Functional Insight into the C-Terminal Extension of Halolysin SptA from Haloarchaeon Natrinema sp. J7

Zhisheng Xu, Xin Du, Tingting Li, Fei Gan, Bing Tang, Xiao-Feng Tang*

State Key Laboratory of Virology, College of Life Sciences, Wuhan University, Wuhan, China

Abstract

Halolysin SptA from haloarchaeon Natrinema sp. J7 consists of a subtilisin-like catalytic domain and a C-terminal extension (CTE) containing two cysteine residues. In this report, we have investigated the function of the CTE using recombinant enzymes expressed in Halofex volcanii WFD11. Deletion of the CTE greatly reduced but did not abolish protease activity, which suggests that the CTE is not essential for enzyme folding. Mutational analysis suggests that residues Cys303 and Cys338 within the CTE form a disulfide bond that make this domain resistant to autocleavage and proteolysis under hypotonic conditions. Characterization of full-length and CTE-truncation enzymes indicates the CTE not only confers extra stability to the enzyme but also assists enzyme activity on protein substrates by facilitating binding at high salinities. Interestingly, homology modeling of the CTE yields a β-jelly roll-like structure similar to those seen in Claudin-binding domain of Clostridium perfringens enterotoxin (clostridial C-CPE) and collagen binding domain (CBD), and the CTE also possesses collagen-binding activity, making it a potential candidate as an anchoring unit in drug delivery systems.

Introduction

Haloarchaea thrive in hypersaline environments and generally require 15-30% NaCl as an optimal condition for their growth. Many of them secrete proteases to degrade protein substrates into small peptides and amino acids which the organisms then metabolize [1]. Halophilic enzymes, including proteases from haloarchaea, are active and stable at high salt concentrations, representing an attractive model for investigating enzymatic adaptation mechanisms at the molecular level [2]. Additionally, halophilic proteases show great potential as biocatalysts in the synthesis of oligopeptides in low water conditions (66%), including organic mixtures [3].

Many extracellular proteases have been isolated from haloarchaea, and almost all of them are serine proteases [1]. So far, the genes encoding the extracellular proteases 172P1 from Natrathia asatica [4], R4 from Halofex mediterranei [5], SptA from Natrinema sp. J7 [6] and Nep from Natrathia magadai [7] have been identified. Analysis of deduced amino acid sequences of these four enzymes indicate that they are members of the thermolysin family of subtilisin-like serine proteases [subtilases] [8], and have been denoted as “halolysins” [1,4]. To fulfill their physiological functions, halolysins should not only be stable in hypersaline environments, but also be efficiently active in spite of substrates undergoing salt-induced conformational change and precipitation. Exploring the molecular basis of halolysin activity at high salinities will provide new insights into the adaptation mechanisms of haloarchaea.

One common feature shared by halolysins is that the mature enzyme is composed of a subtilisin-like catalytic domain, and a C-terminal extension (CTE) with approximately 120 amino acid residues. CTEs have also been found in other subtilases, such as alkaline serine protease KP-43 from Bacillus sp. [9], hyperthermostable subtilisin Tk-SP from Thermococcus kodakaraensis [10], tomato subtilase 3 [11], kexin-like protease from Aeromonas sobria [12], yeast Kex2 [13] and mammalian furin [14]. The CTEs of these enzymes are proposed to be involved in enzyme stability and substrate recognition or binding. In addition, the precursor of aqualysin from Thermus aquaticus YT-1 has a C-terminal propeptide, which is required for enzyme secretion but is later processed to form the mature enzyme [15]. In haloarchaea, the CTE of halolysin R4 has been proposed to be functionally important, possibly by providing structural stability. However, halolysin R4 has proven difficult to purify and biochemically characterize in detail [5] and the function of haloarchaeal CTE domains is yet to be elucidated.

Halolysin SptA (formerly SptA protease) is extracellularly produced by Natrinema sp. J7 [6]. The SptA precursor is composed of a 49 residue signal peptide, a 103 residue propeptide, and a 413 residue mature region containing a CTE. When expressed in Halofex volcanii WFD11, the recombinant SptA was secreted into the medium as an active mature form, and was stable enough to purify for further investigation [6]. In addition, mature SptA has four cysteine residues, two of which are conserved among halolysins and contained in the CTE. In this report, we perform deletion and point mutation analyses of the halolysin SptA C-terminal extension and discuss the possible roles of the CTE and its cysteine residues in enzyme stability, activity and substrate binding.
Results

Expression of C-terminal truncation mutants in *Hfx. volcanii*

In order to probe the function of the CTE, we first constructed two truncation mutants, SptAΔC125 and SptAΔC76, by deleting 125 and 76 residues from the C-terminus of the enzyme, respectively (Fig. 1). As shown in Fig. 2A, hydrolysis halos could be detected around the colonies of *Hfx. volcanii* harboring the expression plasmid for SptA or SptAΔC76 growing on skim milk plates, but was hardly detected for SptAΔC125. This served as a means of detecting successful transformants as well as a basic indicator of activity. For a more robust activity assay, we used suc-AAPF-pNA as the substrate in reactions with culture supernatant containing secreted SptA, SptAΔC76, or SptAΔC125. Substantial levels of protease activity were found in culture supernatants individually expressing SptA and SptAΔC76, and trace activity was detected in that expressing SptAΔC125. In contrast, extracellular protease activity could not be detected in the control strain harboring plasmid pSY1, either by milk plate assay or enzyme activity measurement under the same conditions (Fig. 2A).

In order to identify the fractions containing the pro- and mature forms of the enzymes, we conducted immunoblot analyses. It should be mentioned that these recombinant enzymes migrate (Fig. 2B) higher than their theoretical molecular masses because they are acidic and resistant to SDS denaturation compositions, a common feature of halophilic enzymes [2]. Pro-forms of SptA, SptAΔC76 and SptAΔC125 were detected in the insoluble fractions but not in the soluble fractions of the cells (Fig. 2B). Mature forms of SptA and SptAΔC76 were found in culture supernatants, and mature SptAΔC125 could hardly be detected (Fig. 2B). This is consistent with our assay results that showed the activity of the culture supernatant containing SptAΔC125 was only approximately 1/1400 and approximately 1/500 of that containing SptA and SptAΔC76 (Fig. 2A), respectively. The appearance of other bands might result from the cross reaction between the antibodies and host proteins or/and degraded products of recombinant enzymes. Taken together, these results demonstrate that the CTE is important but not required for correct folding and activity of SptA.

The protective effects of disulfide bond Cys303-Cys338

Our next question regarding the function of the CTE focused on intramolecular disulfide bond formation, which is known to protect against autolysis in subtilisins [16] and is postulated for halolysin R4 [5]. The mature SptA contains four cysteine residues at positions 103, 154, 303 and 338. Of these, Cys303 and Cys338 are located within the CTE (Fig. 1). In order to identify the

---

**Figure 1.** Schematic representation of the primary structures of SptA precursor and its derivatives (A) and amino acid sequence alignment of the CTEs of halolysins SptA (AAX19896) and SptB (AAX19897) from *Natrinema* sp. J7, 172P1 (P29143) from *Nab. asiatica*, Nep (AAV66536), a putative subtilisin (Hly natria) (ADD04299) from *Nab. magadii*, R4 (BAA10958) from *Hfx. mediterranei*, NRC1 (NP_281139) from *Halobacterium* sp. NRC-1, and thermitase (1THM_A) from *Thermoactinomyces vulgaris* (B). A, the signal peptide (SP), the N-terminal propeptide (Pro), the catalytic domain and the C-terminal extension (CTE) of SptA precursor are indicated. The locations of four cysteine residues are shown. The numbers on the right or left side of the boxes represent the positions of the C- or N-terminal residues of the proteins starting from the N-terminus of mature enzyme. B, the number (149) in the sequence (Hly natria) represents the number of inserted residues at the position indicated. The numbers on the right side represent the positions of the residues starting from the N-terminus of precursor or mature enzyme (in parentheses). Conserved cysteine residues are boxed. The vertical arrow indicates the cleavage site of the CTE of SptA under reducing conditions. The underlined sequence represents the N-terminal residues of the major autolysis products of SptA under lower salt conditions (Fig. 4A). The positions of C-termini of truncation mutants (SptAΔC125 and SptAΔC76) are indicated above the sequences. The eight β-strands (β1–8) in the CTE of SptA were predicted using DNAman software (Lynnon Biosoft Inc.).

doi:10.1371/journal.pone.0023562.g001
cysteine residues involved in disulfide bonds, mutants C103S, C154S, C303S and C338S were expressed in Hfx. volcanii WFD11 and the culture supernatants were subjected to SDS-PAGE under reducing and non-reducing conditions. C103S and C154S showed the same results as wild type SptA: in non-reducing conditions, they migrated faster than their reduced forms (Fig. 3A). This result indicates that the wild type, C103S, and C154S are all more compact in non-reducing conditions and have a smaller radius of gyration [17] than in reducing conditions. In contrast, no difference was observed in migration behaviors of C303S or C338S under reducing and non-reducing conditions. This finding suggests that in the wild type SptA, Cys303 and Cys338 formed the disulfide bond that was prevented in the C303S and C338S mutants.

Mature SptA, C103S and C154S in culture supernatants displayed the same apparent molecular mass of approximately 62 kDa, while C303S and C338S showed an apparent molecular mass of approximately 40 kDa (Fig. 3A). N-terminal sequencing identified the first 5 residues of the 40 kDa protein as YTPND, the same as those of mature SptA, suggesting that the C303S and C338S mutants underwent a C-terminal cleavage event. To further clarify this situation, a C-terminal His-tagged version of SptA (SptAH) was expressed in Hfx. volcanii WFD11, and was purified from the culture supernatant by Ni-NTA affinity chromatography (Fig. 3B). The purified SptAH displayed properties similar to SptA [6] in terms of salt dependence, activity and stability (data not shown), indicating that the presence of the His-tag had no significant effect on enzyme function. In the presence of reducing agents, SptAH gradually converted to the 40 kDa protein (named SptAΔC110), accompanied by the appearance of a 25 kDa protein (named SptAC110), which could be fully degraded after heat-treatment at 70°C for 5 min (Fig. 3B). The five N-terminal amino acid residues of SptAC110 were very stable at 40°C for 3 h, suggesting that the cleavage occurred between Cys303-Gly304 (Fig. 1). In addition, the C-terminal cleavage was not observed in PMSF-inactivated SptAH, but its CTE could be processed after treatment with active SptAH under reducing conditions (data not shown), implying the cleavage occurred in an intermolecular autocatalytic manner. These results demonstrate that disulfide bond Cys303-Cys338 plays an important role in stabilizing SptA against autocleavage of the CTE.

The effect of the CTE on enzyme stability

To test the contributions of the CTE to enzyme stability, we first tested activity in low salt. At low enzyme concentrations (e.g., 0.1 μM), both SptAH and SptAΔC110 were very stable at 40°C in the presence of 3 M NaCl, and no loss of activity was observed after incubation at 40°C for 24 h. Meanwhile, the more concentrated enzymes (5 μM) showed slight autolysis at 3 M NaCl after incubation at 40°C for 3 h (Fig. 4A), suggesting only a...
small concentration dependence on autolysis. However, the two enzymes showed significant autolysis under lower salt conditions (e.g., 1 M NaCl), wherein SptAH could convert to two major autolysis products with apparent molecular mass of approximately 30 kDa (Fig. 4A). By N-terminal sequencing, the first five amino acid residues of the two products were identified as TEPGT and TSPDP, which coincided with the amino acid sequence of SptA after Thr287 and Gly291, respectively (Fig. 1). It was observed that the two products could bind to Ni-NTA affinity column (data not shown), implying the His-tag fused at the C-terminus was not processed. Therefore, they represent two forms of undigested CTE that are more resistant to proteolysis than the catalytic domain at low salinities. High temperatures were also used to test the stability limits of the enzymes and heat inactivation profiles of SptAH and SptAAC110 were examined at 70°C in 3 M NaCl (Fig. 4B). Results showed that SptAH autocleaved its CTE and lost its azocaseinolytic activity faster and to a greater degree than SptAAC110, which suggests that the CTE is important to activity.

The CTE interacts with the subtilisin domain via hydrophobic residues and these contacts have been proposed to provide stabilization to the enzymes [10,14]. To test this hypothesis, we used CD to investigate CTE contributions to enzyme structure in solution. Far-UV CD spectra of SptAH, SptAAC110 and CTE* were measured at different salinities or temperatures (Fig. 5). At 40°C, the mean residue ellipticity value of SptAH or CTE* is lower than that of SptAAC11010 in the presence of 3 M NaCl (Fig. 5A, C and E), as expected from the contributions of the β-strands in the CTE (Fig. 1) [18]. When the salinity was decreased to 1 M or the temperature increased to 70°C, SptAAC110 showed a larger global change in secondary structure than CTE* (Fig. 5C to F), indicating that the CTE domain is more stable than the catalytic domain. In contrast to SptAAC110, both SptAH and CTE* showed no significant change in secondary structure in the temperature range of 40–70°C (Fig. 5B, D and F), demonstrating that the CTE domain, in addition to being stable itself, helps the stability of the catalytic domain. At elevated temperatures, CTE* exhibited a more significant change in CD spectrum under reducing conditions than under non-reducing conditions (Fig. 5G and H), suggesting that disulfide bond Cys305-Cys338 confers additional stability to the CTE domain.

The importance of the CTE in enzyme activity

It was previously shown that SptA is most active on 0.5% azocasein in the presence of 2.5–3.0 M NaCl [6]. We investigated the salt dependence of our enzyme constructs to ascertain if the CTE assists the cleavage reaction. Using casein (1%) as the substrate, SptAH was most active at 2.0–3.0 M NaCl (A, C and E), or at different temperatures in the presence of 3 M NaCl (B, D and F), as described in Materials and Methods. In some cases (G and H), the spectra were recorded in the presence of 1 mM DTT. doi:10.1371/journal.pone.0023562.g005

Figure 5. The far UV CD spectra of SptAH, SptAAC110 and CTE*. The measurements were performed at a protein concentration of 0.2 mg/ml, either at 40°C in the presence of different concentrations of NaCl (A, C and E), or at different temperatures in the presence of 3 M NaCl (B, D and F), as described in Methods. In some cases (G and H), the spectra were recorded in the presence of 1 mM DTT.

doi:10.1371/journal.pone.0023562.g005

The Roles of the CTE of Halolysin SptA
effects, the salt dependence of SptAH and SptAΔC110 activities on azocasein were determined at a lower substrate concentration (0.25%). As shown in Fig. 6A, SptAH and SptAΔC110 displayed maximal azocaseinolytic activity above 4.5 M NaCl and at 3.5 M NaCl, respectively. Notably, SptAH showed higher azocaseinolytic activity than SptAΔC110 over the whole range of NaCl concentrations tested (0.5–4.5 M), and the difference became more pronounced as the salinity increased (Fig. 6A). In contrast, no significant difference was observed between the two enzymes in terms of their activities against the small synthetic substrate suc-AAPF-pNA, and these activities increased with rising salinity up to 4.75 M (Fig. 6A). A similar result was observed when suc-AAPL-pNA was used as an alternate small substrate (data not shown).

The variation in enzymatic salt dependence toward the two sizes of substrates may be due differing levels of salt-induced substrate conformation changes, which would be more significant in proteins than in peptides [20]. In order to determine whether the SptA catalytic domain and CTE can act in trans, we tested equimolar mixtures of SptAΔC110 and SptAC110. These showed a lower level of azocaseinolytic activity similar to that of SptAΔC110 alone (data not shown), implying the covalent link between the two domains is necessary for the higher activity of wild-type SptA.

SptAH and SptAΔC110 showed similar kinetic parameters for the hydrolysis of suc-AAPF-pNA (Table 1), indicating the CTE has no significant effect on the enzyme’s affinity for the small synthetic substrate. Kinetic characterization with protein substrates is difficult because these substrates contain more than one hydrolysable peptide bond. Nevertheless, the evidence that the SptAH/SptAΔC110 activities ratio increased as the substrate (azocasein) concentration decreased (Fig. 6B) implies SptAH has a higher affinity for the substrate than SptAΔC110. In addition, β-casein digestion products of both enzymes were the same (Fig. 6C), suggesting the CTE domain does not contribute significantly to the cleavage specificity of the enzyme towards casein. These results demonstrate that the CTE is beneficial for catalytic efficiency of SptA toward the larger protein substrate, most likely by facilitating the binding of the protein for catalysis.

**Binding of the CTE to insoluble substrates**

In order to further ascertain the contribution of the CTE to enzymatic catalysis, and to investigate possible CTE binding interactions with substrates, insoluble azocoll, type I collagen, elastin-orcein and keratin-azure were employed for activity and binding assays. The activity assay results showed that, while both enzymes were inert towards elastin-orcein and keratin-azure,
SptAH showed a higher activity (18.5±0.9 U/nmol) than SptΔC110 (11.5±1.0 U/nmol) toward azocoll at 3 M NaCl, confirming the importance of the CTE in enzyme activity. The binding assays were carried out at 0°C to prevent substrate breakdown as much as possible [21]. In the presence of 3 M NaCl, SptAH significantly bound to azocoll, and showed a weak binding capacity towards elastin-orcein. In contrast, SptΔC110 displayed no binding capacity for the four substrates tested (Fig. 7A), implying that the CTE alone is involved in the binding of the enzyme to these substrates. To test this hypothesis, CTE* was employed for binding assay. As shown in Fig. 7A, the binding capacities of CTE* towards azocoll, elastin-orcein and keratin-azure were similar to those of SptAH. However, CTE* differed from SptAH in that about 50% of CTE* could bind to insoluble type I collagen at 3 M NaCl, while most of SptAH remained in the supernatant under the same conditions. Since various smaller polypeptides were detected in the supernatant of the sample of SptAH but not in that of CTE* (Fig.7A), the difference in type I collagen-binding behaviors between the two proteins was most likely due to partial hydrolysis of the insoluble collagen by SptAH and subsequent products release into the supernatant. CTE* was also able to bind to insoluble type I collagen at low salinities (e.g., 0.15 M NaCl), and its collagen-binding capacity was not significantly affected by the addition of BSA (Fig. 7B). At 40°C, the amount of unbound CTE* slightly increased, probably due to partial solubilization of the substrate as evidenced by the occurrence of background signals in the lanes (Fig. 7B).

Conserved tertiary structure of the CTE

There has been no reported crystal structure of any halolysin so far. By using secondary structure prediction methods, eight β-strands could be found in the CTEs of SptA and other halolysins (Fig. 1). This was supported by the CD analysis of CTE*, which displays a spectrum typical of a β-strand backbone (Fig. 5). In addition, automated homology modeling of the SptA CTE yields a β-jelly roll-like structure (Fig. S1) that suggests a possible distant relationship to clostridial C-CPE (≈17% identity) and CBD (≈16% identity) domains. Although the similarity is low at the sequence level, the three domains have similar β-jelly roll-like structures and some key residues of clostridial C-CPE and CBD involved in receptor or collagen binding are conserved in the SptA CTE. For example, three Tyr residues near the extreme C-terminus of the C-CPE are critical for receptor binding [22], among them the residue Tyr310 is conserved in the SptA CTE (Tyr404) (Fig. S1B). In the case of clostridial CBD, three (Tyr970, Leu992 and Tyr996) of the five residues showing >5-fold reduction in collagen-binding ability upon mutation are present in the SptA CTE (Tyr568, Leu594 and Tyr598) (Fig. S1B). Moreover, three residues (Arg929, Phe952 and Val978) of clostridial CBD were shown to enhance collagen-binding ability upon mutation to Ala [23]. Interestingly, two Ala residues (Ala350 and Ala377) exist naturally in the SptA CTE at the positions corresponding to Phe952 and Val978 in clostridial CBD (Fig. S1B). Our results suggest that the CTE domain of SptA also has the ability to bind to collagen. We speculate that the binding of the CTE to substrates may be in a manner similar to the cases of clostridial CBD and C-CPE because of their structural similarity.

Discussion

Downstream of the subtilisin-like catalytic domain, halolysins 172P, Rh, Nep and SptA all have a respective CTE containing two cysteine residues (Fig. 1). The pioneer work of Kamekura et al. [5] suggested that the removal of the CTE from halolysin R4 abolished protease activity in culture supernatant of recombinant Hfs. volanii WFD11 and, thus, the CTE has been proposed to provide an essential (but as yet unknown) function. In the case of SptA, although the deletion of its CTE led to a sharp decrease in protease activity in culture supernatant of Hfs. volanii WFD11 expressing the mutant SptAAC125, the appearance of active enzyme in the culture supernatants indicates the CTE is not essential for the folding of SptA. Since the amount of the secreted active SptAAC125 is rather low relative to that of wild-type enzyme, we cannot exclude the possibility that the CTE may assist
in the correct folding of the enzyme or/and its secretion. It is observed that inactive pro-form of SptAΔC125 was detected in the insoluble fraction of the recombinant cell. This is unlikely due to a lack of the CTE, because wild-type pro-SptA also existed in the insoluble fraction. Further investigation of the mechanisms of folding and secretion of the enzyme is required to find possible explanation for this observation.

Many subtilases contain disulfide bond(s) that contribute to enzyme stability [9]. For halolysins, it has been reported that the substitution of Cys316 and Cys352 within the CTE of halolysin R4 decreased enzyme stability in hypotonic solutions, possibly owing to disruption of potential disulfide bonds or perturbation of calcium binding site(s). However, the stabilization mechanism of the two cysteine residues has not yet been elucidated because of the difficulty of enzyme purification [5]. In this report, the experiments of Cys→Ser mutations clearly indicate that Cys303 and Cys338 form a disulfide bond in the CTE of SptA, and disruption of the disulfide bond results in the autocleavage of the CTE. The two cysteine residues within the CTE of SptA are conserved in 172P1, Nep and SptB, while those of R4 are conserved in halolysin NRC1 (Fig. 1). Despite variation in their locations, all these cysteine residues reside in the loop regions connecting the secondary structure elements (Fig. 1). The formation of a disulfide bond, as observed in SptA, can endow the CTE with structural stability against proteolysis, which seems to be a common stabilization strategy shared by halolysins.

The presence of the CTE domain increases the affinity of SptA for larger protein substrate rather than smaller peptide substrate. One reasonable explanation for this is that the CTE domain is situated too far away from the catalytic cleft to participate in the binding of small substrate (Fig. S1A). Conversely, the CTE domain may provide additional binding site(s) for protein substrate, and thus be beneficial to hydrolysis activity. However, the CTE domain does not contribute significantly to the cleavage specificity of SptA towards casein, unlike the cases of kexin-like proteases in which the P-domains are not only involved in substrate binding but also play an important role in the regulation of the substrate specificity of these enzymes [12,24]. This distinction may be rationalized by consideration of the differences in physiological function and enzyme structure between SptA and kexin-like proteases. The members of kexin/furin family function as pro-protein processing proteases (convertases), and differ from other subtilases in that they have a high degree of specificity for cleavage after dibasic or multi-basic residues [8]. At the C-terminus of the P-domain in the kexin-like protease from A. sobria, there is an extra occluding region situated close to the active site, which may act as a steric obstacle and is important for substrate recognition (Fig. S1A) [12]. Unlike these convertases, halolysins serve as degradative proteases to provide halophilic archaea with small peptides and amino acids as nutrition [1]. The lack of the occluding region is favorable for SptA to have broad substrate specificity, thereby allowing the enzyme to deal with different protein substrates in the environment. Given that SptA is produced by a halophilic archaeon adapted to extremely saline environments, the role of its CTE domain in substrate binding seems to be of physiological importance. At high salt concentrations, proteolysis of protein substrates may be affected by limited accessibilities and availabilities of the substrates due to salt-induced conformation changes and precipitation ("salting-out" effect). Obviously, the enhanced substrate affinity afforded by the CTE domain would be advantageous for halolysins to fulfill their physiological roles at high salinities.

Recently, several crystal structures of subtilases with CTEs have been determined. Although their CTEs have very low or almost no sequential homology, they nonetheless fold into a β-jelly roll-like structure, such as the P-domains of mammalian furin [14], yeast Kex2 [13] and kexin-like protease from A. sobria [12], the C-domain of the protease Kf-13 from Bacillus sp. [9], the Fn III domain of tomato subtilase 3 [11], as well as the β-jelly roll domain of subtilisin Tk-SP from T. kodakaraensis [10]. Additionally, fervidolysin from Firmicutes from Bacillus subtilis [52] and streptoccocal C5a peptidases [26,27] possess two and three β-jelly roll domains at their C-termini, respectively. β-jelly roll-like structures are common in substrate or receptor-binding domains for other bacterial proteins, such as clodirilcal C-CPE [22] and CBD [23]. It is interesting that the CTE domain of SptA has the ability to bind to collagen at low salinities (e.g., 0.15 M NaCl). CBD has been well studied in mammalian matrix metalloproteinases and bacterial collagenases [28], but has not yet been reported in archaea. A large focus of protein engineering is the creation of molecules that bind to and enhance the delivery of therapeutics across the dermal barrier. The clodirilcal CBD has been used in a drug delivery system by acting as an anchoring unit to collagen, the major constituent of extracellular matrix of animals [29]. Recently, the CBD of the collagenase from thermophile Geobacillus collagenovorans MO-1 has been proposed to have the potential for applications to such a drug delivery system by taking advantage of the non-pathogenicity of the strain [21]. Likewise, the collagen-binding capacity of the CTE domain from a non-pathogenic halophilic archaeon makes it a potential candidate for clinical applications. From a practical view point, durability and sustainability are highly desired for drug delivery systems. The apparent stabilization effect of the disulfide bond Cys303-Cys338 on the CTE domain of SptA might provide a rational basis for improving the stability of the anchoring unit.

**Materials and Methods**

**Strains and growth conditions**

*Natronemina* sp. J7 was isolated from a salt mine in China [6], and has been deposited in the China Center for Type Culture Collection (CCTCC) under the accession number AB91141. *Hfx. volcanii* WFD11 was used as the host for expression. These two halophilic archaea were cultured as described previously [6]. *Escherichia coli* JM110 and E. coli BL21(DE3) were used as the hosts for routing cloning and expression, respectively, and were grown at 37°C in Luria-Bertani (LB) medium supplemented with ampicillin (50 µg/ml) or kanamycin (30 µg/ml) as needed.

**DNA manipulation, plasmid construction and mutagenesis**

The genomic DNA of *Natrenema* sp. J7 was prepared according to the method of Kamekura et al. [4], and was used as the template for PCR. The primers used in this study are listed in Table S1. The gene of wild type SptA was amplified with primers sptA-ATG1 and sptA-R1. The genes encoding the C-terminal truncation mutants (SptAAC125 and SptAAC76) and the C-terminal His-tagged version of halolysin SptA (SptAH), were amplified from the genomic DNA with the primer sptA-ATG1 in combination with the primers sptAAC125-R, sptAAC76-R, and sptAH-R, respectively. Overlapping PCR was employed as described previously [16] to introduce the C103S or C154S mutations into the SptA gene. Briefly, the 5’ end of the gene was amplified with the primer sptA-ATG1 in combination with the primer sptAC103S-R or sptAC154S-R, and the 3’ end of the gene was amplified with the primer sptA-R1 in combination with the primer sptAC103S-F or sptAC154S-F, respectively. The “mega-primer” method of site-directed mutagenesis [30] was used for
substitutions C303S and C338S, with primers sptAC303S-F and sptAC338S-F used as the mutagenic primers. The amplified genes were ligated into the Nde-I-Nco restriction site of pSY1 [6] to construct the expression plasmids pSY1-sptAC303, pSY1-sptAC338, pSY1-sptAC125, pSY1-sptAC103S, pSY1-sptAC154S, pSY1-sptAC303S and pSY1-sptAC338S, respectively. This new expression plasmid pSY1-sptAC for wild type SptA differs from the previously constructed pSPTA1 [6] in that the latter contains additional 104 nucleotides downstream from the stop codon. All recombinant plasmids have been confirmed by DNA sequencing.

Expression and purification

The expression plasmids were amplified in *E. coli* JM110 and then transferred into *Hfx. volcanii* WFD11 [31]. The transformed cells were plated on 18% MGM agar plates containing 1% skim milk and 0.2 mg/ml novobiocin, and incubated at 37 °C for 7 days. Successful *Hfx. volcanii* WFD11 transformants were identified by formation of halos around colonies [6].

*Hfx. volcanii* WFD11 transformants harboring different expression plasmids were subsequently cultivated in 18% MGM liquid medium containing 0.2 mg/ml novobiocin and 0.15 M NaCl, pH 8.0, and then suspended in buffer C (50 mM Tris-HCl, 10 mM CaCl2, 0.15 M NaCl, pH 8.0), and then suspended in buffer C (50 mM Tris-HCl, 10 mM CaCl2, 8 M Urea, pH 8.0) and disrupted by sonication. The supernatant was collected by centrifugation at 13,000 × g for 10 min at 4 °C and then subjected to a Ni-charged Chelating Sepharose Fast Flow column for purification. After washing the column with buffer C containing 40 mM imidazole, the Ni-bound protein was eluted with buffer C containing 200 mM imidazole. The purified protein was dialyzed against buffer A before use.

The culture supernatant containing SptAH was applied to a Ni-charged Chelating Sepharose Fast Flow resin (Amersharm Biosciences, Sweden) column (1.6 × 20 cm) equilibrated with buffer A. After washing the column with buffer A containing 20 mM imidazole, the Ni-bound SptAH was eluted with the same buffer containing 200 mM imidazole, and the eluted fraction was dialyzed against buffer A at 4 °C. The purified SptAH was supplemented with 10 mM dihiothreitol (DTT), and then incubated at 40 °C for 90 min, until SptAH had fully converted to SptAΔC110 (SptA minus the CTE) and SptAΔC110 (the processed CTE). The sample was subsequently re-applied to a Ni-charged Chelating Sepharose Fast Flow column. The wash fractions containing SptAΔC110 were collected and His-tagged SptAΔC110 was eluted from the column as described above. The samples were dialyzed against buffer A at 4 °C and used as the purified proteins. The protein sample was concentrated with a Microcon YM-3 centrifugal filter (Amicon) as needed. The protein concentration was determined using a Bio-Rad Bradford protein assay kit (Bio-Rad Laboratories, Inc.) with BSA as the standard.

Preparation of SptA precursor and the isolated CTE domain (CTE*). The gene fragments encoding SptA precursor and the CTE* (from Thr287 to Arg 413, Fig. 1) were amplified from the genomic DNA of *Natrinema sp.* J7 with primer pairs spta-ATG1 and spta-AI-1, spta-ATG1 and spta-AI-2, SptAC125-F and SptAC125H (Supplemental Table 1), respectively, and then ligated into the Nde-I-HindIII sites of pET26b to construct expression vectors pET26-sptaA and pET26-sptaAΔCTE. Then, *E. coli* BL21(DE3) cells individually containing the expression vectors were cultured in LB medium supplemented with 1% glucose and 30 μg/ml kanamycin at 37 °C until the OD600 reached approximately 0.6. The target proteins were induced with 0.4 mM isopropyl β-D-thiogalactopyranoside (IPTG), and the cultivation was continued at 37 °C for 4 h. The cells were washed with buffer B (50 mM Tris-HCl, 10 mM CaCl2, 0.15 M NaCl, pH 8.0), and then suspended in buffer C (50 mM Tris-HCl, 10 mM CaCl2, 8 M Urea, pH 8.0) and disrupted by sonication. The supernatant was collected by centrifugation at 13,000 × g for 10 min at 4 °C and then subjected to a Ni-charged Chelating Sepharose Fast Flow column for purification. After washing the column with buffer C containing 40 mM imidazole, the Ni-bound protein was eluted with buffer C containing 200 mM imidazole. The purified protein was dialyzed against buffer A or B before use.

Enzyme activity assay

Unless otherwise indicated, (azo)caseinolytic activity of the enzyme was measured at 40 °C for 30 min in 400 μl of reaction mixture containing 0.25% (w/v) azocasein (Sigma A2765, MW = 23600) or 1% (w/v) casein (Sigma) and 200 μl of enzyme sample in buffer A. The reaction was terminated by the addition of 400 μl of 40% (w/v) trichloroacetic acid (TCA) into the reaction mixture. After incubating the mixture at room temperature for 15 min, it was centrifuged at 13,400 × g for 10 min, and the absorbance of the supernatant was measured in a 1 cm cell at 360 nm (azocasein) or 280 nm (casein). One unit (U) of caseinolytic activity was defined as the amount of enzyme required to release 1 μg of soluble azopeptidase per minute. The specific absorption coefficient (A280 = 40) of the azocasein solution was calculated by measuring its absorption after total digestion [32].

The enzymatic activity towards azo dye-impregnated collagen (azocoll), keratin-azure or elastin-orcein (Sigma) was measured as described previously [33], except that the reaction was performed at 40 °C for 60 min in buffer A.

The enzymatic activity on suc-AAPF-pNA (Sigma) was measured at 40 °C in buffer A containing 0.2 mM substrate unless indicated otherwise. The activity was recorded by monitoring the initial velocity of suc-AAPF-pNA hydrolysis at 410 nm in a thermostated spectrophotometer (Cintra 10e, GBC, Australia), calculated on the basis of an extinction coefficient for p-nitroaniline of 8,480 M⁻¹ cm⁻¹ at 410 nm [34]. One unit (U) of enzyme activity was defined as the amount of enzyme that produced 1 μmol of pNA per minute under the assay conditions.

Measurement of kinetic parameters

Suc-AAPF-pNA was used as the substrate to determine the kinetic parameters of the enzymes. Kinetic parameters were calculated from the initial velocity of hydrolysis in 40 °C in buffer A, with a substrate concentration range of 0.1–2.5 mM. K_m and k_cat values were obtained by using the nonlinear regression Table Curve 2D software (Jandel Scientific, version 5.0).

SDS-PAGE, N-terminal sequencing and immunoblot analysis

The SDS-PAGE was carried out employing glycine-Tris [35] or Tricine-Tris buffer systems [36]. The N-terminal sequencing, preparation of antisera against SptA precursor and immunoblot analysis were carried out as described previously [33].

Far-UV circular dichroism (CD) analysis

The far-UV CD spectra were measured on a Jasco J-810 spectropolarimeter (Jasco Corporation, Tokyo, Japan) at a protein concentration of 0.2 mg/ml using a 1 mm path length cell. In the...
cases of SptA and SptAΔC110, the enzymes were first inactivated by treatment with 10 mM phenylmethylsulfonyl fluoride (PMSF) in buffer A for 2 h at 30°C, followed by dialysis overnight against buffer A alone at 4°C. Since the dynode voltage exceeded 0.6 kV below 205 nm due to the high NaCl concentration, CD spectra were recorded from 205 to 260 nm. After subtracting a corresponding solvent spectrum, the sample spectrum was converted to the mean residue ellipticity [θ] deg cm² dmol⁻¹ by using mean residue molecular weights of 103,234 Da (SptA), 101,155 Da (SptAΔC110), and 108,855 Da (CTE*).

Binding assay on insoluble substrates
Azocoll, keratin-azure, elastin-orcinol and type I collagen (Sigma) were used as the insoluble substrates. Substrate-binding assays were performed according to the batch method of Itoi et al. [21]. Briefly, using either buffer A or buffer B, 5 mg of the insoluble substrate was washed with ice-chilled buffer 3 times then suspended in 200 μl of buffer and kept at 0°C for 60 min. After centrifugation at 10,000 × g for 5 min, the pellet was suspended in 80 μl of buffer containing enzyme at a concentration of 15 μg/ml. Unless otherwise indicated, the mixture was incubated at 0°C for 60 min, then the soluble fraction was recovered by centrifugation as described above and subjected to SDS-PAGE analysis. Protein bands were quantitated by densitometry using GeneTools software (Syngene).

Supporting Information

Figure S1 Superimposition of SptA (A) and its CTE (B) with their homologues. The structure models of SptA and its CTE (green) were generated by automated homology modeling using SWISS-MODEL (http://swissmodel.expasy.org), with the kexin-like serine protease (ASP) (cyan) from A. sobria (PDB code 3HJR) and the CBD (yellow) of the collagenase from C. histolyticum (PDB code 1NQD) as the templates, respectively. The figure was prepared by PyMol (http://www.pymol.org). A, the active site residues Asp38, His79 and Ser232 of SptA are indicated by D, H and S, respectively. B, the lower panel shows the sequence alignment of the CTE (AAAX19896), the CBD (BAA77453) and the C-CPE of C. perfringens enterotoxin (AA721210). The residues indicated with stars in the CBD or the C-CPE represent those involved in collagen or receptor binding. Arrow heads indicate the three residues shown to increase collagen-binding ability of the CBD when mutated to Ala.

(TIF)

**Table S1** Oligonucleotide primers used in this study.

(DOC)

**Acknowledgments**

The authors thank Dr. W. F. Doolittle (Dalhousie University, Canada) for kindly providing Hfx. volcanii WFD11.

**Author Contributions**

Conceived and designed the experiments: ZX BT X-FT. Performed the experiments: ZX XD TL FG. Analyzed the data: ZX BT X-FT. Wrote the paper: ZX BT X-FT.
proteins having a collagen-binding domain. Proc Natl Acad Sci U S A 95: 7018–7023.

30. Sarkar G, Sommer SS (1990) The “megaprimer” method of site-directed mutagenesis. Biotechniques 8: 404–407.

31. Cline SW, Lam WL, Charlebois RL, Schalkowyk LC, Deolittle WF (1989) Transformation methods for halophilic archaeabacteria. Can J Microbiol 35: 148–152.

32. Wang CC, Houng HC, Chen CL, Wang PJ, Kuo CF, et al. (2009) Solution structure and backbone dynamics of streptopain: insight into diverse substrate specificity. J Biol Chem 284: 10957–10967.

33. Cheng G, Zhao P, Tang XF, Tang B (2009) Identification and characterization of a novel spore-associated subtilase from Thermoactinomyces sp. CDF. Microbiology 155: 3661–3672.

34. DelMar EG, Largman C, Brodrick JW, Geokas MC (1979) A sensitive new substrate for chymotrypsin. Anal Biochem 99: 316–320.

35. King J, Laemmli UK (1971) Polypeptides of the tail fibres of bacteriophage T4. J Mol Biol 62: 465–477.

36. Schagger H, von Jagow G (1987) Tricine-sodium dodecyl sulfate-polyacrylamide gel electrophoresis for the separation of proteins in the range from 1 to 100 kDa. Anal Biochem 166: 368–379.