The Streptococcal Cysteine Protease SpeB Is Not a Natural Immunoglobulin-Cleaving Enzyme

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The human bacterial pathogen *Streptococcus pyogenes* has developed a broad variety of virulence mechanisms to evade the actions of the host immune defense. One of the best-characterized factors of the streptococcal cysteine protease SpeB, an important multifunctional protease that contributes to group A streptococcal pathogenesis in vivo. Among many suggested activities, SpeB has been described to degrade various human plasma proteins, including immunoglobulins (Igs). In this study, we show that SpeB has no Ig-cleaving activity under physiological conditions and that only Igs in a reduced state, i.e., semimonomeric molecules, are cleaved and degraded by SpeB. Since reducing conditions outside eukaryotic cells have to be considered nonphysiological and IgG in a reduced state lacks biological effector functions, we conclude that SpeB does not contribute to *S. pyogenes* virulence through the proteolytic degradation of Igs.

*Streptococcus pyogenes*, also known as group A *Streptococcus*, is the causative agent of a wide variety of mild infections, like pharyngitis and impetigo, but also the cause of potentially life-threatening diseases, like streptococcal toxic shock syndrome and necrotizing fasciitis (1). *S. pyogenes* is a strictly human pathogen and has evolved a range of surface-attached and secreted virulence factors that allow the pathogen to survive and proliferate in its human host (1, 2). Among secreted proteins, the streptococcal cysteine proteinase SpeB is the most-prominent and probably best-characterized bacterial virulence factor. The vast amount of literature on SpeB transcription, secretion, maturation, and activation has recently been summarized in an excellent review (3). In an overview, SpeB is secreted as an inactive 40-kDa zymogen that is converted into a mature 28-kDa active protease either by autocatalytic cleavage or aided by host or streptococcal proteases (4–7). Regulation of speB transcription and SpeB expression is complex and involves environmental factors, transcriptional and posttranscriptional regulation, and posttranslational regulation, and posttranslational regulation, brought about by an endogenous inhibitor encoded on the streptococcal chromosome (for details, see reference 3 and references therein). Mature SpeB (mSpeB) exhibits promiscuous protease activity toward a large variety of both bacterial and human proteins, including vitronectin, fibronectin (8), fibrinogen (9), and immunoglobulins (10–12). Further examples include the activation of interleukin-1β (13), the matrix metalloproteinase MMP-2 (14), and the release of active proinflammatory kinins from H-kininogen (15). SpeB also targets numerous streptococcal proteins, like M1 protein, protein H, C5a peptidase (4, 16), and EndoS (17). Furthermore, analysis of the secreted streptococcal proteome suggests that SpeB can degrade the majority of abundant secreted streptococcal proteins in vitro (18). It has been suggested that hydrolysis of other streptococcal virulence factors by SpeB leads to less-severe disease progression (18, 19). In fact, SpeB expression is inversely correlated to severity of infection, i.e., the majority of streptococcal isolates from cases with severe invasive infections exhibit low or no SpeB expression (19). A comprehensive summary of target proteins for SpeB has recently been presented in another excellent review (20). Despite experimental variations, and partly conflicting data, there is no doubt that SpeB is an important virulence factor and contributes to group A streptococcal pathogenesis in vivo (20).

A strategy crucial for *S. pyogenes* survival and proliferation is the ability to evade adaptive immune responses, in particular, the recognition by specific, opsonizing antibodies. Specific antibodies mediate activation of phagocytic cells and the complement response. A common mechanism to avoid the detrimental effects of specific antibodies is to degrade immunoglobulins (Igs) by specific IgG or IgA proteases. *S. pyogenes* has been reported to secrete two IgG-degrading enzymes: IdeS (IgG-degrading enzyme of *S. pyogenes*), which is a highly efficient IgG-specific endopeptidase (21), and SpeB, which has been reported to degrade IgA, IgM, IgE, and IgD and to cleave IgG in the hinge region (10–12). However, a role for SpeB as part of the first line of defense against specific antibodies has been questioned, since SpeB is expressed during stationary growth (22), i.e., when infections are already established (23), and since, in contrast to IdeS, IgG proteolysis by SpeB appears to occur slowly over 24 to 48 h (10–12).

As a cysteine protease, SpeB activity is dependent on the reduction of the catalytic-site cysteine (4). In experimental settings, activation of SpeB is usually accomplished by incubation of SpeB with reducing compounds, e.g., dithiothreitol (DTT), β-mercaptoethanol, or l-cysteine. Furthermore, reducing compounds are commonly added in assay buffers or growth medium to maintain SpeB activity (5, 10–12, 17, 24–26). However, the inflammatory state that typically accompanies streptococcal infections leads to infiltration of activated neutrophils and the subsequent release of reactive oxygen species (ROS). Thus, an inflammatory environment has to be considered a rather oxidative environment, imply-
ing that in vitro analyses of SpeB activity in the presence of reducing agents might not be representative of physiological environments (20). In this study, the proteolytic cleavage of immunoglobulins by SpeB under nonreducing conditions was investigated. We demonstrate that in order to cleave and degrade the heavy chains of Ig, SpeB requires Ig in a reduced state, i.e., in a semimonomeric form in which the molecule lacks intact disulfide bonds and is held together only by noncovalent binding forces in the CH1 region (27). We therefore conclude that SpeB is not contributing to IgG cleavage under physiological conditions and that the contribution of SpeB to S. pyogenes virulence is not due to the proteolysis of immunoglobulins. Analyses of SpeB activity in physiological environments revealed that SpeB is not oxidized in the presence of human plasma, due to the antioxidant activity of human serum albumin, and therefore retains activity also in the presence of activated neutrophils.

MATERIALS AND METHODS

**Proteins.** Fibrinogen, fibronectin, human serum albumin (HSA), immunoglobulins, and vitronectin were all purchased from Sigma-Aldrich.

**Purification of SpeB.** For purification of mSpeB, the S. pyogenes strain 544B was grown for approximately 16 h in Todd-Hewitt broth (BD Biosciences) in 5% (vol/vol) CO2 at 37°C. The bacteria were collected by centrifugation (3,800 × g for 10 min at 4°C), and culture supernatant was subjected to ammonium sulfate precipitation (50 to 80% [wt/vol]). Precipitated proteins were dissolved in 1× phosphate-buffered saline (PBS) buffer. After dialysis against 20 mM sodium acetate buffer (pH 5.0), protein samples were sterile filtered, diluted in 20 mM sodium acetate buffer (pH 5.0), and applied to a HiTrap SP FF anion exchange column (GE Healthcare) equilibrated in the same buffer. Proteins were eluted in a gradient of 0 to 2 M NaCl over 20 column volumes at a flow rate of 1 ml/min, and SpeB starts to elute at 0.6 M NaCl. Eluted protein fractions were dialyzed overnight at 4°C against 1× PBS. Protein purity and identity were assayed by SDS-PAGE and Western blot. The amount of active SpeB was determined by active-site titration using various amounts of the substrate.

**SpeB activity assays.** SpeB activity was measured as previously described (29). Briefly, purified SpeB (0.1 mg/ml) was incubated with 2 mM diithiothreitol (DTT) for 30 min at 37°C to reduce the active-site cysteine and activate the enzyme. DTT was removed by using Zeba spin desalting columns, with a 7-kDa molecular mass cutoff, according to the manufacturer’s instructions (Thermo Scientific). Sixty microliters of the synthetic substrate 4-nitroanilide. Phenylalanine-serine-alanine-arginine-hydrochloride (BPFA) (Sigma) was mixed with 90 μl of 0.1 M phosphate buffer (pH 6.0) and added to activated SpeB. For determination of SpeB activity in human plasma or in the presence of human serum albumin (HSA), SpeB was incubated with plasma or increasing concentrations of HSA for 0 to 180 min prior to the addition of BPFA. Changes in absorbance were determined at an optical density of 405 nm (OD405) after 70 min of incubation at room temperature. All assays were performed in duplicate or triplicate.

**SpeB cleavage of IgG.** For IgG cleavage assays with SpeB, 0.025 mg/ml activated SpeB (after removal of DTT) was incubated with 1 mg/ml IgG (reduced or nonreduced) at 37°C for various time periods. Reactions were stopped by addition of reduced loading buffer and incubation at 96°C for 10 min. To obtain reduced IgG substrate, 10 mg/ml IgG in PBS was incubated overnight at 37°C in the presence of 10 mM DTT. DTT was removed by buffer exchange using Zeba spin columns with a 7-kDa molecular mass cutoff.

**SpeB cleavage of plasma proteins.** Purified SpeB (0.1 mg/ml) was activated by addition of 2 mM DTT. Different concentrations of activated, purified SpeB, with and without DTT, were incubated with 3 μg of fibrinogen, fibronectin, IgG, IgM, IgA, or vitronectin at 37°C in PBS for 1 h to 4 h. The reaction was terminated by the addition of an equal volume of SDS-PAGE sample buffer, followed by incubation at 95°C for 10 min. Proteins were separated and analyzed by standard SDS-PAGE and stained with Coomassie blue (R-250) (USB chemicals).

**SpeB activity in plasma and whole human blood.** SpeB activity against IgG in human plasma or whole blood was determined by a Western blot. Purified SpeB (3.5 μg) was incubated in 1 ml human blood lacking specific antibodies against SpeB or IdeS at 37°C under rotation. As controls, human blood and human blood supplemented with recombinant IdeS (21) were used. Samples were diluted 1/100, separated by 8% SDS-PAGE, and transferred to an Immobilon-P polyvinylidene difluoride (PVDF) membrane (Millipore) using a semidyed electrohoresis transfer cell (Bio-Rad). Goat anti-human IgG horseradish peroxidase (HRP)-conjugated antibodies (Bio-Rad) were used for detection of IgG. Immunoreactive proteins were detected using an ECL Plus Western blotting detection system (GE Amersham Biosciences) according to the manufacturer’s instructions.

**SpeB activity in the presence of human polymorphonuclear leukocytes and measurement of ROS.** Human polymorphonuclear leukocytes (PMNs) were isolated from heparinized blood using polymorphon-prep (Nykomed Pharma, Norway) as previously described (30). Briefly, whole blood was layered onto polymorphon prep medium and centrifuged at 400 × g for 30 min. After centrifugation, the neutrophil layer was isolated and washed in PBS and residual erythrocytes were removed by hypotonic lysis in water. Neutrophils were collected by centrifugation, resuspended in 1× PBS, and counted using a counting chamber. For assaying SpeB activity in the presence of activated neutrophils, PMNs were stimulated with phorbol-12-myristate-13-acetate (PMA) at a final concentration of 0.8 μM in the presence of SpeB and BPFA. Generation of extracellular reactive oxygen species (ROS) was measured as chemiluminescence using isoluminol (0.04 mM) (Sigma) and horseradish peroxidase (2.4 units) (Sigma). Chemiluminescence was detected using an Infinite M 200 plate reader instrument (Tecan). SpeB activity against BPFA was measured simultaneously as described previously. For measurements of ROS and SpeB activity in the presence of HSA, increasing amounts of HSA (0 to 15 g/liter) were added to the reactions.

**RESULTS**

**SpeB activity under nonreducing conditions.** The proteolytic activity of activated SpeB in the absence of reducing agents was investigated. SpeB preparations were activated with 2 mM DTT at 37°C. DTT was removed by repeated buffer exchange, and SpeB activity against the synthetic substrate n-benzoyl-proline-phenylalanine-arginine-p-nitroanilide hydrochloride (BPFA) was monitored for up to 5 h in assay buffer and compared to that in similar reaction mixtures containing DTT. In a buffer system and in the absence of reducing compounds, SpeB activity decreases slowly until precipitously dropping after 30 min to baseline levels (Fig. 1A, open squares). In a reducing buffer, however, SpeB activity remains practically unchanged over at least 3 h (Fig. 1A, filled squares). Oxidation and inactivation of the cysteine protease occur much faster at elevated temperatures, as SpeB remains active for at least 3 h at room temperature, also in the absence of reducing compounds (data not shown). Under these experimental conditions, reincubation of oxidized SpeB in the presence of DTT results in less than half of the initial enzyme activity (Fig. 1B), indicating that the oxidation state of the enzyme is not easily reversed.

**SpeB activity in human plasma and in the presence of activated neutrophils.** In the next set of experiments, assay buffer was replaced with human plasma. Activated SpeB preparations were incubated in human plasma with or without DTT at 37°C, and SpeB activity against BPFA was monitored for 4 h. Plasma contains endogenous BPFA-hydrolyzing activity, as evident by back-
ground absorbance readings, but SpeB-mediated BPFA hydrolysis in human plasma clearly remains stable over time independently of whether samples were supplemented with DTT or not (Fig. 2A).

During streptococcal infection, migrating activated neutrophils will generate and degranulate reactive oxygen species (ROS) and create an oxidative environment at the infection site. SpeB activity in the presence of neutrophils was monitored, and SpeB protease activity is significantly inhibited in response to extracellular ROS, most likely due to rapid oxidation of the catalytic-site cysteine (Fig. 2B). However, ROS-associated cell and tissue damage is generally prevented by circulating antioxidants. Most free-radical-quenching activities in serum have been assigned to human serum albumin (HSA), one of the most abundant plasma proteins (31) with multiple antioxidant properties (32). HSA concentrations are elevated during inflammation (33), and therefore, the amount of extracellular ROS and SpeB activity in the presence of HSA was investigated (Fig. 3). HSA concentrations as low as approximately 10% of HSA serum concentrations were capable of efficiently trapping ROS (Fig. 3, open diamonds), and consequently, SpeB activity increased in correlation to the HSA concentration (Fig. 3, filled diamonds). Thus, in the presence of physiological HSA concentrations, SpeB remains active, also in an inflammatory environment, and there is no apparent requirement for reducing conditions to sustain enzymatic activity.

**SpeB is not an immunoglobulin-degrading protease.** Immunoglobulins have previously been described as targets for SpeB proteolytic activity (10–12). In vitro treatment of specific IgG with SpeB was shown to interfere with immunoglobulin-mediated phagocytosis (25), and SpeB was suggested to contribute to survival of *S. pyogenes* in human blood by preferentially cleaving antigen-bound IgG (12). Common to all studies is that the proteolytic assay was performed under reducing conditions using incubation times ranging from 24 h to 48 h to accomplish cleavage of Igs. SpeB proteolytic activity against IgG, IgM, and IgA was reassessed with activated SpeB under reducing and nonreducing conditions. Clearly, while all three immunoglobulin types were efficiently cleaved under reducing conditions, incubation with SpeB under nonreducing conditions did not affect the integrity of IgG, IgM, and IgA.
the immunoglobulins (Fig. 4A to C). The lack of cleavage of IgG and IgM under nonreducing conditions was confirmed by Western blotting with IgG- and IgM-specific antibodies (data not shown). DTT present in experimental assay reactions not only ensures activation of SpeB but also reduces disulfide bridges in Igs, and we hypothesized that reduced, i.e., semimonomeric, IgG, but not intact IgG, is a substrate for SpeB. The IgG cleavage assay was repeated using either native IgG or IgG that had been reduced by DTT (followed by buffer exchange) as the substrate (Fig. 4D). SpeB readily and efficiently cleaved semimonomeric IgG already after 3 min of incubation, while nonreduced IgG remained intact also after 90 min of incubation at 37°C (Fig. 4D). In addition, no cleavage was detected when IgG was allowed to reoxidize prior to the proteolytic assay with activated SpeB, confirming that SpeB activity against IgG is dependent on separated Ig chains (data not shown).

Several other plasma proteins, including fibrinogen (9), fibronectin, and vitronectin (8), have previously been shown to be substrates for SpeB. Degradation of fibrinogen has been demonstrated to occur under nonreducing conditions (9). Cleavage of fibronectin and vitronectin has also previously been assayed in the absence of reducing compounds in the assay buffer, but it was not evident whether reducing compounds had been removed from the SpeB preparation prior to incubation with the plasma proteins (8). However, fast and efficient cleavage of fibrinogen, vitronectin, and fibronectin under either reducing or nonreducing conditions was confirmed (data not shown).

SpeB has no immunoglobulin-degrading activity in human plasma or blood. Initial experiments demonstrated that SpeB enzymatic activity in the absence of reducing compound dropped after 1 h of incubation in assay buffer to baseline levels (Fig. 1A), while activity in plasma samples was present for at least 4 h (Fig.

**FIG 3** ROS production and SpeB activity in the presence of increasing concentrations of HSA. Detectable extracellular ROS (○) are trapped by HSA, allowing SpeB activity (●) to increase under the same conditions. Negative controls are PMNs alone (□) and PMNs and SpeB (■).
The role of SpeB for streptococcal pathogenesis has been thoroughly investigated, and the list of reported substrates for this important virulence factor is growing considerably, reflecting the indiscriminant substrate recognition properties of the enzyme. However, does the fact that SpeB can hydrolyze a substrate in vitro necessarily mean that the enzyme will do so in vivo? Nelson and coworkers stated that a lot of controversy about SpeB function originates from the finding that reported activities would counteract each other. Historically, the identification of novel substrates has been a major objective in order to understand the role of SpeB for streptococcal virulence, and findings have not always been set in a physiological context. This is, in particular, evident for the reports of immunoglobulin-degrading activity of SpeB. Immunoglobulins play a key role in adaptive immune responses, and S. pyogenes cells recognized by specific IgG antibodies are rapidly eliminated. Therefore, the identification of Ig proteolytic activity of SpeB has been seen as a matter of considerable interest. However, a role of SpeB as part of the first-line defense against specific antibodies can also be questioned. SpeB transcripts have certainly been detected at 24 h postinfection in a murine infection model, but for an Ig protease to contribute to protection against opsonizing IgG, it would be essential that the enzyme is instantly secreted to be able to act on circulating specific antibodies. Furthermore, proteases acting against IgG have to be highly effective and should be specific enough to avoid that other, redundant substrates occupy the enzyme and affect the cleavage of specific Ig.

In previous studies, IgG proteolysis by SpeB was carried out for 24 to 48 h to achieve cleavage or degradation of Ig. In the presence of opsonizing antibodies, IgG-mediated phagocytosis and bacterial killing occur within less than 15 min, and a time frame of 24 to 48 h to achieve inactivation of IgG does not represent a protective biological function. In addition, and most importantly, all proteolytic assays were performed under reducing conditions, and it has, in fact, been noted that such conditions are necessary to achieve cleavage of IgG. Although reducing conditions certainly sustain SpeB enzymatic activity, a reducing environment will also lead to disruption of disulfide bonds of immunoglobulins, thereby destroying Ig integrity and creating semimonomeric molecules. An infection site is considered an oxidative environment, and it is highly unlikely that SpeB will encounter semimonomeric IgG in vivo. The presence of reducing compounds in isolated microenvironments, e.g., sulfide in gingival pockets during periodontal disease, has been described. However, gingival pockets are not a natural habitat for S. pyogenes, and infiltrating neutrophils will rapidly oxidize and detoxify sulfide. In fact, even in the reducing environment in gingival pockets, sulfide concentrations are not sufficient to achieve reduction of the disulfide bridges of IgG.

In light of our findings of distinct differences in SpeB activity against immunoglobulins under reducing or nonreducing conditions, we also investigated SpeB activity against several plasma proteins that previously have been reported to be substrates for the streptococcal protease and that all contain inter- and intramolecular disulfide bonds. We corroborated the earlier findings that fibrinogen, fibronectin, and vitronectin are substrates for SpeB under both reducing and nonreducing conditions (data not shown).

In the current study, we demonstrate that SpeB is not a natural Ig protease under physiological conditions. We also show that SpeB, once activated, retains its activity in a plasma environment in the absence of supplemented reducing compounds. In contrast, SpeB is oxidized and rapidly inactivated in the presence of activated neutrophils, but this mechanism can, at least initially, be counteracted by HSA. Notably, in the presence of SpeB protein, i.e., independently of enzymatic activity, the amount of extracellular ROS is clearly diminished compared to that of the control, an observation also reported for IdeS, another streptococcal cysteine protease.

Immunoglobulins are among the most abundant human plasma proteins. In light of the numerous important functions of SpeB and considering that Igs as redundant substrates are likely to occupy the enzyme and affect proteolytic activity against other substrates, the removal of Ig from the list of SpeB substrates some-
what eases the task to understand the role of SpeB during streptococcal infection. However, although SpeB is one of the most investigated prokaryotic virulence factors, investigations of SpeB function are far from being completed and future studies will certainly contribute to further understanding.

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REFERENCES

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