Solution Structure and Backbone Dynamics of Streptopain

**INSIGHT INTO DIVERSE SUBSTRATE SPECIFICITY**

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Streptococcal pyrogenic exotoxin B (SPE B) is a cysteine protease expressed by Streptococcus pyogenes. The D9N, G163S, G163S/A172S, and G239D mutant proteins were expressed to study the effect of the allelic variants on their protease activity. In contrast to the other mutants, the G239D mutant was ~12-fold less active. The Gly-239 residue is located within the C-terminal S230-G239 region, which cannot be observed in the x-ray structure. The three-dimensional structure and backbone dynamics of the 28-kDa mature SPE B (mSPE B) were determined. Unlike the x-ray structure of the 40-kDa zymogen SPE B (proSPE B), we observed the interactions between the C-terminal loop and the active site residues in mSPE B. The structural differences between mSPE B and proSPE B were the conformation of the C-terminal loop and the orientation of the catalytic His-195 residue, suggesting that activation and inactivation of SPE B is involved in the His-195 side-chain rotation. Dynamics analysis of mSPE B and the mSPE B/inhibitor complexes showed that the catalytic and C-terminal loops were the most flexible regions with low order parameter values of 0.5 to 0.8 and exhibited the motion on the ps/ns timescale. These findings suggest that the flexible C-terminal loop of SPE B may play an important role in controlling the substrate binding, resulting in its broad substrate specificity.

Streptococcus pyogenes (group A Streptococcus (GAS)), one of the most common human bacterial pathogens, has developed diverse mechanisms that allow the bacteria to evade the immune system (1, 2). This bacteria causes a variety of human diseases, including pharyngitis, cellulitis, necrotizing fasciitis, streptococcal toxic shock syndrome, scarlet fever, acute rheumatic fever, rheumatic heart disease, and glomerulonephritis (3, 4). Virtually all strains of GAS isolated from patients with invasive disease express an extracellular cysteine protease known as streptopain (EC 3.4.22.10), with synonyms including streptococcal pyrogenic exotoxin B (SpeB or SPE B), streptococcal peptidase A, SPP, and streptococcal cysteine protease (SCP) (5–10). The level of SPE B activity is associated with the extent of gross pathological changes induced by the specific strains displaying varied degrees of virulence (5–7). Many reports also suggest that SPE B is an important virulence factor in streptococcal infections (11–14).

SPE B produced from GAS is released extracellularly to culture medium as a zymogen (proSPE B) with a molecular mass of 40 kDa. The conversion of proSPE B to the 28-kDa active mature SPE B (mSPE B) can be achieved by autoproteolysis and exogenous proteases (8–10). SPE B has diverse substrate specificity and is involved in the processing of host proteins (10, 14). SPE B degrades extracellular matrix proteins fibronectin and vitronectin and increases bacterial attachment to host cells (6, 15). Additionally, it cleaves and activates matrix metalloproteases 2 and 9, which increase bacterial dissemination (16). It also causes severe inflammation in the host by activating interleukin 1β (17). In addition to its ability to cleave host proteins, SPE B releases streptococcal surface proteins, including IgG-binding proteins and properdin. Thereby, SPE B avoids complement activation (18, 19). However, little is known about the substrate specificity of SPE B.

The three-dimensional structure of the 40-kDa C47S mutant of proSPE B was determined by x-ray crystallography. Three regions of the prodomain (the residues D1p-N3p, S22p-G34p, and T114p-K118p) and one region of the protease domain (the residues Ser-230 to Gly-239) are undefined and assumed to be flexible (20). The prodomain has a unique fold and consists of a central four-stranded β-sheet packing by two α-helices. Despite a low sequence identity, the protease domain of SPE B exhibits a canonical papain fold, which consists of a three α-helix N-terminal domain and a four-stranded β-sheet C-terminal domain. The catalytic residues, Cys-47 and His-195, are located at the molecular surfaces between the N- and C-terminal domains (21). Unlike most other cysteine proteases containing a Cys-His-Asn triad, SPE B has a Cys-His catalytic dyad. The His-195 residue is located at the catalytically incompetent posi-
tion, resulting from the position displacement by the N89p residue from the prodomain.

In our mutagenesis study, we found that the Gly-239 residue played an important role in its protease activity, and is located within an undefined S230-G239 loop in the x-ray structure of proSPE B. In our previous study, we have determined the backbone $^1$H, $^{13}$C, and $^{15}$N resonances for the 28-kDa C47S mutant of mSPE B and identified all nuclear magnetic resonance (NMR) resonances of the undefined S230-G239 loop from multidimensional NMR spectroscopy (22). Therefore, we used NMR spectroscopy to determine three-dimensional structures and backbone dynamics of the 28-kDa C47S mutant of mSPE B so that the presence of an undefined S230-G239 loop in the protease domain of proSPE B and the possibility of the loop movements can be resolved. We also identified the binding site of mSPE B to its inhibitor L-trans-epoxysuccinyl-leucylamido (4-guanidino)butane (E-64) by comparing NMR chemical shift difference between the 28-kDa C47S mutant and the mSPE B/E-64 complex. The mutation on SPE B was used to examine the binding site of mSPE B for its inhibitor E-64. Our results provide a structural basis for understanding the mechanism of how the conformation and dynamics properties of the loops of SPE B controls its substrate binding, resulting in its broad substrate specificity.

**EXPERIMENTAL PROCEDURES**

**Expression, Construction, and Purification of SPE B Mutants**—The expression and purification of SPE B mutants were performed as described previously (10). The genomic DNA of GAS was extracted from strain A20. The structural gene of the 371-residue proSPE B was amplified by polymerase chain reaction (PCR) and then cloned into the pET-21a vector. The wild-type construct was used to produce D9N, G136A, G163S, G163S/A172S, V189A, and G239D mutations using overlap extension PCR (23). The recombinant plasmid was transformed into the Escherichia coli BL21(DE3)pLyS strain, and the system was inducibly expressed under the control of a strong T7 promoter.

Recombinant SPE B mutants were produced by growing cells at 28 or 37 °C for 12 h in LB medium (10 g of Bacto tryptone, 5 g of Bacto yeast extract + 10 g NaCl in 1 liter). Cells were harvested by centrifugation and lysed by liquid shear with a French press to obtain the extract. The proteins were purified by Ni$^{2+}$-chelating chromatography (Amersham Biosciences, Piscataway, NJ) with a gradient of 20–200 mM imidazole. The proteins were concentrated by Amicon ultrafiltration using a 10-kDa cutoff membrane and then exchanged with phosphate-buffered saline. To prevent the conversion, mercuric chloride was added to the extract to a final concentration of 1 mM, which was retained throughout the purification. The final solutions were stored at −20 °C, and the recombinant proteins were activated by adding 5 mM dithiothreitol (DTT) and 5 mM EDTA (Sigma-Aldrich) before use. The purification of proteins was greater than 95%, as judged by SDS-PAGE.

Labeled NMR samples were prepared as described previously (22). M9 minimal media was used and 1 gram/liter $^{15}$NH$_4$Cl (99% $^{15}$N), 2 gram/liter $^{15}$C glucose (99% $^{13}$C), and/or 98% D$_2$O were substituted for the unlabeled compounds in the growth media. Selective [$\alpha$-$^{15}$N] - A-, -D-, -E-, -F-, -G-, -H-, -I-, -K-, -L-, -M-, -Q-, -V-, or -Y-labeled proteins were prepared using the protocol described by McIntosh et al. (24). The 28-kDa mutants of mSPE B were obtained by digesting 20–40 mg/ml of the 42-kDa mutants of proSPE B with 50–100 μg/ml of mSPE B. The reaction solution was then purified with Bio-Gel P-30 (Bio-Rad) gel filtration chromatography. The final solution was also concentrated by amicon ultrafiltration using a 10-kDa cutoff membrane. The yields of 28-kDa proteins were >60%, and the purification of proteins was greater than 95% according to SDS-PAGE analysis.

**Mass Spectrometric Measurement**—The molecular weights of proteins were confirmed using an API 365 triple quadrupole mass spectrometer equipped with a TurboionSpray source (PE Sciei, Thornhill, Ontario, Canada). The protein solutions (1–10 μM in 50–90% methanol or acetonitrile with 0.1% formic acid) were infused into the mass spectrometer using a syringe pump (Harvard Apparatus, South Natick, MA) at a flow rate of 12–20 μl/min to acquire the full scan mass spectra. The electrospray voltage at the spraying needle was optimized at 5000–5300 V. The molecular weights of proteins were calculated by computer software provided with the API 365 mass spectrometer.

**Activity Measurement**—The azocasein assay was used to test for proteolytic activity of streptopain and mutant proteins. The assay was modified as previously described (25). Activity was determined by measuring the hydrolysis of azocasein based on the increase in absorbance at 366 nm against time as described. The reaction was initiated by adding 20 μl of streptopain or mutant proteins to 160 μl of the reaction mixture containing 2.7 mg/ml of azocasein and 5 mM DTT and 5 mM EDTA in a phosphate-buffered saline buffer. After incubating the solution at 37 °C for designated time intervals ranging from 0 to 24 h, the reaction was stopped by adding 40 μl of 15% of ice-cold trichloroacetic acid. Absorbance was measured using a Beckman Model DU 640 spectrophotometer. One enzyme unit was defined as the amount of protease required to release 1 μg of soluble azopeptides per minute. The specific absorption coefficient, $A_{366}^{\text{iso}} = 40$, of the azocasein solution was calculated by measuring its absorption after total digestion (25).

The proSPE B C47S mutant was also used as the substrate for activity assay because it does not exhibit any enzyme activity and exists as a 42-kDazymogen, (8). The reaction was carried out in a total of 20 μl of phosphate-buffered saline buffer containing 5 mM EDTA and 5 mM DTT. A final concentration of 1.2 μM of purified or recombinant protein was incubated with 24 μM 42-kDa C47S mutant at 37 °C with designated time intervals ranging from 0 to 14 h. The reactions were quenched by adding 5 μl of 50 μM E-64 with subsequent incubation at 37 °C for 30 min. The solution was then heated at 100 °C for 10 min and a 12% SDS-PAGE was used for analysis. Gels were scanned by a Vilber Lourmat model CN-TCX imaging system, and the band intensities were integrated with the software BIO-1D version 5.07. Proteolytic activity was measured as the disappearance of proSPE B. Relative reaction rates were obtained from the change in intensity of the 42-kDa C47S proSPE B band against time.

**NMR Spectroscopy**—Triple resonance experiments were recorded with 1.2–2.5 mM of 2H (70%), $^{15}$N/$^{13}$C (99%)-labeled...
proteins, and deuterium decoupling was applied. The samples were made to 10 or 100% D₂O at pH 6.0 in 20 mM phosphate buffer. NMR experiments were performed on a Bruker Avance 600 spectrometer at 27 °C. Triple resonance experiments of TROSY-HNCA, TROSY-HN(CO)CA, TROSY-HN(CA)CB, TROSY-CBCA(CO)NH, HBHA(CBCA)NH, HBHA(CBCA-CO)NH, H(CCCO)NH, (H(C(CCO)NH. ¹⁵N-edited, and ¹³C-edited TOCSY-HSQC, HCCH-TOCSY, and -COSY were carried out for the purpose of proton, carbon, and nitrogen resonance assignments (26–29). Two-dimensional COSY, TOCSY, and NOESY experiments with an unlabeled sample in D₂O provided the basis for aromatic proton assignments and hydrogen bond determination (30). Distance restraints were obtained from three-dimensional ¹⁵N-edited and ¹³C-edited NOESY-HSQC with mixing times of 80 and 120 ms. TheHNHA experiment was performed to measure JH-NH coupling constants and to obtain φ dihedral angles (31). The φ and θ backbone dihedral angles were also calculated by the TALOS program (32). Data were processed and analyzed using the XWINNMR and Aurelia programs on an O2 Silicon Graphics workstation.

Chemical shift perturbation was used to map the interactions between SPE B and the inhibitor E-64. The weighted average change in the chemical shift of the amide nitrogen and proton atoms was measured through ¹H-¹⁵N HSQC spectra and treated according to Equation 1 (33).

$$\Delta \delta = (0.5 \times (0.25 \Delta \delta_{NN}^2 + \Delta \delta_{NN}^2))^{1/2} \quad \text{(Eq. 1)}$$

In this equation, Δδ_N and Δδ_H represent the changes in nitrogen and proton chemical shifts (in parts per million), respectively.

Structure Calculations—Structures were calculated using the program X-PLOR Version 3.851, and by utilizing the hybrid distance geometry-constrained simulated annealing method (34–35). NOESY cross peak intensities were categorized into strong, medium, and weak. These intensities were then converted into distance constraints of 1.8–2.8, 1.8–5.0, and 2.5–6.0 Å, respectively. Pseudoatom corrections were used for methylene, methyl, and aromatic protons, and an additional 0.5 Å was added to the upper limit distances involving methyl protons. Hydrogen bond restraints were included in calculations only if the amide protons were slowly exchanging and if the β-strand interstrand NOE cross-peaks were observed. Each hydrogen bond was enforced by two distance restraints of 1.8–2.3 Å (amide proton to carbonyl oxygen) and 2.8–3.3 Å (amide nitrogen to carbonyl oxygen). A family of 100 structures was generated using NOE distance, dihedral angle, and hydrogen bond restraints. During the first phase of dynamics at 2000 K, the value of the force constant of the NOE term was kept constant at 50 kcal·mol⁻¹·Å⁻². The repulsion term was gradually increased from 0.03 to 4.0 kcal·mol⁻¹·Å⁻², and the torsion angle term from 5 to 200 kcal·mol⁻¹ rad⁻². The simulated annealing refinement consisted of a 9-ps cooling dynamics followed by 200 cycles of Powell minimization. The 20 lowest energy structures were accepted based on violations of distance restraints of less than 0.5 Å, dihedral angle restraints of less than 5°, a van der Waals energy cut-off value of 45 kcal/mol, and an NOE energy cutoff value of 70 kcal/mol. The structure figures were prepared using the program MOLMOL (36).

Measurements of NMR Dynamics—Backbone dynamics of SPE B and its E-64 inhibitor complex were studied by two-dimensional proton-detected heteronuclear NMR spectroscopy. The ¹⁵N spin-lattice (R₁) and spin-spin (R₂) relaxation rate constants and steady-state ¹H-¹⁵N NOEs were measured from ¹H-detected ¹H-¹⁵N correlation spectra recorded with sensitivity-enhanced pulse sequences (37). A series of 10 experiments with relaxation delays of 20, 100, 250, 400, 500, 600, 800, 1000, 1500, and 3000 ms were carried out to measure T1. A series of 10 experiments with relaxation delays of 30, 50, 70, 90, 110, 150, 200, 300, 500, and 1000 ms were carried out to measure T₂. The longitudinal and transverse relaxation rate constants, R₁ and R₂, were obtained from exponential fits of the peak height data using least-squares fit software SigmaPlot (Jandel Scientific Software, San Rafael, CA). The reported R₁ values are the mean values of two independent data sets. In NOE experiment, two spectra one with the NOE and one without were collected. A recycle delay of 6 s was used, and 256 complex t₁ increments of 32 scans were acquired. The NOE effect was calculated as the ratio of peak heights in spectra collected with and without NOE. The reported NOE value was the average value from three pairs of NOE experiments.

The heteronuclear ¹⁵N relaxation rate constants, R₁ and R₂, and the ¹H-¹⁵N steady state NOE values were analyzed using the TENSOR2 program (38). This program is based on the model-free formalism that was pioneered by Lipari and Szabo and further extended by Clore et al. (39–41). In this approach, the overall and internal molecular motions were assumed to be independent, and the spectral density function for a molecule undergoing isotropic tumbling was calculated using the appropriate expression in Equation 2,

$$J(\omega) = 2/5[(S²\tau_m/1 + (\omega\tau_m)^2)] + (S² - S²\tau/1 + (\omega\tau)^2)] \quad \text{(Eq. 2)}$$

where 1/τ = 1/τ_m + 1/τ_φ and S² = S²φS²_φ, τ_m is overall rotational correlation time of the molecule, τ_φ is the effective correlation time for the motions on the slower of the two time scales, S²φ is the square of the generalized order parameter, and S²_φ and S² are the squares of order parameters for the motion on the fast and slow time scale, respectively (39–41).

Protein Data Bank Accession Number and NMR Assignment—The coordinates of 20 calculated structures of the SPE B C47S mutant have been deposited in the Protein Data Bank under accession number 2JT. ¹H and ¹⁵N resonances of Rho have been deposited in the BioMagResBank data bank under accession number BMRB-5547.

RESULTS

Expression and Protease Activity of SPE B Mutants—To identify functional mutation from allelic polymorphism in the SPE B gene, the D9N, G163S, G163S/A172S, and G239D mutant proteins were expressed in E. coli and purified to homogeneity. We also expressed G136A, V189A, W212A, and W214A to examine the residues near the active site. All recombinant proteins contained eleven extra vector residues (ASMTGGQQMGS) and six histidine residues to simplify the purification procedures. They were purified to apparent homogeneity in a single
step by Ni²⁺-chelating chromatography. 42-kDa recombinant wild-type and polymorphic mutants of SPE B were purified by adding 1 mM HgCl₂ as an inhibitor to prevent the conversion to 28-kDa form. In contrast, C47S mutant was purified as a 42-kDa zymogen without adding HgCl₂ as an inhibitor. The final yields of mutant proteins were about 40–250 mg per liter. Based on SDS-PAGE analysis, the wild-type and mutant proteins were homogenous (data not shown). The experimental molecular weights of SPE B mutants were determined by mass spectrometric measurement. The deviations of their molecular weights were less than 4 Da when compared with the theoretical values.

We used both azocasein and the 42-kDa inactive C47S mutant as the substrates to examine the proteolytic activities of the mutant proteins of SPE B (Table 1 and Fig. 1). The D9N, G163S, and G163S/A172S mutants were varied from 0 to 38% reduction in proteolytic activity compared with the wild-type enzyme. Interestingly, the G239D mutant was 14-fold less active. It was shown that the maturation of proSPE B goes through multiple step processing involving seven intermediates and two final products (10). Similar intermediates are involved in the trans-processing of the proSPE B C47S mutant by 28 kDa mSPE B. The trans-processing of the proSPE B C47S mutant by D9N, G163S, and G163S/A172S mutants also produced similar patterns, and Fig. 1B showed a representative example of D9N mutant. In contrast, only two final products were found from the proSPE B C47S mutant by G239D mutant (Fig. 1C), which is similar to the processing patterns identified from the mutation on the residues near active site, such as the W212A mutant (10). These results indicate that the residues in the C-terminal loop may also play an important role in its activity. However, the G239 residue is located within the C-terminal loop (R223-Q245), and the S230-G239 region cannot be observed in an x-ray structure. To understand the role of the C-terminal loop in SPE B, we used NMR spectroscopy to determine three-dimensional structures and backbone dynamics of 28-kDa mSPE B and its inhibitor complexes.

Structure Determination—The solution structure of mSPE B C47 mutant was determined by NMR spectroscopy and the hybrid distance geometry-dynamical simulated annealing method. NOE-derived distance restraints were obtained from the ¹⁵N- and ¹³C-edited NOESY data, and the x-ray structure of proSPE B was used to resolve ambiguous NOE. We found that the undefined S230-G239 region in the x-ray structure of SPE B-C47S and SPE B.

| Proteins       | Substrates | Azocaseinᵃ | C47S mutantᵇ |
|----------------|------------|------------|--------------|
| Wild-type      |            | 341 ± 22   | 378 ± 38     |
| D9N            |            | 256 ± 15   | 250 ± 25     |
| G163S          |            | 348 ± 15   | 385 ± 24     |
| G163S/A172S    |            | 286 ± 19   | 236 ± 23     |
| G239D          |            | 35.3 ± 2.8 | 20.0 ± 1.1   |
| G136A          |            | 22.3 ± 1.7 | 8.1 ± 0.3    |
| V189A          |            | 2.6 ± 0.1  | 0.9 ± 0.1    |
| W212A          |            | 1.2 ± 0.2  | 0.9 ± 0.1    |
| W214A          |            | 0.8 ± 0.1  | 0.8 ± 0.1    |

ᵃ One enzyme unit was defined as the amount of soluble protease required to release 1 µg of soluble azopeptides per minute.
ᵇ One enzyme unit was defined as the amount of soluble protease required to release 1 µg of SPE B C47S mutant per minute.

FIGURE 1. Proteolytic activity of SPE B and its mutants. SDS-PAGE analysis of the digestion of the proSPE B C47S mutant by (A) wild-type, (B) D9N, (C) G239D, (D) G136A, and (E) V189A mutants of streptopain are shown. A final concentration of 1.2 μM of native or recombinant streptopain protein was incubated with 24 μM of the proSPE B C47S mutant for various time intervals. Lane M, M.W. marker (94, 67, 43, 30, and 20.1 kDa). Lanes 1–5 are the proSPE B C47S mutant incubated at 37 °C in the presence of active enzymes. Lane 6 is the active enzyme. Arrows designate the cleavage products of the reaction.
proSPE B interacted with the active-site residues in mSPE B. For example, NOEs between side chain NH of W214 and the protons from the C-terminal loop residues, such as Hβ of A231, Hβ and H8 of L232, and Hβ of T234, were identified. Additional NOEs were also found between NH of A238 and H8 of K191, between NH of A238 and Hβ of V192, and between NH of A238 and NH of K191 (Fig. 2). A total of 4744 experimentally derived restraints, including 4504 NOEs, 137 hydrogen bonds, and 240 dihedral angles, were used for structure calculation. An average of 18.8 restraints per residue was identified (Table 2). NMR analysis revealed that mSPE B C47 mutant were 1.01 Å, indicating that no significant difference was found between NMR and x-ray structures resulted in an R.M.S.D. value of 0.96 Å, and 0.49 Å, respectively. According to our Ramachandran analysis, all dihedral angles of mSPE B C47 mutant were in the allowed region. A summary of the restraint and structural statistics was presented in Table 2. NMR analysis revealed that mSPE B C47 mutant and its E-64 complex was the H195 residue from mSPE B NMR structure rotates from a catalytically competent position (20). In this study, the H195 residue from mSPE B NMR structure rotates from a catalytically incompetent position in the proSPE B to a catalytically competent position (Fig. 3D). The rotation of the side chain of H195 was found from the interactions of H195 with F197, G213, and W214, which were identified by the NOEs between He1 of H195 and NH of W214 (Fig. 2), between He2 of H195 and He1 of W214, between He1 of H195 and Ha of G213, and between He1 of H195 and H81 of F197. Additional NOEs were also found between H81 of F197 and NH of G213 and between H81 of F197 and NH of G213 (data not shown). These results were consistent with that of NMR chemical shift perturbation, and the only shifted residue of seven histidine residues between mSPE B C47S mutant and its E-64 complex was the H195 residue (Fig. 4A).

Chemical shift perturbation was used to compare structural differences between mSPE B C47S mutant and its E-64 complex. A superimposition of their two-dimensional 1H-15N HSQC spectra was shown in Fig. 4B. Seven regions, including Y16-G18, T45-M56, S134-G136, Q187-G190, G194-F197, W212-W214, and L227-Q245, exhibited the values of chemical shift perturbation greater than 0.5 (Fig. 4C). This result was consistent with the reported active site of SPE B that is involved in Q17, C47, H195, W212, and W214. To confirm the results, we expressed G136A and V189A mutants, whose residues have the largest perturbation, and determined their protease activity.

Protein Analysis

The three-dimensional structure and backbone dynamics of SPE B were determined by NMR, and the backbone dynamics were compared with those of the protease domain of the proenzyme. The structural comparison of SPE B with the protease domain of the proenzyme is shown in Fig. 2. A stereo-view of the 20 best structures from 100 initial structures are shown. The short and long range NOEs are shown on the left and right, respectively.

Three-dimensional Structure and Backbone Dynamics of SPE B

![Figure 2](image-url)
Three-dimensional Structure and Backbone Dynamics of SPE B

The G136A and V189A mutants exhibited 15.3–46.7-fold and 131–420-fold less activity, respectively (Table 1). Similar to the processing patterns identified from the mutation on the residues near the active site, only two final products were found from the proSPE B C47S mutant by G136A and V189A mutants (Fig. 1, D and E). These findings suggested that the S134-G136 and Q187-G190 regions also may be involved in the binding of inhibitor and substrate.

Backbone Dynamics Determination of SPE B and Its E-64 Complex—$^1$H,$^{15}$N correlated NMR spectroscopy was used to measure $^{15}$N R$_1$, $^{15}$N R$_2$, and $^1$H--$^{15}$N NOE parameters of mSPE B C47S mutant and its E-64 complex (Fig. 5). The square of the generalized order parameter ($S^2$), the effective internal correlation time ($\tau_m$), and a conformational exchange broadening parameter (Rex) for each backbone amide NH vector were determined using the model-free formalism (Fig. 6). The optimized values of $\tau_m$ for mSPE B C47S mutant and its E-64 inhibitor were determined to be 13.9 and 15.2 ns, respectively. The obtained diffusion tensors of mSPE B C47S mutant and its E-64 complex were symmetric with $D_\parallel/D_\perp = 1.19$ and 1.18, respectively. The isotropic, axial, and anisotropic models were used to fit the diffusion tensor, and the isotropic model yielded the best results. These results were consistent with our structural analysis that the tertiary fold of mSPE B C47S mutant and its E-64 inhibitor had a symmetric shape.

Comparison of Dynamical Properties of mSPE B and Its E-64 Complex—All three relaxation parameters of mSPE B and its E-64 complex were very similar throughout the sequence. However, they exhibited lower NOE values in the K191-G194 and A231-G240 regions, and higher R1 value and lower R2 value in the A231-G240 region (Fig. 5). A summary of the average NOE, R$_1$, R$_2$, and $S^2$ values of all, the K191-G194, and A231-G240 regions are presented in Table 3. The only observable differences between mSPE B and its E-64 complex were found in the K191-G194 and A231-G240 regions. The average NOE values of mSPE B and its E-64 complex were 0.84 and 0.82, respectively. In contrast, the average NOE values of the K191-G194 region for mSPE B and its E-64 complex were 0.52 and 0.34, respectively. The average NOE values of the A231-G240 region for mSPE B and its E-64 complex were 0.60 and 0.43, respectively. The only negative NOE value was found in the G193 residue of the SPE B/E-64 complex. Upon binding to the E64 inhibitor, the differences in the average NOE values of the K191-G194 and A231-G240 regions were 0.18 and 0.17, respectively. The average R1 and R2 values of the K191-G194 region of mSPE B were very similar to those of the rest of molecule. In contrast, the average R1 and R2 values of the A231-G240 region of the SPE B/E-64 complex were 1.48 ± 0.07 and 13.43 ± 0.79 s$^{-1}$, which are 32% higher and 12% lower than those of mSPE B.

The optimized values of $S^2$, $\tau_m$, and Rex for mSPE B and its E-64 complex are shown in Fig. 6. The average $S^2$ values of mSPE B and its E-64 complex were 0.95 and 0.94, respectively (Table 3). The average $S^2$ values of the K191-G194 region of mSPE B and its E-64 complex were the same with a value of 0.79. The average $S^2$ values of the A231-G240 region of mSPE B and its E-64 complex were 0.8 and 0.55, respectively. They were lower than the average $S^2$ value of 0.95. The order parameter variation of the dynamics associated with the E-64 inhibitor can be seen more clearly in the sausage plot shown in Fig. 7. Upon binding of the inhibitor, the A231-G240 region of SPE B became more flexible. Most of the residues of mSPE B and its E-64 complex exhibited extensive flexibility on slow conformational exchange motion on the $\mu$s/$ms$ time scale. Interestingly, the fast motion on the ps/ns time scale was mainly found in the K191-G194 and A231-G240 regions. Our results showed that SPE B became more flexible and exhibited fast motion in the K191-G194 and A231-G240 regions upon binding to the E-64 inhibitor.
FIGURE 4. NMR analysis of C47S mutant and the E-64 complex of mSPE B. Superimposition of the $^1$H-$^1$N HSQC spectra for (A) $[\alpha$-$^1$N]His and (B) $^1$N-labeled C47S mutant (black) and the E-64 complex (red) of mSPE B. The residues with chemical shift perturbation greater than 0.5 are shown. They are labeled according to the residue types and numbers. C, chemical shift difference between C47S mutant and the E-64 complex of mSPE B. The chemical shift differences are calculated according to the equation: $\Delta \delta = [0.5 \times (\Delta^{(1)}H)^2 + 0.2 \times (\Delta^{(15)}N)^2]^{1/2}$. 

Three-dimensional Structure and Backbone Dynamics of SPE B
DISCUSSION

It is known that SPE B plays an important role in GAS infection, and mature SPE B is the active form of SPE B under physiological conditions (5–19). To understand the molecular basis of SPE B in GAS pathogenesis, we determined three-dimensional structure and backbone dynamics of the mature form of SPE B.
We found that the C-terminal loop of mSPE B, which cannot be observed in the x-ray structure of proSPE B, interacts with the active site residues. The contacts are predominantly hydrophobic. This is very similar to the interactions between prodomain and the active site residues. Thus, the 87p-90p residues of the prodomain insert into the active site close to W214.

Superimposition of the protease domain of proSPE B and mSPE B showed no differences in their secondary structures. However, main differences were found from the conformation of the R223-Q245 loop and the orientation of the catalytic H195 residue (Fig. 3). In contrast to three-dimensional structure of proSPE B, structural analysis of mSPE B showed that the H195 residue is located at catalytically competent position. Structural comparison of proSPE B and mSPE B reveals that an inactivation mechanism may be involved in the displacement of the catalytic H195 residue by the Asn89p residue of the prodomain. Thus, the side chain of the Asn89p residue from an α-helix of the prodomain inserts into the active site cleft, at a position that corresponds to the catalytically competent position of the H195 residue. Therefore, the proteolytic removal of the proregion would allow the side chain of a catalytic histidine to rotate into a competent position so that it can interact with the catalytic cysteine residue. A similar activation mechanism involving in a torsional rotation of the catalytic histidine is also found from the interpain A cysteine protease (InpA) of Prevotella intermedia (42). The competent position of the catalytic histidine residue is displaced by the S88p residue of an α-helix of prodomain and is trapped by a zymogenic hairpin and a latency flap from the protease domain. Sequence alignment between SPE B and InpA showed that they contain a deletion and insertion in these regions (42). In SPE B, the zymogenic hairpin region does not exist and the latency flap region contains the additional residues L232-G237. These results suggested that the activation and inactivation of SPE B and InpA are involved in a rotation of the side chain and a different trapping mechanism for the catalytic histidine residue, which are distinct from other families of cysteine proteases (43, 44).

The information on substrate-binding sites should be defined by not only proSPE B but also mSPE B structures (20). In our previous study we found that the substrate specificity of SPE B was similar to the substrate preference of the papain-like family, with preference for a hydrophobic residue (isoleucine [5/9], tyrosine [2/9], methionine [1/9], or valine [1/9]) at the P2 site. The analysis of the cleavage sites also reveal trends, with a asparagine residue (3/9) at the P3 site, a lysine residue (3/9) at the P1 site, a glycine residue (3/9) at the P2’ site, a glycine or alanine residue (2/9) at the P3’ site, and a glutamate residue (3/9) at the P4’ site (10). To identify possible substrate binding sites of SPE B, we mapped the amide nitrogen and hydrogen chemical shift changes for each residue upon the inhibitor E-64 binding. Chemical shift perturbation studies showed that seven regions, including Y15-G18, T45-A51, S135-S141, Q187-G190, orange, model 4; red, model 5; blue, proline residues; black, resonances that were not used to deduce the dynamic information because of weak intensities or multiple decay or an inability to fit them satisfactorily to any model. These images were generated using the MOLMOL program.
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involved in the active site of SPE B, the processing of the 40-kDa SPE B C47S mutant by G136A, V189A, W212A, W214A, and G239D mutants gave rise to only two visual intermediates. In addition to the information about the active site provided by the x-ray structure, this result suggests that the S134-G136, G188-F197, and A231-A246 regions also may be involved in the binding of inhibitor and substrate.

It is known that enzymes behave as dynamic machines, and the motions on various time scales play important roles in promoting enzyme catalysis (45). The connection between structure and dynamics is essential to the functions of many enzymes (45–48). NMR spectroscopy is unique in its ability to characterize protein motions over a wide range of biologically relevant time scales with atomic resolution (46). NMR dynamics analysis showed that most regions of SPE B and its E-64 inhibitor complex are fairly rigid, except for the catalytic and C-terminal loops with low order parameter values of 0.5–0.8. It is consistent with many studies, showing that the flexible catalytic loops are related to enzyme catalysis and the substrate-binding regions are flexible (45–48). Most residues of mSPE B and its E-64 complex throughout the molecule exhibit extensive regions are flexible (45–48). Most residues of mSPE B and its E-64 inhibitor complex are fairly rigid, except for the catalytic and C-terminal loops with low order parameter values of 0.5–0.8. It is consistent with many studies, showing that the flexible catalytic loops are related to enzyme catalysis and the substrate-binding regions are flexible (45–48). Most residues of mSPE B and its E-64 complex throughout the molecule exhibit extensive flexibility on slow conformational exchange motion on the ps/ns time scale. It is known that slow motion on the ps/ns time scale is usually related to enzyme catalysis (45–48). Interestingly, the fast motion on the ps/ns time scale was found mainly from the K191-G194 and A231-G240 regions. SPE B became more flexible and exhibited fast motion in the K191-G194 and A231-G240 regions upon binding to the E-64 inhibitor. Our functional analysis also showed that the G239 residue within the C-terminal loop plays an important role for its protease activity. These findings suggested that the activation and inactivation of SPE B are involved in a rotation of the side chain of the catalytic histidine residue, which is distinct from other families of cysteine proteases. This study may serve as the basis for gaining insight into streptococcal infections by exploring the structure and function relationships of SPE B. This study also extends our understanding of the molecular basis of SPE B under physiological conditions.

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