Staphylococcal Enterotoxin C Subtypes Are Differentially Associated with Human Infections and Immunobiological Activities

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ABSTRACT Staphylococcus aureus causes significant infections, responsible for toxic shock syndrome (TSS), hemorrhagic pneumonia, and many other infections. S. aureus secretes virulence factors, which include superantigens such as staphylococcal enterotoxins (SEs). We examined differences in immunobiological activities and disease associations among the four human SEC subtypes. We sequenced the sec gene from 35 human isolates to determine SEC subtypes. Upon finding differences in disease association, we used a [3H]thymidine uptake assay to examine SEC-induced superantigenicity. We also employed a rabbit model of SEC-induced TSS. SEC-2 and SEC-3 were associated with menstrual TSS and vaginal isolates from healthy women, whereas SEC-4 was produced by USA400 isolates causing purpura fulminans and hemorrhagic pneumonia. SEC subtypes differed in potency in a TSS rabbit model and in superantigenicity. There was no difference in superantigenicity when tested on human peripheral blood mononuclear cells. Despite differences, all SECs reacted with polyclonal antibodies raised against the other SEC subtypes. The associations of SEC subtypes with different infections suggest that S. aureus produces virulence factors according to host niches.

IMPORTANCE Staphylococcal enterotoxin C has four subtypes that cause human diseases, designated SEC-1 to -4. This study shows that SEC-2 and SEC-3 are the most toxic subtypes in a rabbit model and are associated with human vaginal infections or colonization in association with another superantigen, toxic shock syndrome toxin 1. SEC-4 is associated with purpura fulminans and hemorrhagic pneumonia. SEC-1 is uncommon. The data suggest that there is some selective pressure for the SEC subtypes to be associated with certain human niches.

KEYWORDS Staphylococcus aureus, enterotoxin, superantigen, toxic shock syndrome

Although Staphylococcus aureus can exist as a benign constituent of the human microbiome, it is capable of causing serious diseases such as toxic shock syndrome (TSS), purpura fulminans, and hemorrhagic pneumonia (1–3). The Centers for Disease Control and Prevention previously reported that S. aureus is a highly significant cause of serious infectious diseases in the United States (4). Its numerous secreted exotoxins contribute to the ability of S. aureus to cause pathology in the host. A group of these exotoxins, the superantigens (SAgs), is known to induce massive T-cell proliferation, leading to large amounts of T-cell and macrophage cytokine production (1–3). This resultant “cytokine storm” induces fever, hypotension/shock, and rash, possibly resulting in death. SAg interaction with major histocompatibility complex (MHC) class II molecules on antigen-presenting cells is nonspecific and does not require SAg processing prior to interaction with T lymphocytes through the variable part of the β-chain of the T cell receptor (Vβ-TCR). SAgs stimulate ~10,000-fold more T lymphocytes than typical antigens (1–3, 5, 6).
A. Signal Peptides

SEC1  M N K S R F I S C V I L I A L L I L V L F T P N V L A
SEC2  M N K S R F I S C V I L I A L L I L V L F T P N V L A
SEC3  M Y K R L F I S R V I L A L L I L V I S T P N V L A
SEC4  M N K S R F I S C V I L I A L L I L V L F T P N V L A

B. Maturity Proteins

SEC1  E S Q P D P T P D E L H K A S K F T G L M E N M K
SEC2  E S Q P D P T P D E L H K S S E F T G T M G N M K
SEC3  E S Q P D P M P D D L H K S S E F T G T M G N M K
SEC4  E S Q P D P T P D E L H K S S E F T G T M G N M K

SEC1  V L Y D D H Y V S A T K V K S V D K F L A H D L I
SEC2  A T K V M S V D K F L A H D L I
SEC3  Y L Y D D H Y V S A T K V K S V D K F L A H D L I
SEC4  Y L Y D D H Y V S A T K V K S V D K F L A H D L I

SEC1  Y N I S D K K L K N Y D K V K T E L L N E G L A K
SEC2  Y N I S D K K L K N Y D K V K T E L L N E D L A K
SEC3  Y N I S D K K L K N Y D K V K T E L L N E D L A K
SEC4  Y N I S D K K L K N Y D K V K T E L L N E D L A K

FIG 1 Alignment of SEC subtypes 1 to 4. (A) Alignment of the signal peptide, which is the first 27 amino acid residues of the entire SEC protein, which are cleaved to yield the mature SEC proteins. Differences from the consensus are in red boldface type. Only SEC-3 has residues in the signal peptide that differ from the others. (B) Alignment of the mature SEC proteins. Differences from the consensus are in red boldface type. Most of the differences between the SEC subtype proteins were in the N-terminal region of the mature proteins.

One subfamily of SAgs produced by S. aureus is the staphylococcal enterotoxins (SEs), SAgs that are responsible for food poisoning and nonmenstrual TSS, including infections that result secondarily to respiratory viral infections, allergies, and asthma (1, 2). SEs characterized thus far include SEA-SEE and SEG; numerous SE-like SAgs have also been described (1, 2, 7).

Among the SEs is SEC, which exists as 4 known human subtypes (SEC-1 through SEC-4); they differ from one another by at most 15 amino acids. Most differing amino acids lie in the signal peptide (Fig. 1A) and the N-terminal region of the mature proteins (Fig. 1B). The C-terminal end is more conserved between subtypes and is suggested to be structurally responsible for major biological activities (8). The molecular masses of all SEC subtypes differ only modestly from one another, falling between 24 and 28 kDa (9).

Baba et al. (10) sequenced the community-associated methicillin-resistant S. aureus (CA-MRSA) strain MW2 and noted that its genome carried a new allelic variant of the SEC gene, sec4. It has been shown that USA400 strains like MW2, most of which are methicillin resistant, are associated with SEC-4 carriage (11, 12). This led us to hypothesize
that the SEC subtypes might each associate with different human infection types based on
differences in immunobiological activities. Our current study uses sequencing to deter-
mine SEC subtypes carried by *S. aureus* isolates causing a variety of human diseases. We
found associations of certain SEC subtypes with particular diseases, prompting us to ascer-
tain whether there are disparities in SEC subtype immunobiological activities. We observed
differences in activities. Currently, the most documented method to neutralize SAgs is in-
travenous immunoglobulin (IVIG). IVIG was able to neutralize all 4 SEC subtypes.

**RESULTS**

**Association of SEC subtypes with *S. aureus*-induced infections.** Recent studies
(10–12) observed that USA400 strains such as MW2 and MNKN, both of which are
MRSAs associated with hemorrhagic pneumonia, produce SEC-4. This raises the possi-
bility that SEC-1 to SEC-3 also associate with certain clonal strains of *S. aureus* and
human infection types.

To determine whether such an association is present among isolates, we sequenced
the sec gene from 35 clinical *S. aureus* isolates, most of which were from patients with
*S. aureus* infections, with a few isolates from the vaginal flora of healthy women.
Interestingly, SEC-1 was the least frequent subtype found among the isolates, observed
in only two strains, one responsible for a case of nonmenstrual TSS and one associated
with atopic dermatitis (Table 1). The most common subtypes encountered were SEC-2
and SEC-3; the majority of *S. aureus* isolates producing these SEC subtypes caused
menstrual TSS, with coproduction of TSS toxin 1 (TSST-1), or they were from the vaginal
flora of healthy patients along with TSST-1. SEC-4 was associated with USA400 *S. aureus
isolates responsible for purpura fulminans, hemorrhagic pneumonia, and mastitis; all
were MRSA. These strains did not produce TSST-1. USA400 strains are likely to cause
lethality in such patients (10, 13). We have never observed USA300 strains to produce
any SEC subtype, and thus, no USA300 strains were examined in this study.

When we compared SEC-2/SEC-3 association with menstrual TSS/healthy-person
vaginal isolates to SEC-4 association with nonmenstrual TSS, hemorrhagic pneumonia,
and other infections (Table 2), the difference in SEC subtype was statistically signifi-
cant (*P* < 0.0001). In contrast to the above-mentioned diseases, atopic dermatitis *S. aureus*
isolates had no particular association with any SEC subtype; all subtypes were found
among these strains. The overall trends demonstrated that SEC-2 and SEC-3 are associ-
ated with vaginal isolates, whether from healthy vaginal microflora or from menstrual
TSS patients, where TSST-1 was coproduced, and SEC-4 was associated with MRSA
USA400 strains primarily causing serious pulmonary *S. aureus* infections.

**SEC subtypes stimulate rabbit splenocyte proliferation.** Because we observed
that certain SEC subtypes were associated with different *S. aureus* infections, we exam-
ined whether there might be differences in the immunobiological properties of the
SEC subtypes. One possible difference between the SEC subtypes would be in superan-
tigenicity, or the ability to stimulate T-cell proliferation. In a 
[^3H]thymidine uptake assay with rabbit splenocytes, we observed that all SEC subtype toxins were able to promote
lymphocyte proliferation in a dose-dependent manner (Fig. 2A). However, proliferation
levels differed among the 4 subtypes; SEC-2 stimulated the lowest levels of lymphocyte
proliferation at most concentrations of toxin. At 0.001 μg/well, SEC-2-induced prolifera-
tion dropped to background levels, whereas this decrease was not observed for other
SEC subtype toxins until 0.00001 μg/well. Strikingly, SEC-3 toxin stimulated the highest
proliferation levels, remaining at 15-fold-higher levels at 0.00001 μg/well, when other
SEC subtypes stimulated only background levels of lymphocyte proliferation. Although
we observed differences in splenocyte proliferation levels induced by each SEC sub-
type, all SEC subtypes demonstrated superantigenicity.

**SEC subtypes 1 to 4 stimulate proliferation of human lymphocytes.** Since we
observed that SEC-1 to -4 induced rabbit splenocyte proliferation and that differences
were present among the proliferation levels stimulated, we investigated the effect of the
SAgs on human lymphocytes. The 
[^3H]thymidine uptake assay performed with pe-
ripheral blood mononuclear cells (PBMCs) demonstrated that the SEC subtypes were
all able to generate proliferative responses, also in dose-dependent manners (Fig. 2B). However, the attenuated superantigenicity of SEC-2 seen with rabbit splenocytes was not observed with human PBMCs. The increased superantigenicity of SEC-3 in comparison to the other subtypes was less apparent in the human PBMC assay, but at all concentrations of SEC, SEC-3-induced PBMC proliferation was slightly higher than those of the other subtypes. Thus, we found that all SEC subtypes induce proliferative responses

### TABLE 2 SEC subtypes 1 to 4 associated with human infections

| Isolate type(s)                                                                 | No. of isolates | SEC-1 | SEC-2 | SEC-3 | SEC-4 | PFGE designation* |
|---------------------------------------------------------------------------------|-----------------|-------|-------|-------|-------|-------------------|
| Menstrual TSS/healthy vaginal microflora                                          |                 | 0     | 7     | 13    | 0     | USA200 TSST-1*    |
| Purpura fulminans, hemorrhagic pneumonia, mastitis                               |                 | 1     | 0     | 0     | 9     | USA400            |
| Atopic dermatitis                                                                |                 | 1     | 2     | 1     | 1     | Mixed             |

*PFGE, pulsed-field gel electrophoresis; TSST-1*, TSST-1 positive.
in human PBMCs, but the differences in superantigenicities were not as apparent as the rabbit splenocyte response to SEC subtypes.

We treated PBMCs with IVIG along with SECs, as IVIG can also neutralize *S. aureus* SAgs (14). IVIG was effective in neutralizing SEC-induced lymphocyte proliferation (Fig. 3). There were small differences in IVIG neutralization abilities, which may reflect small epitope differences among the SECs, which are known to exist.

SEC-3 demonstrates the greatest lethality in a TSS rabbit model. To continue elucidating the basis for SEC subtype disease associations (Table 3), we examined the ability of each SEC subtype to enhance lipopolysaccharide (LPS)-induced lethality in Dutch-belted rabbits. Staphylococcal SAgs are known to synergize with LPS, causing fever, diarrhea, and lethal shock (15). When Dutch-belted rabbits were administered 0.001, 0.01, 0.1, and 10 μg/kg of body weight of SEC with the subsequent injection of 10 μg/kg of LPS, the most lethal SAgs were SEC-3, which yielded the lowest 50% lethal dose (LD₅₀), at 0.001 μg/kg (Table 3), and SEC-2, which exhibited an LD₅₀ of 0.002 μg/kg, while the LD₅₀s for SEC-1 and SEC-4 were 0.01 μg/kg.

**Structural studies of SEC subtypes.** In preliminary studies to hypothesize the reason for differences in the biological activities of SEC subtypes, we mapped the locations of the amino acid differences to assess if these changes are in regions of known biological activities. Figure 1A and B show the location of the amino acid variations in the SEC-1 to -4 subtypes. Figure 4 shows a cartoon protein ribbon image of the structural locations of these variations based on data in the RCSB Protein Data Bank (structure of staphylococcal enterotoxin C2 under accession number 1STE). There are no variations in amino acids in the central diagonal α-helix that appears to control the
interaction with CD40 on epithelial cells (16). SEC-2 has one amino acid difference, which is located in the MHC class II-binding region (Lys39 to Met) (17). It is possible that the increased toxicity of this SEC variant is the result of increased MHC class II binding. The SEC-3 subtype has multiple amino acid differences, and the majority of the differences are in the general location of the Vβ-TCR-binding site (17). It is again presumed that these changes in amino acids increase the interaction with the TCR. Such amino acid differences could account for the increased biological activities of these two SEC subtypes. SEC-1 and -4 have reduced toxicity, and changes are located in regions not shown to affect activity. SEC-1 and -4 thus may have baseline SEC toxicity. We have not attempted mutagenesis of these amino acids individually because of the large number of individual changes needed and multiple LD₅₀s required in rabbits.

### DISCUSSION

In this study, we sought to dissect possible differences among the four human-associated SEC subtypes, examining their relationships to human infections and their ability to stimulate the immune system. In addition to finding no additional human SEC subtypes other than the already described four subtypes, we found two groups of SEC subtype associations with human illnesses: (i) SEC-2/SEC-3 association with menstrual TSS and vaginal isolates from expectedly healthy women and (ii) SEC-4 association with USA400 strains responsible for primarily serious pulmonary illnesses, including purpura fulminans and necrotizing pneumonia. Interestingly, atopic dermatitis S. aureus isolates had no association with any single SEC subtype; all SEC subtypes were found among these strains. These data suggest that persons with damaged skin, as in atopic dermatitis, are susceptible to colonization/infection with nearly any type of S. aureus strain. These skin isolates likely serve as reservoirs that give rise to a myriad of

![FIG 3 IVIG neutralizes the activity of all four SEC subtypes. Human PBMCs were incubated in the presence of 0.01 μg/well SEC and differing concentrations of IVIG. Bars indicate levels of lymphocyte proliferation as the incorporation of (³H)thymidine into DNA in counts per minute (CPM). Both IVIG concentrations neutralized superantigenicity with a P value of <0.01 as determined by Student’s t test.](image)

### TABLE 3 LD₅₀s of SEC subtypes in a rabbit model of TSS

| SEC subtype administered | LD₅₀ (μg/kg) |
|--------------------------|-------------|
| SEC-1                    | 0.01        |
| SEC-2                    | 0.002       |
| SEC-3                    | 0.001       |
| SEC-4                    | 0.01        |
other infections (18, 19). The same has been proposed for mucosal isolates, but the unique association of vaginal mucosal isolates with SEC-2/SEC-3 suggests that these isolates are more restricted in the host niche; the isolates in our study also coproduced TSST-1 that was the cause of menstrual vaginal-associated TSS.

During a skin isolate’s transition to a mucosal surface (i.e., the vaginal surface), transcriptional regulatory mechanisms could alter the expression profile of toxins, surface proteins, and other virulence factors to better adapt the strain to its new environment (19). We know that nearly all menstrual TSS S. aureus isolates are positive for TSST-1. TSST-1 is capable of easily crossing epithelial barriers, whereas SEs are not as able to do so (20). However, coproduction of TSST-1 with SEC-2 or SEC-3 may facilitate SEC penetration of mucosal surfaces. To cause systemic illness in the vaginal environment, the toxin must cross the epithelial barrier. Most atopic dermatitis isolates in this study coproduced TSST-1, and thus, this SAg’s expression could be upregulated once the skin isolate reaches the vaginal mucosa, enabling the strain to cause TSS.

At least two possibilities can explain the disease association with certain SEC subtypes. It is possible that unique activities of some SEC subtypes allow them to facilitate certain infection types. Alternatively, different SEC subtypes could be markers for some other property responsible for the association. The SEC-2 and SEC-3 association with menstrual TSS and vaginal isolates from healthy women indicates that these SEC subtypes might synergize with TSST-1 during TSS or simply accompany strains that produce TSST-1. SEC does not cross epithelial barriers readily, but TSST-1 has this ability when these toxins are administered vaginally in a rabbit model (20). Thus, it is possible that as TSST-1 opens the vaginal mucosal barrier to cause menstrual TSS, SEC-2 or SEC-3 could facilitate disease causation. In our unpublished studies, we have shown that similar doses of multiple superantigens are more toxic to rabbits than single SAgs. Additionally, our laboratory has shown that alpha-toxin can inflict damage on the vaginal epithelial barrier by direct cytotoxicity and inducing localized inflammation, allowing for enhanced penetration of TSST-1 (21) and, thus, possibly of SEC-2 and SEC-3. In the current study, we also observed that when SEC-2 and SEC-3 were administered intravenously (i.v.) to Dutch-belted rabbits, these two subtypes yielded the lowest LD_{50} in the presence of LPS. The enhanced virulence of SEC-2 and SEC-3 seen in this model could support their association with menstrual TSS. One way for S. aureus to inactivate the immune system through SAg production is to select for SAgs with greater activity such that even small amounts may have large effects on the host. We also found that SEC-2 and SEC-3 were associated with vaginal isolates of healthy women; the presence of these isolates in the vaginal microbiome is consistent with these isolates giving rise to menstrual TSS.

**FIG 4** Protein Data Bank three-dimensional (3-D) ribbon model of SEC-2 (accession number 1STE) showing amino acid variations (colored residues). Yellow, amino acid differences from the consensus in SEC-1; magenta, difference from the consensus in SEC-2; blue, differences from the consensus in SEC-3; red, difference from the consensus in SEC-4. The MHC class II-binding domain on SECs includes multiple amino acids on the right side (in the standard view) of the molecules. The Vβ-TCR-binding sites are in a groove at the top of the SECs. The epithelial cell-binding site (CD40) encompasses the central diagonal α-helix with surface amino acids exposed on the back side of the SECs.
As for the SEC-4 association with USA400 isolates, our findings confirm results from Shukla et al. (11). USA400 strains are particularly known for their lethality, and in the study by Baba et al. (10), CA-MRSA USA400 strain MW2 was sequenced to determine genetic reasons for this strain’s increased lethality. Several new virulence genes were found in the genome, including sec4. In the current study, SEC-4 was not the most toxic in activity compared to other SEC subtypes. It stimulated levels of human lymphocyte proliferation similar to those of SEC-1, whereas with rabbit splenocytes, SEC-4 was not as potent as SEC-1. Similarly, the LD50 for SEC-4 and SEC-1 was 0.01 μg/kg in our rabbit model, implying that these SEC subtypes are not as lethal as SEC-2 and SEC-3 in causing TSS, at least in that model. These results suggest that SEC-4 is likely to contribute importantly to human necrotizing pneumonia, as demonstrated by Strandberg et al. (22), but also serves as a marker for USA400 strains. In those necrotizing pneumonia studies, histological evidence revealed hemorrhage and the absence of air spaces in lung tissue when either MW2 (USA400 strain) or purified SEC-4 was administered intratracheally to rabbits. Immunization with SEC-4 prevented the occurrence of respiratory distress and lethality.

Our studies also demonstrated differences among subtypes in superantigenicity as tested with rabbit splenocytes and human PBMCs. All 4 SEC subtypes were superantigenic. In rabbits, SEC-2 was less active than the other 3 SEC subtypes, and SEC-3 was more active in both rabbit and human systems. These differences in superantigenicity do not appear to result from direct amino acid changes altering interactions with TCRs and MHC class II molecules since the observed changes are outside the contact areas. Instead, the amino acid changes likely alter superantigen stability or cause minor structural alterations that result in activity differences.

Finally, IVIG was successful in neutralizing SEC-induced lymphocyte proliferation and has been used in treating SAg-related S. aureus infections (14). It has become a less expensive treatment to neutralize SEC than a few years ago.

**MATERIALS AND METHODS**

**SEC subtype determination in clinical S. aureus isolates.** S. aureus isolates were grown in Todd-Hewitt (TH) broth until stationary phase. Cultures were pelleted (14,000 × g for 5 min) and resuspended in lysis buffer (20 mM Tris–10 mM EDTA, 2.7 mg/ml lysozyme, 1.4 mg/ml lysostaphin, 1 μl per 91 μl Triton X-100). Using the Qiagen (Valencia, CA) DNeasy blood and tissue kit, genomic DNA was purified from cell lysates. The sec gene from each isolate’s DNA was amplified by PCR with sec-specific primers (forward primers GCGTAATTTTGATATTCGCACTTATA, CAAGATGCTTAGAAATCCTCTG, and CCTTGAGAAAGA GTTGTGTATAAAG and reverse primers TTATCCATTCTTTGTTGTAAGGTG, TCACTGTATAATACCGCCTTT TC, and TTATGTCCTCTTTAAATAATCCCGCCTTTTTC, PCR-amplified DNA was sequenced by the University of Minnesota Biomedical Genomics Center.

**SEC toxin purification.** As described previously (20), cultures of S. aureus strains MNDON, FR1361, FR1913, and MW2 (strains expressing SEC-1, -2, -3, and -4, respectively) (Table 1) were grown overnight aerobically in 25 ml of TH broth at 37°C. Cultures were individually diluted 1:120 into dialyzable beef heart medium supplemented with 1% glucose-phosphate buffer, with growth at 37°C until stationary phase (23). Exoproteins in cultures were precipitated with an 80% final concentration of ethanol, and precipitates were collected by centrifugation, resuspended in 75 ml pyrogen-free water, and dialyzed overnight against pyrogen-free water. The dialyzed toxin solution was subjected to isoelectric focusing overnight against pyrogen-free water. The dialyzed toxin solution was subjected to isoelectric focusing twice, in pH gradients of 3.5 to 10. Double immunodiffusion was utilized to identify SEC-containing fractions; reactive fractions were pooled and dialyzed against pyrogen-free water. Pooled fractions were electrophoresed on SDS-PAGE gels to verify purity, and toxins were stored lyophilized until used.

**Isolation of rabbit splenocytes and human PBMCs.** The spleen from a New Zealand White rabbit was removed aseptically and placed into RPMI 1640 medium (Lonza, Walkersville, MD) supplemented with 2% fetal bovine serum, 1% penicillin-streptomycin, and 200 μg/ml glutamine (24). Cell suspensions were obtained by teasing into RPMI 1640 medium, with cell washing in RPMI 1640 medium. Spleen cells were resuspended in RPMI 1640 medium to obtain a concentration of 1.0 × 10^6 cells/ml and aliquoted (200 μl/well) into a 96-well flat-bottom culture plate (Becton, Dickinson, Franklin Lakes, NJ) for lymphocyte proliferation assays.

Heparinized (10 U/ml) human blood was obtained in compliance with an approved University of Minnesota IRB protocol (1004M80313). Blood was layered onto a Histopaque-1077 gradient solution (Sigma, St. Louis, MO), and cells at the interface (PBMCs) were collected and washed with RPMI 1640 medium (6). PBMCs were suspended in a volume of RPMI 1640 medium for a final concentration of between 5.0 × 10^5 and 1.0 × 10^6 cells/ml.

**Rabbit and human lymphocyte proliferation assays.** Purified rabbit and human lymphocytes were incubated with different concentrations of SEC subtypes (0.1 pg to 1 μg per well) for 3 days at 37°C in...
5% CO₂, as described previously (6). For some experiments, we used IVG (1.74, 1.74, and 174.0 μg/well) to neutralize SEC subtypes, as it has been shown to neutralize S. aureus SaG activity (14). After 3 days, 1 μCi of [3H]thymidine was added to each well of lymphocytes, with exposure for an additional 24 h. A Mash II apparatus (Microbiological Associates, Bethesda, MD) was used to collect cellular DNA onto glass fiber filters; thymidine uptake was measured using a liquid scintillation counter (model LS; Beckman Instruments, Fullerton, CA). Data were expressed as averages from four replicates, in counts per minute. 

Experiments testing the effects of different SEC subtype concentrations on PBMCs were repeated at least 4 times; plots shown represent total data.

**SEC-induced enhancement of lipopolysaccharide shock in Dutch-belted rabbits.** A rapidly fatal model of TSS in rabbits can be tested based on the ability of SaG to amplify the lethal effects of lipopolysaccharide (LPS) through synergistic tumor necrosis factor alpha (TNF-α) production. We used this model to compare the lethal properties of all 4 SEC subtypes. Young adult rabbits were administered 0.001, 0.01, 0.1, or 10.0 μg/kg of SEC toxin (SEC-1, SEC-2, SEC-3, or SEC-4) dissolved in pyrogen-free phosphate-buffered saline (PBS) intravenously via the marginal ear vein, as described previously (25, 26). At least three rabbits were administered each SEC dose. Four hours after the administration of SEC subtypes, 10 μg/kg of LPS from *Salmonella enterica* serovar Typhimurium was given intravenously. Rabbits were monitored for lethality and euthanized with phenytoin-pentobarbital (Beuthanasia D; Schering-Plough Animal Health Corporation, Union, NJ) (1 ml/kg) if they exhibited failure to right themselves and escape behavior. All experiments were conducted according to the guidelines of the University of Minnesota Institutional Animal Care and Use Committee. The lethal dose 50% endpoint (LD₅₀) values were calculated according to the technique of Reed and Muench (27).

**Statistical analysis.** Fisher’s exact test was utilized to calculate the statistical significance of SEC subtype associations with human illness. Student’s t test was used to assess differences in the results of lymphocyte proliferation assays.

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**REFERENCES**

1. Spaulding AR, Salgado-Pabón W, Kohler PL, Horswill AR, Leung DYM, Schlievert PM. 2013. Staphylococcal and streptococcal superantigen exotoxins. Clin Microbiol Rev 26:422–447. https://doi.org/10.1128/CMR.00104-12.

2. McCormick JK, Yarwood JM, Schlievert PM. 2001. Toxic shock syndrome and bacterial superantigens: an update. Annu Rev Microbiol 55:77–104. https://doi.org/10.1146/annurev.micro.55.1.77.

3. Schlievert PM, Davis CC. 2020. Device-associated menstrual toxic shock syndrome. Clin Microbiol Rev 33:e00323-19. https://doi.org/10.1128/CMR.00323-19.

4. Klevens RM, Morrison MA, Nadle J, Petit S, Gershman K, Ray S, Harrison LH, Lynfield R, Dumyati G, Townes JM, Craig AS, Zell ER, Fosheim GE, McDougal LK, Carey RB, Fridkin SK, Active Bacterial Core Surveillance MRSA Investigators. 2007. Invasive methicillin-resistant *Staphylococcus aureus* infections in the United States. JAMA 298:1763–1771. https://doi.org/10.1001/jama.298.15.1763.

5. Schlievert PM, Orwin PM, Schlievert PM. 2000. Exotoxins of *Staphylococcus aureus*. Clin Microbiol Rev 13:36–34. https://doi.org/10.1128/cmr.13.1.36-34.2000.

6. Pindexter NJ, Schlievert PM. 1985. Toxic-shock-syndrome toxin 1-mediated proliferation of lymphocytes: comparison of the mitogenic response of human, murine, and rabbit lymphocytes. J Infect Dis 151:65–72. https://doi.org/10.1093/infdis/151.1.65.

7. Fraser JD, Prof T. 2008. The bacterial superantigen and superantigen-like proteins. Immunol Rev 225:226–243. https://doi.org/10.1111/j.1600-065X.2008.00681.x.

8. Bohach GA, Schlievert PM. 1989. Conservation of the biologically active portions of staphylococcal enterotoxins C1 and C2. Infect Immun 57:2249–2252. https://doi.org/10.1128/IAI.57.7.2249-2252.1989.

9. Couch JL, Betley MJ. 1989. Nucleotide sequence of the type C3 staphylococcal enterotoxin gene suggests that intergenic recombination causes antigenic variation. J Bacteriol 171:4507–4510. https://doi.org/10.1128/JB.171.8.4507-4510.1989.

10. Baba T, Takeuchi F, Kuroda M, Yuzawa H, Aoki K, Oguchi A, Nagai Y, Iwama N, Asano K, Naimi T, Kuroda H, Cui L, Yamamoto K, Hiramatsu K. 2002. Genome and virulence determinants of high virulence community-associated MRSA. Lancet 359:1819–1827. https://doi.org/10.1016/S0140-6736(02)08713-5.

11. Shukla SK, Karow ME, Brady JM, Stemper ME, Kislowsky J, Moore N, Wroblewski K, Chyou P-H, Warshauer DM, Reed KD, Lynfield R, Schwab WR. 2010. Virulence genes and genotype associations in nasal carriage, community-associated methicillin-susceptible and methicillin-resistant *Staphylococcus aureus* isolates. J Clin Microbiol 48:3582–3592. https://doi.org/10.1128/JCM.00657-10.

12. Fey PD, Said-Salim B, Rupp ME, Hinrichs SH, Boxrud DJ, Davis CC, Kreiswirth BN, Schlievert PM. 2003. Comparative molecular analysis of community- or hospital-acquired methicillin-resistant *Staphylococcus aureus*. Antimicrob Agents Chemother 47:196–203. https://doi.org/10.1128/AAC.47.1.196-203.2003.

13. Voyich JM, Otto M, Mathema B, Braughton KR, Whitney AR, Welty D, Long RD, Dorward DW, Gardner DJ, Lina G, Kreiswirth BN, DeLeo FR. 2006. Is Panton-Valentine leukocidin the major virulence determinant in community-associated methicillin-resistant *Staphylococcus aureus* disease? J Infect Dis 194:1761–1770. https://doi.org/10.1086/509506.

14. Schlievert PM. 2001. Use of intravenous immunoglobulin in the treatment of staphylococcal and streptococcal toxic shock syndromes and related illnesses. J Allergy Clin Immunol 108:S507–510. https://doi.org/10.1016/j.jaci.2001.11.8782.

15. Lee PK, Derenger JR, Kreiswirth BN, Novick RP, Schlievert PM. 1991. Fluid replacement protection of rabbits challenged subcutaneously with toxic shock syndrome toxins. Infect Immun 59:879–884. https://doi.org/10.1128/IAI.59.3.879-884.1991.

16. Schlievert PM, Cahill MP, Hostager BS, Brosnahan AJ, Klingelhoft AJ, Gourmonc FA, Bishop GA, Leung DYM. 2019. Staphylococcal superantigens stimulate epithelial cells through CD40 to produce chemokines. mBio 10:e00214-19. https://doi.org/10.1128/mBio.00214-19.

17. Leder R, Llera A, Lavoie PM, Lebedeva MI, Li H, Sekaly RP, Bohach GA, Gahr PJ, Schlievert PM, Karjalainen K, Mariuzza RA. 1998. A mutational analysis of the binding of staphylococcal enterotoxins B and C3 to the T cell receptor beta chain and major histocompatibility complex class II. J Exp Med 187:823–833. https://doi.org/10.1084/jem.187.6.823.

18. Schlievert PM, Case LC, Strandberg KL, Abrams BB, Leung DY. 2008. Superantigen profile of *Staphylococcus aureus* isolates from patients with steroid-resistant atopic dermatitis. Clin Infect Dis 46:1562–1567. https://doi.org/10.1086/586746.

19. Schlievert PM, Strandberg KL, Lin YC, Peterson ML, Leung DY. 2010. Secreted virulence factor comparison between methicillin-resistant and methicillin-sensitive *Staphylococcus aureus* and its relevance to atopic dermatitis. J Allergy Clin Immunol 125:39–49. https://doi.org/10.1016/j.jaci.2009.10.039.

20. Schlievert PM, Jablonski LM, Roggiani M, Sadler I, Callantine S, Mitchell DT, Ohlendorf DH, Bohach GA. 2000. Pyrogenic toxin superantigen site

USA400 *Staphylococcus aureus* isolates. J Clin Microbiol 48:3582–3592. https://doi.org/10.1128/JCM.00657-10.
21. Brosnahan AJ, Mantz MJ, Squier CA, Peterson ML, Schlievert PM. 2009. Cytolysins augment superantigen penetration of stratified mucosa. J Immunol 182:2364–2373. https://doi.org/10.4049/jimmunol.0803283.

22. Strandberg KL, Rotschafer JH, Vetter SM, Buonpane RA, Kranz DM, Schlievert PM. 2010. Staphylococcal superantigens cause lethal pulmonary disease in rabbits. J Infect Dis 202:1690–1697. https://doi.org/10.1086/657156.

23. Blomster-Hautamaa DA, Schlievert PM. 1988. Preparation of toxic shock syndrome toxin-1. Methods Enzymol 165:37–43. https://doi.org/10.1016/s0076-6879(88)65009-9.

24. Barsumian EL, Schlievert PM, Watson DW. 1978. Nonspecific and specific immunological mitogenicity by group A streptococcal pyrogenic exotoxins. Infect Immun 22:681–688. https://doi.org/10.1128/IAI.22.3.681-688.1978.

25. Kim YB, Watson DW. 1970. A purified group A streptococcal pyrogenic exotoxin. Physiochemical and biological properties including the enhancement of susceptibility to endotoxin lethal shock. J Exp Med 131:611–622. https://doi.org/10.1084/jem.131.3.611.

26. Schlievert PM. 1982. Enhancement of host susceptibility to lethal endotoxin shock by staphylococcal pyrogenic exotoxin type C. Infect Immun 36:123–128. https://doi.org/10.1128/IAI.36.1.123-128.1982.

27. Reed LJ, Muench H. 1938. A simple method of estimating fifty per cent endpoints. Am J Hyg 27:493–497. https://doi.org/10.1093/oxfordjournals.aje.a118408.

28. Schlievert PM, Osterholm MT, Kelly JA, Nishimura RD. 1982. Toxin and enzyme characterization of Staphylococcus aureus isolates from patients with and without toxic shock syndrome. Ann Intern Med 96:937–940. https://doi.org/10.7326/0003-4819-96-6-937.

29. Schlievert PM, Shands KN, Dan BB, Schmid GP, Nishimura RD. 1981. Identification and characterization of an exotoxin from Staphylococcus aureus associated with toxic shock syndrome. J Infect Dis 143:509–516. https://doi.org/10.1093/infdis/i43.4.509.

30. Rizkallah MF, Tolaymat A, Martinez JS, Schlievert PM, Ayoub EM. 1989. Toxic shock syndrome caused by a strain of Staphylococcus aureus that produces enterotoxin C but not toxic shock syndrome toxin-1. Am J Dis Child 143:848–849. https://doi.org/10.1001/archpedi.1989.02150190098031.

31. Bohach GA, Schlievert PM. 1987. Nucleotide sequence of the staphylococcal enterotoxin C1 gene and relatedness to other pyrogenic toxins. Mol Gen Genet 209:15–20. https://doi.org/10.1007/BF00329830.

32. MacDonald KL, Osterholm MT, Hedberg CW, Schrock CG, Peterson GF, Jentzen JM, Leonard SA, Schlievert PM. 1987. Toxic shock syndrome: A newly recognized complication of influenza and influenzalike illness. JAMA 257:1053–1058. https://doi.org/10.1001/jama.257.8.1053.

33. CDC. 1999. Four pediatric deaths from community-acquired methicillin-resistant Staphylococcus aureus—Minnesota and North Dakota, 1997-1999. MMWR Morb Mortal Wkly Rep 48:707–710.

34. Kravitz GR, Dries DJ, Peterson ML, Schlievert PM. 2005. Purpura fulminans due to Staphylococcus aureus. Clin Infect Dis 40:941–947. https://doi.org/10.1086/428573.

35. Avena RM, Bergdoll MS. 1967. Purification and some physicochemical properties of enterotoxin C, Staphylococcus aureus strain 361. Biochemistry 6:1474–1480. https://doi.org/10.1021/bi00857a033.

36. Marr JC, Lyon JD, Roberson JR, Lupher M, Davis WC, Bohach GA. 1993. Characterization of novel type C staphylococcal enterotoxins: biological and evolutionary implications. Infect Immun 61:4254–4262. https://doi.org/10.1128/IAI.61.10.4254-4262.1993.