SPM could be used for development of a self-excising affinity tag that allows purification of untagged recombinant proteins in a single chromatographic step (7). However, the structural and mechanistic aspects of the Ca²⁺-dependent clip-and-link activity of large RTX proteins remained unknown.

Here, we determined the solution structure of the calcium-loaded SPM (Ca-SPM) domain derived from the FrpC protein of Neisseria meningitidis and propose the mechanism of Ca²⁺-dependent autocatalytic processing of the D₄₁₄-P₄₁₅ peptide bond. We further show that abrogation of the Ca²⁺-dependent clip-and-link activity of ApxIVA, an exclusively in vivo-expressed RTX protein of Actinobacillus pleuropneumoniae, results in attenuation of the virulence of the pathogen in experimental lung infection in pigs. This is the first demonstration that the Ca²⁺-dependent clip-and-link activity of large RTX proteins plays a role in the virulence of Gram-negative pathogens.

RESULTS

NMR structure of Ca-SPM. To gain insight into the catalytic mechanism of Asp-Pro bond processing, we determined the solution structure of calcium-loaded SPM (Ca-SPM) by nuclear magnetic resonance (NMR) spectroscopy. As shown in Fig. 1B, the ¹H-¹⁵N heteronuclear single-quantum coherence (HSQC) spectrum of Ca-SPM exhibited the broad dispersion of backbone amide cross-peaks typical for structured proteins. The spectrum did not contain residual peaks typical for the ¹H-¹⁵N HSQC spectrum of the Ca²⁺-depleted SPM (Fig. 1B, black contours), thus documenting that the Ca-SPM preparation was free of residual unfolded protein. Conventional triple resonance and nuclear Overhauser effect spectroscopy (NOESY) NMR spectra provided structural information for the whole protein, except of a region of 21 residues between T₄₃₀ and G₄₅₀. These residues were affected by conformational or chemical exchange and likely formed a loop lacking a unique structure. Moreover, deletion of the loop consisting of residues T₄₃₀ to G₄₅₀ did not affect the cleavage capacity of the respective glutathione S-transferase (GST)-SPM fusion protein, indicating that the loop does not contribute to structuring and autocatalytic activity of SPM (see Fig. S1 in the supplemental material). The overlay of 20 low-energy structures based on the 3,412 NOE and 234 dihedral angle...
restraints is displayed in Fig. 1C. The Ca-SPM structure has a compact fold composed of eight anti-parallel \( \beta \)-strands connected by a single helix-turn-helix motif and several surface-exposed turns and loops (Fig. 1D). These structures appear to be stabilized by calcium ions as the overall structure of Ca-SPM is well defined with a backbone root square mean deviation (RMSD) of 0.70 \( \pm \) 0.16 Å (Table 1). However, the most abundant calcium isotope, \( ^{42}\text{Ca} \), is not observable by NMR, and the number and exact position of calcium ion(s) could not be identified by this NMR approach.

**Calcium stoichiometry for Ca-SPM.** In order to determine the number of \( \text{Ca}^{2+} \)-binding sites in SPM, Ca-SPM was dialyzed against a calcium-free buffer to remove free \( \text{Ca}^{2+} \) ions and was next subjected to NMR titration with the metal-chelating agent EDTA. The extensive dialysis of Ca-SPM did not affect the overall fold of the protein, indicating a high stability of the Ca-SPM structure at residual free \( \text{Ca}^{2+} \) concentrations. For precise quantification of bound \( \text{Ca}^{2+} \), we took advantage of the fact that complexion of free EDTA with \( \text{Ca}^{2+} \) ions is manifested by specific peak shifts in the \(^1\text{H}\) NMR
spectra that are associated with changes in the chemical environment of methylene and ethylene protons of the acetyl and ethylenediamine moieties, respectively (8) (Fig. 2A). As demonstrated in Fig. S2, titration of Ca-SPM by EDTA resulted in progressive accumulation of singlet (2.47 ppm) and quartet (3.05 to 2.95 ppm) peaks in the 1H NMR spectra, corresponding to a gradual displacement of Ca2+ ions from Ca-SPM and formation of Ca2+-chelated EDTA (EDTA.Ca2+) complexes. This was further accompanied by a decrease of peak intensities in the aromatic, backbone amide and methyl regions of the 1H NMR spectra, illustrating an EDTA-induced unfolding of Ca-SPM. No peaks of partially folded SPM intermediates were detectable in the 1H-15N HSQC spectra, indicating a cooperative binding of Ca2+ ions (9). Once Ca-SPM was unfolded, additions of EDTA did not further affect the 1HNMR spectra, except of two singlet peaks at 3.5 and 3.15 ppm, which corresponded to proton signals from free EDTA (Fig. S2).

Changes in relative intensities of proton signals were used to determine Ca2+ stoichiometry for Ca-SPM. As shown in Fig. 2B, ~4 molar equivalents of EDTA were required to completely unfold the Ca-SPM molecule. Moreover, the slope of the titration curve was close to $-1/4$, indicating a Ca2+ stoichiometry of 4:1 (Ca2+/SPM). In parallel, four calcium equivalents were required to saturate the signal of methylene protons of the Ca2+-EDTA complex, confirming that Ca-SPM contains four Ca2+-binding sites.

**Structural model of Ca-SPM.** The amino acid sequence of SPM has been predicted to contain three putative Ca2+-binding regions located between residues D462 and E474, D499 and D511, and D521 and E532 (2). These regions exhibit sequence homology to a 12-residue-long canonical EF-hand motif (D-X-[D/N]-X-[D/N]-G-X,-[D/E]) that binds the Ca2+ ion via side chain carboxyl (Glu or Asp) and carbonyl (Gln or Asn) groups in the conserved positions 1, 3, 5, and 12 (10). Visual inspection of the Ca-SPM structure revealed that the listed regions form very stable loops. In particular, the regions of D499 to D511 and of D521 to E533 form well-defined helix-loop-helix and helix-loop-strand

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**TABLE 1** NMR and refinement statistics for protein structures

| Parameter | Ca-SPM | Ca-SPM-P415A | Ca-SPM +4Ca2+ ions | SPM-P415A +5Ca2+ ions |
|-----------|--------|--------------|--------------------|-----------------------|
| NMR distance and dihedral constraints | | | | |
| Distance constraints (no.) | | | | |
| Total NOE | 3,412 | 2,250 | 3,412 | 2,250 |
| Intraresidue | 726 | 597 | 726 | 597 |
| Interresidue | 2,686 | 1,653 | 2,686 | 1,653 |
| Sequential (|i – j| = 1) | 853 | 552 | 853 | 552 |
| Medium-range (|i – j| < 4) | 460 | 293 | 460 | 293 |
| Long-range (|i – j| > 5) | 1,373 | 808 | 1,373 | 808 |
| Total dihedral angle restraints (no.) | | | | |
| $\Phi$ | 117 | 119 | 117 | 119 |
| $\Psi$ | 117 | 119 | 117 | 119 |
| Structure statistics | | | | |
| Violations (mean ± SD) | | | | |
| Distance constraints (Å) | 0.03 ± 0.004 | 0.03 ± 0.007 | 0.03 ± 0.005 | 0.03 ± 0.006 |
| Dihedral angle constraints (°) | 0.40 ± 0.084 | 0.45 ± 0.099 | 0.50 ± 0.067 | 0.50 ± 0.097 |
| Max dihedral angle violation (°)$^a$ | 4.30 | 5.54 | 6.36 | 7.03 |
| Max distance constraint violation (Å) | 1.31 | 1.88 | 1.19 | 0.92 |
| Deviation from idealized geometry$^b$ | | | | |
| Bond lengths (Å) | 1.14 ± 0.002 | 1.14 ± 0.002 | 1.14 ± 0.002 | 1.14 ± 0.002 |
| Bond angles (°) | 0.78 ± 0.001 | 0.79 ± 0.008 | 0.79 ± 0.007 | 0.79 ± 0.009 |
| Improvers (°) | 0.82 ± 0.03 | 0.83 ± 0.033 | 0.83 ± 0.027 | 0.83 ± 0.022 |
| Avg pairwise RMSD (Å)$^c$ | | | | |
| Heavy | 0.95 ± 0.15 | 1.27 ± 0.21 | 0.94 ± 0.14 | 1.08 ± 0.16 |
| Backbone | 0.70 ± 0.18 | 0.97 ± 0.23 | 0.70 ± 0.16 | 0.80 ± 0.18 |

$^a$Max, maximum.

$^b$Calculated by WHAT IF.

$^c$RMS Z-scores were calculated among 20 refined structures. The calculation includes the assigned residues 415 to 430 and 451 to 584.
structural motifs, respectively. Moreover, the SPM sequence contains a stretch of conserved residues (D419, D421, D423, and G424) that strongly resemble the conserved residues (positions 1, 3, 5, and 6) of an incomplete EF-hand-like motif. These residues form a stable loop in the Ca-SPM structure. Along with the adjacent D456 residue, these can provide two oxygens for calcium coordination and are likely to constitute the fourth Ca$^{2+}$/M$^+$-binding site in Ca-SPM.

In order to build a reliable model of Ca-SPM, four Ca$^{2+}$/M$^+$ ions were docked into four putative Ca$^{2+}$/M$^+$-binding sites of the Ca-SPM structure and subjected to a series of molecular dynamics (MD) simulations. Analysis of the final Ca-SPM structures after 200 ns of unrestrained MD runs revealed that all four Ca$^{2+}$/M$^+$ ions bind within the structure without triggering any significant conformational change at the level of the polypeptide backbone and without violating NMR restraints. However, calcium coordination provoked local structural changes at the level of side chains within Ca$^{2+}$/M$^+$-binding loops, where the surface-exposed Asp or Glu residues were reoriented toward the Ca$^{2+}$/M$^+$ ions. The reorientation and coordination of Ca$^{2+}$/M$^+$ by the negatively charged residues was associated with a significant shift of the electrostatic surface potential to less negative values, except for the binding site of D419 to E426 that remained negative after Ca$^{2+}$/M$^+$ binding (Fig. S3).

**FIG 2** Ca-SPM binds four Ca$^{2+}$/M$^+$ ions. (A) The $^1$H NMR spectra of free EDTA (blue) and the Ca$^{2+}$/M$^+$-chelated EDTA (red). The positions of methylene and ethylene protons of the acetyl and ethylenediamine moieties are indicated by circles and triangles, respectively. (B) Titration of Ca-SPM by EDTA. Ca-SPM was dialyzed overnight at 4°C against the Ca$^{2+}$/M$^+$-free buffer before being titrated by stepwise addition of EDTA. Relative intensities of methyl (circles) and Ca$^{2+}$/M$^+$-chelated EDTA (triangles) peaks in the $^1$H NMR spectra were plotted against the molar EDTA/SPM ratio, with the maximal intensity of peak set arbitrarily to 1. (C) Structural model of the lowest energy structure of Ca-SPM. (D) Detailed view of the Ca$^{2+}$/M$^+$-binding sites in Ca-SPM.
The final structural model of Ca-SPM is displayed in Fig. 2C. It is composed of eight anti-parallel β-strands, forming two β-sheets oriented perpendicularly to each other. The N-terminal β-strand is adjacent to the C-terminal β-strand (in antiparallel orientation), with the N and C termini of the molecule in close contact. Both β-strands form an interface between two β-sheets, constituting a hydrophobic core of the protein. This is enclosed by surface-exposed turns, a helix-turn-helix motif, and a long flexible loop at the N terminus of the protein. The structure is stabilized by binding of four Ca\(^{2+}\) ions, each coordinated by six oxygen atoms from side chains or backbone carbonyl groups of the engaged residues (Fig. 2D). Three of the four Ca\(^{2+}\)-binding sites are made up of sequential motifs (D\(_{462}\) to E\(_{474}\), D\(_{499}\) to D\(_{511}\), and D\(_{521}\) to E\(_{532}\)) that closely resemble that of EF hands, while the fourth Ca\(^{2+}\)-binding site consists of a structural motif, where the position 12 of an incomplete EF-hand-like motif (D\(_{419}\) to G\(_{424}\)) is structurally supplemented by D\(_{456}\). The calcium-binding site at D\(_{521}\) to E\(_{532}\) is adjacent to a specific pair of two tryptophan residues (W\(_{451}\) and W\(_{519}\)), whose aromatic rings are arranged in the T-shaped or edge-to-face orientation (center-to-center distance of 6.1 ± 0.3 Å; angle between the ring planes of 85 ± 5°).

Ca-SPM adopts a novel fold. Structural similarity searches using CATH (11), COFACTOR (12), DALI (13), and PDBeFold (14) revealed that the Ca-SPM structure does not resemble any of the structures deposited in the Protein Data Bank (PDB) (Fig. S4). No significant hits were found using DALI and COFACTOR (template-modeling [TM] score of <0.5 and Z score of <2). The closest structural match was identified between the Ca-SPM structure and the human split pleckstrin homology domain of phospholipase C-γ (PDB 2W2W). However, the overall homology was very low, with a Q-score of 0.12. Hierarchic classification of protein domain structures using CATH revealed that the Ca-SPM structure shows signs of similarity with a β-barrel of the lipocalin fold (CATH superfamily 2.40.128.50) of d-aminopeptidase from the Gram-negative bacterium *Ochrobactrum anthropi* and a two-layer sandwich domain (CATH superfamily 3.30.60.30) of a wild turkey serine protease inhibitor. However, the CATH analysis yielded SSAP (sequential structure alignment program) scores of 71.6 and 72.1 (with RMSDs of 5.1 Å and 9.8 Å for 82 and 51 residues, respectively), indicating that the Ca-SPM structure does not contain the same fold and only belongs to the same protein class with common structural motifs. Hence, the structure of Ca-SPM reveals a novel fold.

Backbone dynamics of Ca-SPM. In order to assess the dynamic properties of Ca-SPM on a fast (subnanosecond) timescale, we measured the steady-state \(^{1}H-^{15}N\) nuclear Overhauser enhancement and the longitudinal (\(R_1\)) and transverse (\(R_2\)) relaxation rates (Fig. S5). Based on a model-free approach, the analysis yielded the overall rotational correlation time (\(\tau_\theta\)) of 7.54 ± 0.01 ns. The generalized order parameter \(S^2\), describing the amplitude of fast (picosecond) internal motions, was higher than 0.8, indicating that the Ca-SPM backbone is rigid, similar to that of other well-folded proteins. Lower \(S^2\) values (<0.8) were observed only for six residues (G\(_{422}\), L\(_{463}\), D\(_{484}\), S\(_{500}\), V\(_{551}\), and Q\(_{563}\)), illustrating high flexibility of these residues within the Ca\(^{2+}\)-binding sites. Moreover, the NMR relaxation experiments revealed a conformational exchange on the micro- to millisecond timescale: (i) the \(R_2\) values were elevated (>15 s\(^{-1}\)) for residues V\(_{428}\) and I\(_{468}\) to A\(_{493}\), and (ii) the ratio between the intensities of the \(^{1}H-^{15}N\) cross-peaks at 30°C and 10°C (I\(_{30°C}/I_{10°C}\)) was decreased for residues in the regions of I\(_{425}\) to L\(_{456}\), I\(_{468}\) to D\(_{493}\), and Q\(_{520}\) to R\(_{534}\) against the expected I\(_{30°C}/I_{10°C}\) value. Most of these residues are in close proximity to the invisible portion of the structure, documenting that the region of T\(_{430}\) to G\(_{450}\) is in a slow conformational exchange.

Solution structure of a catalytic intermediate of SPM (SPM-P415A). To unravel the mechanistic details of the D\(_{414}\)-P\(_{415}\) cleavage, we prepared a cleavage-incompetent mutant of SPM, where the proline residue (P\(_{415}\)) of the D\(_{414}\)-P\(_{415}\) cleavage site was replaced by an alanine residue (P\(_{415}\)A). Unlike Ca-SPM, which represents a stabilized low-energy conformation of SPM after the cleavage and in which the N-terminal
sequence begins with the P415 residue (415PLALD419), the SPM-P415A construct possesses the GSDALALD419 sequence at its N terminus. This is not processed, and it is likely to represent a structure of the SPM precursor before cleavage. As shown in Fig. 3A, the overlay of the HSQC spectra demonstrated that the chemical shift patterns of the Ca2+/H11001-saturated SPM-P415A (Ca-SPM-P415A) and Ca-SPM almost overlap, indicating that the cleavage of the D414-P415 peptide bond is preceded by Ca2+/H11001-induced folding of the SPM polypeptide and that the overall structure of SPM remains unaffected after the cleavage.

Calcium titration followed by far-UV circular dichroism (CD) spectroscopy revealed that the SPM-P415A protein undergoes Ca2+-induced structural transition from an unfolded to a folded conformation that is undistinguishable from that of SPM (Fig. 3B). Moreover, Ca-SPM-P415A exhibited a strong CD exciton couplet at 230 nm, indicating that the T-shaped orientation of two tryptophan residues was maintained in the Ca-SPM-P415A structure. However, thermal unfolding experiments performed by nano-differential scanning fluorimetry (nanoDSF) revealed that Ca-SPM-P415A unfolds at a lower temperature than Ca-SPM, with the midpoint of the thermal melting curves (or melting temperature, \( T_m \)) being shifted by about 10°C (from 89°C to 79°C) (Fig. 3C). The \( T_m \) values were dependent on Ca2+ concentrations in the sample buffer, and reversible unfolding of the proteins enabled us to determine thermodynamic parameters of the folding/unfolding process. The values of Gibbs free energy of Ca-SPM and Ca-SPM-P415A unfolding, calculated for the Ca2+-saturated protein (10 mM CaCl2) at 30°C, were determined to be approximately 80 kJ/mol and 40 kJ/mol, respectively. Thus, the three extra residues at the N terminus negatively affected the thermal stability of Ca-SPM-P415A, indicating structural destabilization of the overall structure prior to cleavage.

The number of Ca2+-binding sites in Ca-SPM-P415A was determined by NMR titration experiments. Unlike results for Ca-SPM, the dialysis of Ca-SPM-P415A against the Ca2+-free buffer yielded a partial unfolding of the protein, indicating a lower conformational stability of the Ca-SPM-P415A structure. In order to avoid inaccurate

![FIG 3](https://example.com/fig3.png)
interpretation of the titration data, the dialyzed Ca-SPM-P415A sample was supplemented with an excess of two molar equivalents of Ca$^{2+}$ (to ensure folding of 100% molecules) and was next titrated with EDTA. As shown in Fig. 3D, the EDTA-induced unfolding of Ca-SPM-P415A was characterized by a nonlinear equilibrium and was completed after addition of \(-6.5\) equivalents of EDTA (including two Ca$^{2+}$ equivalents added prior to the EDTA titration). The titration data fit well to binding models describing the cooperative binding of both four and five Ca$^{2+}$ ions, but none of the models could unambiguously assign the exact number of Ca$^{2+}$-binding sites in the Ca-SPM-P415 structure. However, the overall calcium stoichiometry for Ca-SPM-P415A was higher than that for Ca-SPM (Fig. 2B), indicating the presence of an additional Ca$^{2+}$-binding site in the Ca-SPM-P415A structure.

The solution structure of Ca-SPM-P415A was determined using a procedure identical to that for Ca-SPM. A total of 2,250 distance and 119 dihedral angle restraints were used for the structure calculations (Table 1), and the overlay of final structures is depicted in Fig. 3E. Comparison of the Ca-SPM-P415A structure with that of Ca-SPM revealed that the two structures are highly similar across the models, with the average RMSD of 1.12 Å for the assigned residues. Differences in chemical shifts in most of the residues were very low (<0.1 ppm), except for residues located at the N and C termini (L416 and G578 to L582) and the loop adjacent to the N terminus (G477 to N479 and H480). Close examination of the Ca-SPM-P415A structure revealed that the N terminus of the protein adjoins two carboxy side chains of D478 and D579, which were proposed to be involved in the catalysis of the D414-P415 peptide bond cleavage by promoting protonation of P415 (1). Moreover, D478 and D579 along with D414 form a region with a negative net charge in the vicinity of the cleavage site, which is likely to form the fifth Ca$^{2+}$-binding site in the Ca-SPM-P415A structure. Therefore, the fifth Ca$^{2+}$ ion, coordinated by D414, D478, and D579, was modeled within the Ca-SPM-P415A structure.

D478 and D579 residues do not catalyze the D414-P415 peptide bond cleavage. To assess whether the carboxylate anions of D478 and D579 residues catalyze the D414-P415 peptide bond cleavage, the D478 and D579 residues were replaced by alanine or asparagine residues, and the Ca$^{2+}$-dependent self-processing activity of the mutants was examined using the respective GST-SPM fusion proteins. As shown in Fig. S6, the replacement of D478 with alanine (D478A) resulted in a significant reduction of the self-processing activity, while only a moderate reduction in the cleavage rate was observed upon replacement of D478 by asparagine (D478N). In contrast, the removal of the carboxyl group from D579 completely abolished the Ca$^{2+}$-dependent processing of the GST-SPM-D579A protein, while the D579N substitution had no significant impact on the cleavage of the GST-SPM-D579N protein. The double D478N D579N substitution reduced but did not eliminate the cleavage activity of SPM, indicating that the carboxyl groups of the D478 and D579 residues do not catalyze the cleavage of the D414-P415 peptide bond but, rather, play a key structural role.

Structural destabilization of the SPM precursor in simulations. To get insight into structure of the autoproteolytic site prior to cleavage, the A415 residue in the Ca-SPM-P415A structure was replaced in silico by a proline, and the resulting structural model of an SPM precursor (DP-SPM) was subjected to molecular dynamics (MD) simulations. The purpose of the analysis was not to determine the exact value of the $\omega_{414}$ dihedral angle (which is usually planar and fixed to 180°) but to monitor the balance between violation of an ideal peptide bond geometry (strongly imposed by the force field regardless of the actual chemical context) and the overall structure derived from the experimental data and force field requirements. The MD simulations revealed that the noncleavable D414-A415 peptide bond in the Ca-SPM-P415A structure containing four Ca$^{2+}$ ions is almost completely in trans conformation ($\omega_{414} = 180^\circ$). This is in line with the experimental data and validates our computational approach (Fig. 4). In contrast, the Ca-SPM-P415A structure containing five Ca$^{2+}$-binding sites exhibited a significant distortion of the D414-A415 peptide bond geometry, with a deviation from a near planar geometry by about 25° and 15° (depending on the NMR experimental
restraints used in the calculations). In line with that, the D414-P415 peptide bond in the DP-SPM structure containing four Ca\(^{2+}\) ions had a deviation of about 20° and even 40° in a structural model containing five Ca\(^{2+}\) ions. The cis isomer of H9275 occasionally appeared during simulations but with experimental restraints violated (Fig. 4). Taken together, these data suggested that the autocatalytic cleavage of the D414-P415 peptide bond is associated with a ground-state destabilization of the D414-P415 peptide bond through a twisted amide.

The Ca\(^{2+}\)-dependent clip-and-link activity of the ApxIVA protein plays an important role in A. pleuropneumoniae infection. Since N. meningitidis is an exclusively human pathogen, the existing animal models do not accurately simulate meningococcal disease. It has thus proved difficult to study the role of individual proteins involved in the pathogenesis of N. meningitidis in vivo. However, the Ca\(^{2+}\)-dependent clip-and-link activity has been observed also for ApxIVA (1), one of the RTX proteins of the animal pathogen A. pleuropneumoniae that causes porcine pleuropneumonia infections (15). Like FrpC, ApxIVA comprises a highly conserved SPM segment (residues 639 to 815 of the full-length ApxIVA) that mediates Ca\(^{2+}\)-dependent cleavage of the D638-P639 peptide bond and a covalent linkage of the processed N-terminal segment to adjacent proteins (1). To assess the biological importance of the Ca\(^{2+}\)-dependent clip-and-link activity of ApxIVA in vivo, we introduced an in-frame deletion of codons 629 to 827 in the apxIVA gene on A. pleuropneumoniae chromosome (Fig. 5B) and examined the virulence of the mutant strain (ΔSPM) in a pig infection model.

Intranasal administration of high doses (~10\(^9\) CFU per animal) of the wild-type and mutant strains provoked an acute form of pleuropneumonia, causing deaths of most of piglets within 24 h. The surviving animals were in the terminal stage of the disease and were euthanized at day 1 postinfection. Postmortem examination revealed that the lungs of infected animals exhibited the characteristic, dark red-to-black necro-hemorrhagic areas of consolidation accompanied by fibrinous pleuritis, with an average lung score of 28 and 32% for the wild-type and the mutant strains, respectively (Fig. 5C).
However, at lower bacterial challenge doses (~10^6 CFU per animal), moderate clinical symptoms were elicited, including lethargy, respiratory distress, and tachypnea, with no pleuropneumonia-related deaths during the time frame of the experiment (3 days). The body (rectal) temperature was largely above the physiological range (>40.5°C) and remained elevated for about 48 h. Regular hematological examinations showed neutrophilic leukocytosis, reaching the maximum at day 1 postinfection in all infected animals (Fig. 5D). However, compared to the challenge with the wild-type bacteria, no monocytosis was observed in pigs challenged with the ΔSPM mutant, and the overall monocyte counts were the same as for uninfected controls. Moreover, the average lung score for the ΔSPM-challenged pigs was lower than for pigs challenged with the wild-type strain (6.8 versus 14.7%), although the wild-type challenge strain was reisolated from lungs of only 3 out of 6 animals (compared to 5 of 6 animals for the ΔSPM strain). This result showed a significant reduction of lung lesions in pigs challenged with the ΔSPM strain, suggesting that the Ca^{2+}-dependent clip-and-link activity of ApxIVA plays a specific role in the pathogenesis of porcine pleuropneumonia.

**DISCUSSION**

Ca^{2+}-dependent protein clip-and-link activity of large RTX proteins represents a unique mechanism of posttranslational processing of proteins that involves a rearrangement of a polypeptide backbone through the highly specific Ca^{2+}-dependent autocatalytic cleavage of an Asp-Pro peptide bond and a nonspecific covalent linkage of the released carboxyl of the Asp residue to ε-amino groups of adjacent Lys residues. The cleavage of the Asp-Pro peptide bond is catalyzed by the Ca^{2+}-dependent structural transition of the SPM domain, which is highly conserved and defines a specific subclass of RTX exoproteins (1, 2). Very low Ca^{2+} concentrations in the bacterial cytosol (<100 nM) maintain the RTX proteins in an unfolded state that is required for single-step translocation across the bacterial envelope via the T1SS apparatus (16). Once exported to the calcium-rich extracellular environment (>100 μM), the proteins fold and acquire their biological activity. Given the fact that the binding affinity of SPM for Ca^{2+} is about 150 μM (2), the Ca^{2+}-dependent clip-and-link activity of the RTX proteins occurs exclusively outside the bacteria.
In this work, we determined the solution structure of SPM of the *N. meningitidis* FrpC protein and provided the structural insight into the Ca\(^{2+}\)-dependent cleavage of the D\(_{414}\)-P\(_{415}\) peptide bond. The Asp-Pro bond belongs to the most labile peptide bond, which is known to be selectively hydrolyzed within several days under acidic conditions at higher temperatures (17–20). This is due to a higher basicity of the secondary amine group in the proline than that of the primary amine group of other amino acids. At low pH, the free electron pair on the proline nitrogen may polarize the carbonyl C-N bond and make the carbonyl carbon more susceptible to nucleophilic attack. The increased basicity of the nitrogen atom may promote protonation of the nitrogen atom of the intermediate, resulting in the cleavage of the Asp-Pro peptide bond. In contrast, the SPM-mediated cleavage of the Asp-Pro peptide bond is a highly specific catalytic reaction that occurs quickly at physiological pH and with reaction rates of about 2 orders of magnitude higher than the uncatalyzed chemical cleavage. Osicka and coworkers (1) proposed that the protonation of the proline nitrogen in SPM could be promoted by an as yet uncharacterized residue(s), which can interact with the proline residue. Inspection of the Ca-SPM-P415A structure revealed two aspartate residues (D\(_{478}\) and D\(_{579}\)) positioned in close proximity to P\(_{415}\). However, site-directed mutagenesis ruled out that any of the aspartates acted as the proton-donating amino acid (see Fig. S6 in the supplemental material). Instead, both aspartates appear to play a structural role in binding of an additional (fifth) calcium ion. The fifth calcium-binding site was deduced to reside only in the SPM-P415A structure, indicating the essential structural role of the D\(_{414}\) residue prior to cleavage. Molecular docking revealed that the fifth calcium ion could be coordinated by the carbonyl oxygen of the D\(_{414}\) residue and together with D\(_{478}\) and D\(_{579}\) would constitute a transient calcium-binding site that is formed during the Ca\(^{2+}\)-dependent folding of SPM and falls apart upon the cleavage of the D\(_{414}\)-P\(_{415}\) peptide bond. The binding of the fifth calcium ion enhances distortion of the C-N bond of the carboxyamide group, generating a twisted amide in a scissile bond, which destabilizes the overall structure of SPM. This is well documented by the experimental data showing that the thermal stability of the Ca-SPM-P415A structure before cleavage is much lower than that of the Ca-SPM structure after the cleavage.

Twisted amides represent one of the approaches for activation of amide bonds, which are usually stable due to formation of a resonating structure provided by the conjugation of the nitrogen lone pair with the carbonyl group (21, 22). The distortion of amide bonds causes a loss of double-bond character and thus increases their reactivity compared to planar counterparts. Such an unusual conformation of the scissile amide bond is the driving force for a nucleophilic or electrophilic attack that is proposed to be a central design element of a variety of enzymatic processes, such as cis-trans isomerization (23), amide hydrolysis (24), N-linked glycosylation of proteins (25), and intein-based protein splicing (26). Based on the data above, we propose a mechanism for the SPM-mediated processing of the Asp-Pro peptide bond (Fig. 6). Upon calcium binding, the SPM polypeptide undergoes a highly cooperative structural transition from an intrinsically unstructured conformation to the compact protein fold that is stabilized by binding of four calcium ions in EF-hand-like Ca\(^{2+}\)-binding sites. The scissile D\(_{414}\)-P\(_{415}\) peptide bond adopts a highly strained turn that is structurally constrained by the adjacent \(\beta\) strand and the residues of a narrow port (W\(_{451}\), L\(_{475}\), G\(_{477}\), G\(_{491}\), and D\(_{579}\) to L\(_{582}\)) that shields the D\(_{414}\)-P\(_{415}\) cleavage site from the solvent (Fig. 6A and B). Coordination of the fifth calcium ion by the oxygen atom of the carbonyl group of D\(_{414}\) provokes rotation of the D\(_{414}\)-P\(_{415}\) peptide bond over the C-terminal \(\beta\) strand that apparently accentuates rotation of the C-N bond to produce a twisted amide. Such a distortion generates a reactive lone electron pair at the nitrogen that abolishes the conjugation of the nitrogen electrons with the carbonyl group and facilitates a nucleophilic attack of the carboxylic acid side chain of D\(_{414}\) on its carbonyl carbon (Fig. 6C). This results in formation of a cyclic imide intermediate and D\(_{414}\)-P\(_{415}\) peptide bond cleavage, with subsequent formation of a reactive cyclic D\(_{414}\) anhydride. This can next be attacked by a free amino group of a lysine residue to generate an isopeptide bond.
The SPM represents a highly conserved domain that appears to be present in many RTX proteins of Gram-negative bacteria. However, the SPM-mediated processing of the Asp-Pro peptide bond has as yet been reported only for the \textit{N. meningitidis} FrpC and \textit{A. pleuropneumoniae} ApxIVA proteins \textit{in vitro} \cite{1}. Sviridova et al. \cite{6} demonstrated that incubation of meningococci with human alveolar A549 cells resulted in the processing of the D414-P415 peptide bond of the freshly secreted FrpC and a covalent linkage of the FrpC1–414 segment to cell surface proteins of the epithelial cells. In view of the fact that FrpC1–414 exhibits a very high affinity (dissociation constant \(K_d\) of \(\approx 0.2\) nM) for FrpD, the meningococcal outer membrane lipoprotein encoded in the same operon upstream of the \textit{frpC} gene \cite{27}, the interaction of FrpC1–414 with FrpD could be involved in tight adherence of \textit{N. meningitidis} to the host cell surface. Nevertheless, deletion of the \textit{frpC} gene from the genome of \textit{N. meningitidis} MCS8 did not affect the mortality rate of rats after a systemic meningococcal infection, indicating that FrpC is dispensable for virulence in the infant rat model \cite{28}. However, this does not preclude the role of the SPM-mediated processing of FrpC in the attachment of meningococci to mucosal surfaces that are the colonization niche of \textit{N. meningitidis}.

Our \textit{in vivo} pig experiments showed that the abrogation of the Ca\(^{2+}\)-dependent clip-and-link activity of ApxIVA led to significant reduction of necrotizing lung lesions after \textit{A. pleuropneumoniae} infection (Fig. 5C). These data are in good agreement with...
the previous result showing that the deletion of the apxIVA gene attenuates the virulence of *A. pleuropneumoniae* in a pig infection model (29). *A. pleuropneumoniae* produces four different RTX proteins, where only Apxl, ApxlII, and ApxlIII are toxins possessing both hemolytic and cytotoxic activity (30). With respect to the fact that the fourth RTX protein, ApxIVA, does not exert any of these activities, the biological activity of ApxIVA could be ascribed to the Ca$^{2+}$-dependent clip-and-link activity. It is tempting to speculate that colonization of pig respiratory airways by *A. pleuropneumoniae* is associated with the Ca$^{2+}$-dependent processing of the D$_{638}$P$_{639}$ peptide bond of the type I-secreted ApxIVA and covalent linkage of the ApxIVA$_{630-638}$ fragment to the host respiratory epithelium, which would serve as a high-affinity target for an as yet unidentified protein structure exposed on the bacterial cell surface. This would allow tight adherence of the bacterial cells to the host respiratory epithelia, which may be essential for the full virulence of *A. pleuropneumoniae*. Hence, covalent attachment of the N-terminal fragments of specific RTX proteins to the host proteins through the Ca$^{2+}$-dependent protein clip-and-link activity could represent an unconventional strategy for pathogenic microorganisms to adhere to the target host cell surface.

**MATERIALS AND METHODS**

**Reagents.** The suicide vector for allelic exchange on the *Actinobacillus pleuropneumoniae* chromosome (pEMOC2) and the *Escherichia coli* β2155 (ΔdapA) strain were generously provided by Gerald F. Gerlach (Institute for Innovative Veterinary Diagnostics, Hannover, Germany).

**Plasmid construction.** The wild-type SPM protein was expressed from the pET28b-dHis-GST-SPM plasmid as a C-terminal glutathione S-transferase (GST) fusion protein as previously described (31). The SPM-P415A protein was expressed from the pET42b-SPM-P415A plasmid generated by PCR amplification of the nucleotide sequence of SPM from the pET28b-dHis-SPM construct using a pair of forward and reverse primers (5′-TTTACATATGAAACTGCTACTTACAAGGCACCGCTCCGCTAGCCC-3′ and 5′-TTCCTCCATG GTACTCGAGGAAAGGCT-3′) and ligation of the SpeI/Ncol-restricted PCR fragment into the SpeI/Ncol-cleaved pET42b vector. The pET28b-dHis-SPM$_{630-638}$ construct was prepared by PCR mutagenesis using a pair of primers (5′-GGGTCTGGGATCTGACTTCT-3′ and 5′-AAACGGCTAACTTTGATCGTGACGCGC-3′) and cloning of the BamH/Algl-restricted PCR fragment into the BamH/Algl-cleaved pET28b-dHis-SPM plasmid. The pET28b-dHis-SPM-D579A construct was prepared by PCR mutagenesis, whereby two PCR fragments each carrying the linking ApaI restriction site were amplified by PCR using two pairs of PCR primers (5′-GCCACCGGTTGGTATTG-3′ and 5′-TTTGGGCCCATTATTTGCGGTTG-3′). The pET28b-dHis-SPM-D579A and 5′-GGCCACACTAGTGAACC-3′, cleaved with the respective pair of restriction enzymes (AlpI/Aeil and ApaI/DraiI), and collectively ligated into the ApaI/DraiI-cleaved pET28b-dHis-SPM plasmid. The pET28b-dHis-SPM-D579N construct was prepared using the same conditions except that the PCR primer pair 5′-GCCACCGGTTGGTATTG-3′ and 5′-TTTAA ATTCCTCATATGCCGCC-3′ and the pair 5′-TTTGAATTCTTATTTAGCAACGC-3′ and 5′-GGCC ACTACGTGAACC-3′ and the Apol restriction site were used. The PCR products were purified by on-column overlap PCR mutagenesis using two pairs of PCR primers (5′-GAC AACGCTCATATGCCGCC-3′ and 5′-CTGCCAGTTTTGATCTGCGCCGAAGTGTCGCGCCGC-3′, 5′-GGCGGAACTCGCCTGCCGAATACCAACCTGACGCGCGCG-3′ and 5′-GGCCACACTAGTGAACC-3′), whereby the resulting HindIII DRall-restricted PCR product was cloned into the HindIII/DRall-cleaved pET28b-dHis-SPM-GST plasmid. The pET28b-dHis-SPM-D478A plasmid was constructed under the same conditions except that the PCR primer pair 5′-GACAAGCTCTATATGCCGCC-3′ and 5′-CTGCCAGTTTTGATCTGCGCCGAAGTGTCGCGCCGC-3′ and the pair 5′-GCCGCGAATCTTCGCGCACAATCCAAACTGGCGCAG-3′ and 5′-GGCCACACTAGTGAACC-3′ were used. The double-substitution construct pET28b-dHis-SPM-D478H+D579N was prepared by replacement of the ApaI/DraiI DNA fragment of the pET28b-dHis-SPM-D478H by that of the pET28b-dHis-SPM-D579N construct.

For construction of the pEMOC2-ApxIVAΔ629-827 allelic exchange vector, two DNA fragments corresponding to the 5′ and the 3′ flanking regions of the in-frame deletion were amplified from genomic DNA purified from the Czech field isolate Actinobacillus pleuropneumoniae KL2-2000 (biotype 1, serotype 9) using PCR, as follows: the 998-bp DNA fragment upstream of the deletion was amplified from DNA by using the forward primer 5′-TTTGGCAGCCGCTTGGGGAAGAAGTGTAAGC-3′ containing the NotI restriction site and the reverse primer 5′-TTTCTCGCAGATTTCGCCGAATTCCATCG-3′ containing the Xhol site. Similarly, the 1,004-bp DNA fragment downstream of the deletion was amplified from DNA by using the forward primer 5′-TTTGGCAGCCGCTTGGGGAAGAAGTGTAAGC-3′ containing the Xhol site and reverse primer 5′-TTTGGCCCCATTAAAATGATGCGTCAATCGCCG-3′ containing the Apol site. The PCR products were cut with appropriate restriction enzymes and collectively ligated with the NotI/Apol-cleaved pEMOC2 vector. All constructs were confirmed by DNA sequence analysis with an ABI Prism 3130XL analyzer (Applied Biosystems, USA) using a BigDye Terminator cycle sequencing kit.

**Protein expression and purification.** All GST-SPM fusion constructs were expressed in *E. coli* strain BL21 (DE3) transformed with the appropriate plasmid. Exponential 20-ml cultures were grown in a shaking incubator at 37°C in M9 minimal medium supplemented with trace metals, vitamins, and kanamycin (60 μg/ml). For NMR experiments, the cells were grown in M9 medium supplemented with 0.5 g/liter [13C]glucose (Cambridge Isotope Laboratories, USA) and 2 g/liter [15N]glucose (Cambridge Isotope Laboratories, USA).
Expression of proteins was induced by adding 1 mM isopropyl-β-D-thiogalactopyranoside (IPTG) at an optical density at 600 nm (OD600) of 0.6 to 0.8, and bacteria were grown for an additional 4 h. The cells were harvested by centrifugation (1,500 × g for 15 min), washed in TNE buffer (50 mM Tris-HCl, pH 7.4, 150 mM NaCl, 5 mM EDTA), and disintegrated by sonication in TN buffer (50 mM Tris-HCl, pH 7.4, 150 mM NaCl) at 4°C. The cell extracts were centrifuged at 20,000 × g for 30 min, and the supernatants were used for protein purifications.

The purification of the wild-type Ca-SPM was previously described (31). Briefly, the cleared cell lysate was loaded onto Ni-Sepharose 6 Fast Flow beads (GE Healthcare) and washed with TN buffer, and the dHis-GST-SPM protein was eluted with TN buffer supplemented with 500 mM imidazole. The collected fractions were mixed with dithiorthiolether (DTT) to a final concentration of 10 mM before the protein solution was dialyzed overnight at 4°C in TN buffer supplemented with 10 mM DTT and 10 mM CaCl2. Addition of calcium ions induced a self-processing activity of SPM, resulting in the cleavage of the GST-SPM fusion proteins. The cleaved GST protein was precipitated by incubation of the protein mixture at 70°C for 15 min, and the resulting suspension was centrifuged at 5,000 × g for 20 min. The supernatant was loaded onto a PLRP-S reverse-phase column (Agilent Technologies) in buffer containing 50 mM triethylammonium (pH 8.5) and 5% acetonitrile, and SPM was eluted from the column by the gradient of acetonitrile (5 to 95%). The SPM fractions were concentrated by rotary vacuum evaporator and loaded onto a Superdex HR 200 gel filtration column (GE Healthcare) equilibrated with TN buffer supplemented with 10 mM CaCl2. For purification of SPM-P415A, the cell lysate containing GST-SPM-P415A was loaded onto a glutathione agarose column (Life Technologies), extensively washed with TN buffer, and eluted with TN buffer supplemented with 10 mM i-glutathione (reduced). Collected fractions were pooled, mixed with the purified recombinant tobacco etch virus (TEV) protease (1:20, wt/wt), and dialyzed at 4°C overnight against TN buffer. The protein mixture was incubated at 70°C for 15 min, and the resulting suspension was centrifuged at 5,000 × g for 20 min. The supernatant was loaded onto a PLRP-S reverse-phase column (Agilent Technologies) in buffer containing 50 mM triethylammonium (pH 8.5) and 5% acetonitrile, and SPM-P415 was eluted from the column by the gradient of acetonitrile (5% to 95%). The collected fractions were concentrated by rotary vacuum evaporator and loaded onto a Superdex HR 200 gel filtration column (GE Healthcare) equilibrated with TN buffer supplemented with 10 mM CaCl2. The single (D478A, D478N, D579A, and D579N) and double (D478N D579N) substitution mutants of GST-SPM constructs were purified by Ni-Sepharose 6 Fast Flow column chromatography as described above for the wild-type SPM. The collected fractions were mixed with dithiorthiolether (DTT) to a final concentration of 10 mM before overnight dialysis at 4°C in TN buffer. The purity of the proteins was monitored by SDS-polyacrylamide gel electrophoresis (SDS-PAGE), and protein concentrations were determined by the Bradford assay (Bio-Rad) using bovine serum albumin as a standard.

**NMR spectroscopy.** NMR spectra were acquired at 30°C on Bruker Avance III 600 MHz spectrometer equipped with triple-resonance (1H-13C-15N) inverse and quadruple-resonance (1H-13C-15N) inverse cryoprobes, a Bruker Avance III HD 700 MHz NMR spectrometer equipped with a triple-resonance (1H-13C-15N) inverse cryoprobe optimized for 15N detection, a Bruker Avance III HD 950 MHz NMR spectrometer equipped with a triple-resonance (1H-15N) inverse cryoprobe, and a Bruker Avance III HD 950 MHz NMR spectrometer equipped with a triple-resonance (1H-15N) inverse cryoprobe. The uniformly 15N-labeled samples were measured in buffer containing 10 mM Tris-HCl (pH 7.4), 50 mM NaCl, 10 mM CaCl2, 10% D2O, and 0.1% Na2Mo4 at protein concentrations of 0.35 mM and 0.35 mM for Ca-SPM and Ca-SPM-P415A samples, respectively. Two-dimensional (2D) 1H-13C HSQC, three-dimensional (3D) HNCO, 3D CBCA(CO)NH, and 3D H/C/CH total correlation spectroscopy (TOCSY) spectra measured at 600 MHz and 3D aliphatic 1H-13C-edited NOESY spectra measured at 950 MHz were obtained for the SPM-P415A sample. 3D HNCO, 3D HNCACB, 3D CBCA(CO)NH, 3D 13C-NOEDY spectra, 3D H/C/CH-TOCSY, and 3D aromatic 1H-13C-edited NOESY spectra measured at 600 MHz, 2D aliphatic 1H-13C HSQC spectrum measured at 700 MHz, 2D 2H-15N HSQC, 13N R ratios, 13N R N, steady-state 1H-15N NOE enhancement (ssNOE) spectra measured at 850 MHz, and 3D aliphatic 1H-13C-edited NOESY spectrum measured at 950 MHz were obtained for the SPM sample. In addition, 2D 1H-13N HSQC, 3D HN(CO)CA, and 3D HNCA spectra of Ca-SPM were recorded at 10°C at 850 MHz, and 3D HN(CO)CA and 3D CBCA(CO)NH were measured at 10°C at 950 MHz. Sequential assignment of backbone and side chain 1H, 13C, and 15N resonances of SPM was performed by standard triple-resonance NMR techniques (32). All NMR spectra were processed and analyzed with NMRPipe, version 8.7 (33), and Sparky, version 3.115 (T.D. Goddard and D. G. Kneller, University of California, San Francisco, USA). The automated assignment of NOESY spectra and the structure calculation were performed in CYANA, version 3.97 (34).

**Structure calculation of SPM and SPM-P415A.** Torsion angle restraints were determined from chemical shifts by TALOS-N (35) using the 1HN, 1H, 15N, 13C, 13C, and 15C resonances. Distance restraints were derived from the automatically assigned NOESY spectra with a manual check of the results. Distance and torsion restraints were used to generate 100 structures in CYANA, version 3.97. Twenty structures with the lowest target functions were used as an input for the molecular dynamics (MD) simulations. In calculations with calcium ions, four Ca2+ ions were introduced in CYANA, version 3.97, by defining three 3-Å distance restraints between Ca2+ and oxygen OD1 of D419, D421, D423, D462, N464, N468, D499, N501, D503, D521, N523, and D525. The fifth Ca2+ ion was introduced into the constructs containing D419, where the distance restraints were to oxygen OD1 of D414, D478, and D579. The Ca2+ ions were not restrained in the final refinement during the MD runs.

**MD simulations.** Molecular dynamics (MD) simulations were performed in GROMACS, version 5.0.5 (36, 37) using the force field parameters AMBER99SB-ILDN (38). On average, the systems were solvated in 8,200 explicit TIP3P (39) water molecules and 12 Cl− and 18 (minus two times number of Ca2+ ions introduced) Na+ ions in a periodic dodecahedron simulation box mimicking the electroneutral system.
with a salt concentration of 70 mM. The MD step of 1 fs was used in the leapfrog integration scheme. Van der Waals interactions were calculated using a triple-range cutoff scheme, with the updated interaction distance of 1 nm. Electrostatic interactions were calculated by the particle mesh Ewald (PME) method with a Coulomb cutoff of 1 nm and a relative dielectric constant of 1. The temperature and pressure were controlled by a Berendsen thermostat (40) with a coupling constant of 0.1 ps and a Parrinello-Rahman barostat (41) at the constant pressure of 10^6 Pa with the coupling constant of 1 ps. Simulated annealing of the system was initiated with a relaxation phase of 100 ps, equilibration at 300 K, and subsequent heating to 1,000 K in 50 ps. The temperature was kept constant for 3 ns and then slowly decreased to 0 K over an additional 12 ns. The structure was restrained during MD by experimental distances and dihedral angles with force constants of 1,000 kJ mol^-1 nm^-2 and 350 kJ mol^-1 rad^-2, respectively. The quality of the structures was analyzed by PROCHECK (42), WHAT IF (43), and CING (44), available as the online service CING. Ramachandran statistics by program PROCHECK categorized 91.3 and 92.4% residues in the most favored regions, 8.6 and 7.4% in the additional allowed regions, and 0.1 and 0.1% in the disallowed regions. Secondary structure motifs were identified from the models by the program DSSP (45) with a threshold of 75% of occurrence. The electrostatic potentials were calculated at 303.2 K in APBS (46). The chemical shifts were predicted from calculated structures by SPARTA+ (47).

**Titration of Ca-SPM and Ca-SPM-P415A by EDTA.** An unlabeled SPM or SPM-P415A sample was prepared in the same buffer as used for the structure determination and was dialyzed overnight against a CaCl2-free buffer, consisting of 50 mM Tris-HCl (pH 7.4) and 50 mM NaCl; 10% D2O and 0.1% NaN3 were added to the sample prior to the measurement. The final volume and protein concentration were 0.54 ml and 0.175 mM or 0.52 ml and 0.707 mM for SPM or SPM-P415A, respectively. The titration was monitored by a 1D proton NMR experiment with water suppression using the WATERGATE W5 pulse sequence with gradients in a double-echo at 700 MHz, and the delays for water suppression were 205 or 169 ms. The total number of scans was 192 or 80, and the spectral width was 20 or 30 ppm for SPM or SPM-P415A, respectively. The delay for the interscan was 5 s, and spectra were sampled by 64,000 points. The samples were titrated with the solution of 50 mM EDTA, 50 mM NaCl, and 50 mM Tris-HCl (pH 7.4). The 1D spectra were processed with apodization exponential 1Hz function and phased.

**15N relaxation experiments.** The NMR 15N relaxation experiments were carried out at 30°C using 15N heteronuclear single-quantum coherence (HSQC)-based pulse schemes on a uniformly 15N-labeled Ca-SPM (0.33 mM). The temperature was calibrated with a 100% methanol sample, where difference between peaks was set up to 1.518 ppm (48). The following relaxation delays were used in the 15N R1 experiment: 11.2, 78.4, 168.0, 268.8, 380.8, 492.8*, 638.4, 795.2, 985.6, and 1,232.0 ms; the relaxation delays in the 15N R2 experiments: 0, 14.4, 28.8, 43.2, 57.6, 72.0°, 86.4, 100.8, and 115.2 ms. Asterisks denote spectra recorded twice in order to estimate experimental error. Each 2D 15N R1 and 15N R2 experiment was composed of 320 by 2,048 complex points in the indirect 15N and direct 1H dimensions, respectively, corresponding to respective acquisition times of 61.5 ms and 80.2 ms. Experiments were acquired with 8 scans per free induction decay (FID) and an interscan delay of 2 s. The relaxation rates were obtained by fitting peak intensities to a mono-exponential decay by using Relax (49). The 1H-15N ssNOE measurements were achieved with a uniformly 15N-labeled Ca-SPM (0.6 mM) under a steady-state condition, achieved by a 5-ms 1H irradiation with 225 repeats of 200-µs pulses separated by 22.2-µs delays and with a 15-s interscan delay. The recorded spectra consisted of 320 by 2,048 complex points in the indirect 15N and direct 1H dimensions, respectively, corresponding to respective acquisition times of 57.7 ms and 80.3 ms for a reference and steady-state spectra, which were measured in an identical manner. Experiments were acquired with 16 scans per free induction decay (50). Fast backbone motions of Ca-SPM were calculated by a model-free approach (51). The program ROTDIF (52) was used to calculate a rotational diffusion tensor in order to perform the influence of the slow exchange from the effect of the rotational diffusion. The axially symmetric model was used for description of the experimental data.

**CD spectroscopy.** The far-UV circular dichroism (CD) spectra were recorded on a Jasco-815 spectropolarimeter in rectangular quartz Suprasil cells of 1-mm path length (110-QS; Hellma). Protein samples (100 µg/ml) were diluted in 5 mM Tris-HCl (pH 7.4) plus 50 mM NaCl in the absence or presence of 10 mM CaCl2, and measured for wavelengths from 195 to 280 nm at 25°C. Two spectrum accumulations with standard instrument sensitivity and scanning speed of 10 nm/min with response time of 16 ms were acquired. The spectra of the buffers were subtracted from the protein spectra, and mean molar ellipticity ([θ]) was expressed in degrees per square centimeter per decimole.

**Thermal stability assay.** Thermal stability of SPM proteins was performed by nano-differential scanning fluorimetry (nanoDSF) using a Prometheus NT.48 instrument (NanoTemper Technologies, Munich, Germany). Ca-SPM and Ca-SPM-P415A were dialyzed overnight at 4°C against TN buffer, and the freshly prepared protein samples were supplemented with increasing concentrations of calcium ions (0 to 10 mM CaCl2) before 10 µl was used for filling the Prometheus NT.48 high-sensitivity capillaries (NanoTemper Technologies). The measurements were conducted from 20 to 95°C (with a temperature ramp of 2.5°C/min) under constant monitoring of tryptophan fluorescence at 350 and 330 nm. The melting temperature (Tm) values, corresponding to the inflection points of the unfolding curve, were determined via the first derivative of the curve.

**Construction of the apxIVA mutant strain of A. pleuropneumoniae.** In-frame deletion in the apxIVA gene (codons 629 to 827) was performed on chromosome of the Czech field isolate Actinobacillus pleuropneumoniae KL2-2000 (biotype 1, serotype 9) by homologous recombination using a PEMOC2 allelic exchange vector as previously described (53). Briefly, A. pleuropneumoniae cells grown in a 5% CO2 atmosphere at 37°C on Bacto tryptic soy broth agar (BD Biosciences) supplemented with 10 µg/ml
β-NAD (NAD; Sigma) were mated on fresh agar plates supplemented with 10 μg/ml NAD, 1 mM diaminopimelic acid (DAPA; Sigma), and 10 mM MgSO₄ with E. coli pJ2155 grown on LB agar supplemented with 1 mM DAPA and 25 μg/ml chloramphenicol (Cm) before being transformed with the pEMOC2-ApxIVAΔ629-827 plasmid construct. After 4 h at 37°C, the A. pleuropneumoniae transconjugants were selected on Bacto tryptic soy broth agar plates supplemented with 10 μg/ml NAD and 5 μg/ml chloramphenicol and incubated in 5% CO₂ for 24 h at 37°C. For sucrose counterselection, the single Cm⁺ colonies were inoculated into salt-free LB broth supplemented with 10% sucrose, 10% horse serum, and 10 μg/ml NAD, incubated with shaking at 37°C for 2 h, and plated on salt-free LB broth supplemented with 10% sucrose, 10% horse serum, and 10 μg/ml NAD. Sucrose-resistant colonies were screened for the truncated variant of the apxIVA gene by a colony PCR and restriction analysis of the resulting PCR products (XhoI).

Animal experiments. The animal experiments were carried out according to the guidelines of the Animal Care Act (no. 246/1992 Coll.) of the Czech Republic, approved by the Animal Welfare Commission of the Ministry of Agriculture of the Czech Republic, and conducted in accredited barrier-type stables (accreditation certificate no. 5843/2007-10001). Thirty 4-week-old Large White pigs with body weights of 7 to 10 kg were purchased from a porcine reproductive and respiratory syndrome virus-free and A. pleuropneumoniae infection-free herd (as tested by serological reaction in enzyme-linked immunosorbent assays [ELISAs]) and randomly assigned to five groups, each containing six animals. Pigs were regularly monitored throughout the housing period at least three times per day.

Experimental infection. The virulence of A. pleuropneumoniae strains was assessed in an intranasal infection model as described previously (54). Briefly, pigs were anesthetized by intravenous injection of ketamine (4 mg/kg of body weight) and xylazine (2 mg/kg) and intranasally administered 4 ml (2 ml per nostril) of phosphate-buffered saline (PBS) solution (control group) or infected with two doses of 2 ml bacterial suspension corresponding to an infection dose of 10⁶ and 10⁹ CFU per animal. The CFU number was measured spectrophotometrically in the bacterial culture grown at 37°C in brain heart infusion broth (HiMedia, India) supplemented with 10 μg/ml NAD, in which an optical density at 550 nm (OD₅₅₀) of 1.0 corresponded to 5 × 10⁶ CFU/ml. Pigs were monitored for 3 days after infection, and the A. pleuropneumoniae-related clinical signs (respiratory rate, dyspnea, coughing, anorexia, and lethargy) were recorded. At −24, 0, 24, 48, and 72 h postinfection, body temperature was measured in the rectum. Blood samples were collected by jugular venipuncture into heparinized tubes (30 IU/ml) at −24, 24, and 72 h postinfection and immediately used for cytological examination. Cell counts were determined by a hemocytometer for macroscopic pathological changes and for the presence of bacteria. The lung score was determined as the extent of lung tissue damage (lung lesions) according to the percentages for the following anatomical segments: left apical lobe, 8%; right cranial lobe, 12%; left cardiac lobe, 8%; right medium lobe, 12%; left diaphragmatic (caudal) lobe, 25%; right diaphragmatic (caudal) lobe, 30%; and accessorius lobe, 5%. For bacterial examination, nasal and tracheal swabs were inoculated on Columbia agar supplemented with 5% sheep blood with a streak of S. aureus (as a source of NAD) and cultivated aerobically at 36°C for 18 h.

Data availability. NMR chemical shifts and distance restraints have been deposited in the Biological Magnetic Resonance Bank (BMRB) under entries 28058 (http://www.bmrb.wisc.edu/data_library/summary/index.php?bmrbld=28058) and 34424 (http://www.bmrb.wisc.edu/data_library/summary/index.php?bmrbld=34424) for Ca-SPM and Ca-SPM-P415A, respectively. Structural coordinates of Ca-SPM and Ca-SPM-P415A have been deposited in the Protein Data Bank under accession codes PDB 6JW (https://www.rcsb.org/structure/6jw) and PDB 6JSX (https://www.rcsb.org/structure/6jsx), respectively. All other data are available from the authors upon request.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

FIG S1, TIF file, 0.6 MB.

FIG S2, TIF file, 1 MB.

FIG S3, TIF file, 1.3 MB.

FIG S4, TIF file, 2.4 MB.

FIG S5, TIF file, 0.9 MB.

FIG S6, TIF file, 1.6 MB.

TABLE S1, DOCX file, 0.01 MB.

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Ladislav Bumba prepared the proteins; Vojtech Kuban, Pavel Macek, and Lukas Zidek performed the NMR experiments; Vojtech Kuban calculated the NMR structures; Vojtech Kuban and Jozef Hritz ran the molecular dynamics simulations; Katerina Nechvatalova, Katerina Nedbalcova, and Martin Faldyna performed and analyzed the animal experiments; Ladislav Bumba, Peter Sebo, and Lukas Zidek designed the experiments. Peter Sebo edited and Vojtech Kuban, Lukas Zidek, and Ladislav Bumba wrote the manuscript.

We declare that we have no competing interests.

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