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Functional Analysis of the TCR Binding Domain of Toxic Shock Syndrome Toxin-1 Predicts Further Diversity in MHC Class II/Superantigen/TCR Ternary Complexes

John K. McCormick,* Timothy J. Tripp,* Andrea S. Llera,† Eric J. Sundberg,† Martin M. Dinges,* Roy A. Mariuza,† and Patrick M. Schlievert†

Superantigens (SAGs) aberrantly alter immune system function through simultaneous interaction with lateral surfaces of MHC class II molecules on APCs, and with particular variable regions of the TCR (Vβ). To further define the interface between the bacterial SAG toxic shock syndrome toxin-1 (TSST-1) and the TCR, we performed alanine scanning mutagenesis within the putative TCR binding region of TSST-1 along the central α helix adjacent to the N-terminal α helix and the β7–β9 loop as well as with two universally conserved SAG residues (Leu137 and Tyr144 in TSST-1). Mutants were analyzed for multiple functional activities, and various residues appeared to play minor or insignificant roles in the TCR interaction. The locations of six residues (Gly16, Trp116, His135, Gln136, and Gln139), each individually critical for functional activity as well as direct interaction with the human TCR Vβ2.1-chain, indicate that the interface occurs in a novel region of the SAG molecule. Based on these data, a model of the MHC/TSST-1/TCR ternary complex predicts similarities seen with other characterized SAGs, although the CDR3 loop of Vβ2.1 is probably involved in direct SAG-TCR molecular interactions, possibly contributing to the TCR Vβ specificity of TSST-1. The Journal of Immunology, 2003, 171: 1385–1392.

Superantigens (SAGs) are class of bacterial or viral proteins that aberrantly alter immune system function through simultaneous interaction with lateral surfaces of MHC class II molecules on APCs, and with particular variable regions of the TCR (Vβ) (reviewed in Refs. 2 and 3). Although SAGs bind the TCR with affinities similar to those of typical TCR/peptide/MHC interactions (4–8), SAGs bind primarily based on specific TCR Vβ-chains without the requirement for Ag processing and can thus stimulate large fractions of both CD4+ and CD8+ T lymphocytes. This activity, defined as superantigenicity (1), results in the extensive activation and proliferation of T cells with the corresponding massive release of cytokines, including TNF-β, IL-2, and IFN-γ from T cells and TNF-α and IL-1 from APCs. It is the subsequent actions of these and other cytokines that are believed to mediate the pathological consequences leading to capillary leakage and the toxic shock syndrome (TSS). Ultimately, following the initial expansion, SAG-specific T lymphocytes may be rendered anergic (9).

TSS toxin-1 (TSST-1) is a bacterial SAG secreted by strains of Staphylococcus aureus that belongs to the pyrogenic toxin class of SAGs (3). TSST-1 has been associated with TSS since 1981 (10, 11), and it has been estimated that this exotoxin is responsible for ~50% of nonmenstrual-associated TSS cases due to S. aureus and essentially all cases of menstrual-associated TSS (12). This latter association is probably due to the apparently unique ability of TSST-1 among the pyrogenic toxin SAGs to cross mucosal surfaces (13). The crystal structure of this SAG (14, 15) as well as those of many others have been determined (reviewed in Ref. 2), and each shows a similar architecture, divided into two domains. The small domain (residues 18–89 in TSST-1) is composed of a five-strand mixed β-barrel, and the large domain (residues 1–170 and 90–194 in TSST-1) is built around a long central α helix (residues 125–140 in TSST-1) lying against a five-strand β-sheet. The crystal structures of MHC class II complexed with TSST-1 (16) and staphylococcal enterotoxin B (SEB) (17) have been determined, indicating that the interface occurs through the small domain within the O/B fold motif. These studies also revealed that although the binding regions of SEB and TSST-1 on MHC class II overlap, the interaction of TSST-1 is partially peptide dependent (18), while SEB binds more to the edge of the peptide binding groove of MHC class II. The crystal structure of MHC class II complexed with the high affinity, zinc-dependent binding site within the β1b domain on the large domain of streptococcal pyrogenic exotoxin C (SPE C) has also been determined, revealing extensive contacts with the class II bound peptide and indicating that the interaction of SAG with MHC class II can occur through diverse surfaces (19). Furthermore, the structure of SEH bound to HLA-DR1 revealed an interaction similar to that of SPE C although with unrelated peptides, indicating the possibility that the...
The interactions between the relevant TCR β-chain with the SAGs staphylococcal enterotoxin C (SEC) (21) and SEB (22) as well as streptococcal pyrogenic exotoxin A (SPE A) and SPE C (23) have been structurally characterized. These studies have revealed that the orientations of SEB, SEC, and SPE A, when bound to their respective TCR β-chains, are highly similar (23); however, the SPE C/TCR β-chain complex had an entirely different binding mode for SAG-TCR interaction, with a more extensively buried interface and numerous specific interactions. Significantly, the SPE C/TCR complex modeled with the SPE C/MHC complex indicated a complete dissociation between the TCR and MHC molecules (23). In other work, staphylococcal enterotoxin A (SEA) has been shown to stimulate γδ T cells expressing Vγ2, similar to SAG activation of αβ T cells, and mutagenesis and domain swapping suggested that the SEA recognition site for γδ TCR was similar to the SEC recognition site for Vβ8.2 (24). Finally, TCR interaction with the SAG from Mycoplasma arthritidis is stabilized by the β-chain CDR3 region (25), a region not involved in other characterized SAG/TCR interactions, providing further evidence for the diversity of SAG interaction with the TCR.

The cocrystal structure of TSST-1 complexed with the human Vβ2.1 TCR (hVβ2.1) has yet to be determined, and although various residues have been implicated as important for this interaction (26–32), the structure of the TSST-1/hVβ2.1 interface remains to be characterized. In sum, these mutagenesis studies indicate that the TCR binding domain of TSST-1 resides in the cleft located along the central α helix between the N-terminal α helix and the β7-β8 loop (32) in contrast to SEB, SEC, SPE A, and SPE C, where this binding domain is located in a cleft on the other side of the molecule between the large and small domains (21–23). A key issue regarding previous TSST-1 mutagenesis studies is that each of these mutations is hypothesized to reside in the TSST-1/TCR interface because they are located outside the MHC binding domain, although direct interaction with the TCR has not been demonstrated. Our current study was designed to further define residues important for the functional interaction of TSST-1 with the TCR by alanine scanning mutagenesis. Our data suggest that residues Tyr13, Ser15, His137, and Tyr144 are irrelevant for TCR recognition or play a minor stabilizing role in the interaction. Our data also indicate that residues Gly16, Trp116, Gly132, His135, Gly136, and Gly139 are individually crucial for direct binding to hVβ2.1. Due to the location of these residues, we propose a model in which the MHC/TSST-1/TCR ternary complex is similar to the ternary complex of SEB, SEC, and SPE A, yet the TSST-1 complex may allow for direct interactions between the CDR3 loop of the TCR β-chain and TSST-1. Based on this model and in agreement with the variability of Vβ-specificity seen with different SAGs, it is likely that β-chain ligation by TSST-1 represents a distinct mode of SAG/TCR engagement.

### Materials and Methods

#### Reagents

All chemicals were of analytical grade. Oligonucleotides were obtained from Integrated DNA Technologies (Coralville, IA). Pfu DNA polymerase and dNTPs were purchased from Stratagene (La Jolla, CA). Water and materials were maintained pyrogen free for all in vivo and biological activity experiments.

#### TSST-1 mutants

The wild-type TSST-1 gene (tstH; where H stands for human isolate) (33) was obtained from a chromosomal DNA preparation of *S. aureus* MNS digested with HindIII and SalI. The fragment containing tstH was cloned into pCE104 (a shuttle vector containing pE194 and pUC18), also digested with HindIII and SalI to create pCE107. This plasmid was used to express wild-type TSST-1 from *S. aureus* RN4220 and was also used as the template for mutagenesis. Mutations were generated by the inverse PCR Quik-Change mutagenesis procedure (Stratagene), and mutagenic primers are listed in Table I. The double-mutant Y115A/W116G was created because primers were based on the original published sequence that contained a mutation indicating Gly at position 116 (33) when in reality there is a Trp at this position. Mutated plasmids were transformed into *Escherichia coli* DH5α, and the tstH gene was sequenced to confirm that each mutation was incorporated and no second site mutations were generated. Correct plasmids were transformed into *S. aureus* RN4220 by protoplast transformation (34).

#### Protein purification

Mutant toxins were produced and purified from *S. aureus* RN4220 harboring recombinant plasmids as previously described (11). Briefly, strains were grown to stationary phase in pyrogen-free beef heart medium containing 5 µg/ml of erythromycin for plasmid maintenance, and culture supernatants were concentrated by precipitation with a final concentration of 80% ethanol. Precipitates were resolubilized in pyrogen-free water, and purification was achieved by preparative flatbed isoelectric focusing. The purity of the toxin preparations was assessed by SDS-PAGE.

The cDNA for RSD3 TCR was a gift from U. Utz and R.-P. Sekaly (University of Montreal, Montreal, Canada). This human TCR is specific for HTLV-I TAX11–19 peptide in the context of HLA-A2.1. The expression and purification of recombinant human TCR Vβ2.1/Jβ2.3D/Jβ2.1Cβ2 (herein referred to as hVβ2.1) were accomplished as previously described (23).

#### T cell proliferation assays

Gradient purified human PBMCs or unfractionated rabbit spleen cells were stimulated in vitro in 96-well (2 × 10⁴ cells/well) microtiter plates with serial 1/10 dilutions (in quadruplicate) of purified wild-type or mutant TSST-1 molecules. RPMI medium (BioWhittaker, Walkersville, MD) supplemented with 10% FCS (Sigma-Aldrich, St. Louis, MO), 100 units/ml of penicillin G, 100 µg/ml of streptomycin, and 2 mM of L-glutamine were used. After 3 days of incubation, proliferation of T cells was measured by [3H]-thymidine uptake.

### Table I. Altered amino acid residues of TSST-1 and primers used for construction of various mutants by site-directed mutagenesis

<table>
<thead>
<tr>
<th>Targeted Amino Acid</th>
<th>Mutant</th>
<th>Corresponding Primers*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tyr13</td>
<td>Y13A</td>
<td>5′-TTGCTAGACTGCGCCAGGCTGTTGCTC-3′</td>
</tr>
<tr>
<td>Ser15</td>
<td>S15A</td>
<td>5′-GACTGGTTAGTGGGCTGCAGACT-3′</td>
</tr>
<tr>
<td>Tyr21</td>
<td>Y21A</td>
<td>5′-AGCCCTTTAAGGAGGCCTGCAAATTC-3′</td>
</tr>
<tr>
<td>Gly22</td>
<td>G22A</td>
<td>5′-CTGCTAAATGTTGGCGGCAAGTCGAT-3′</td>
</tr>
<tr>
<td>Tyr115/Tyr144</td>
<td>Y115A/W116G</td>
<td>5′-AGCCCTTTAAGGAGGCCTGCAAATTC-3′</td>
</tr>
<tr>
<td>Leu137</td>
<td>L137A</td>
<td>5′-ATTCGTCATCAAGCCCTGCTAACGACAT-3′</td>
</tr>
<tr>
<td>Gln139</td>
<td>G139A</td>
<td>5′-CATACGCTACATGCGATGACT-3′</td>
</tr>
<tr>
<td>His141</td>
<td>H141A</td>
<td>5′-CTAATGATATCCGATGATGACT-3′</td>
</tr>
<tr>
<td>Tyr144</td>
<td>Y144A</td>
<td>5′-ATACGCTGATTACGGTCAAGCGAT-3′</td>
</tr>
<tr>
<td>Gly7/Ser32</td>
<td>G31S/S32</td>
<td>5′-TTGATGATTTGTGCTGCTGCTGCT-3′</td>
</tr>
</tbody>
</table>

*Underlined bases are areas in the primer that have been mutated to elicit the corresponding amino acid mutation. A complementary primer for each primer listed here was also made to perform the mutagenesis method outlined in Methods and Materials.
rabbits were pretreated with low doses of wild-type or mutated TSST-1 to determine the pyrogenic activity of the mutants. Animals were monitored for fever and lethal shock in American Dutch belted rabbits (Birchwood Farms, WI). In the first model, mini-osmotic pumps (Alza Pharmaceuticals, Palo Alto, CA) were preloaded with 200 μg of wild-type or mutated TSST-1 proteins in PBS (pH 7.2) (35). These devices are designed to release toxin at a constant rate over a period of 7 days to mimic toxin exposure during infection. Mini-osmotic pumps were s.c. implanted in American Dutch belted rabbits, and three (or more) rabbits were used for each toxin preparation. Temperatures were recorded on days 0 and 2 to determine the pyrogenic activity of the mutants. Animals were monitored for symptoms of TSS, and mortality was recorded over a 10-day period. In the enhancement of endotoxin shock model (36), American Dutch belted rabbits were pretreated with low doses of wild-type or mutated TSST-1 proteins (5 μg/kg of body weight) administered i.v. by the marginal ear vein. After 4 h animals were challenged with a sublethal dose (10 μg/kg) of endotoxin from Salmonella typhimurium. Animals were monitored for symptoms of TSS, and mortality was recorded over 48 h. In both models, animals displaying clear signs of lethal shock were prematurely euthanized in accordance with our animal experimentation protocols. All animal experiments performed in this study have received prior approval by the University of Minnesota institutional review board.

**Results**

**TSST-1 mutants**

To further define residues important for the functional interaction of TSST-1 with the TCR, we performed alanine scanning mutagenesis of various residues located near the putative TCR binding region located along the central α helix adjacent to the N-terminal α helix and the ββ-β9 loop (32). Alanine substitution was chosen because this technique removes the targeted amino acid side chain while minimizing steric or electrostatic constrains on the tertiary structure of the protein (37). Mutations encompassed residues Tyr115 and Ser115 located in the N-terminal α helix region, residues Tyr115 and Trp116 located between strands β8 and β9, and residues Leu137, Gln310, His311, and Tyr144 located near the distal end of the central helix that joins the large and small domains. In this work we also included the previously constructed mutants G16V (30), which is located on the N-terminal α helix, as well as E132K and Q136A, located along the central helix (31). A control MHC class II binding mutant was also created (G31S/S32P) that lacks significant proliferative activity (30), and these residues are known to be located at the interface with the MHC class II molecule HLA-DR1 (16). The Ser12 mutation is believed to account for most of the impaired interaction with MHC class II (30), although other data suggest that Gly31 is also critical for MHC class II binding (38). All the mutant genes (Table I) were constructed using plasmid pCE107 as a template, and mutated proteins were expressed and purified from S. aureus RN4220. Each protein was purified to apparent homogeneity as determined by SDS-PAGE, with a typical yield of ~5 mg/liter.

**Mitogenic capacity of various TSST-1 mutants**

Initial experiments determined the effect of each individual mutation on superantigenic activity by measuring the ability of each protein to induce the proliferation of unfractionated rabbit splenocytes or gradient purified human PBMCs (Table II). Mitogenic

**Table II. Mitogenic capacity and dissociation constants for binding of wild-type TSST-1 and mutated proteins to the human Vβ2.1 TCR.**

<table>
<thead>
<tr>
<th>TSST-1 Mutant</th>
<th>Human</th>
<th>Rabbit</th>
<th>$K_d$ (nM)</th>
<th>$\Delta G$ (kCal/mol)</th>
<th>$\Delta \Delta G$ (kCal/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild type</td>
<td>50</td>
<td>50</td>
<td>2.3</td>
<td>$-7.7$</td>
<td></td>
</tr>
<tr>
<td>Y113A</td>
<td>50</td>
<td>5–50</td>
<td>2.1</td>
<td>$-7.7$</td>
<td></td>
</tr>
<tr>
<td>S15A</td>
<td>50</td>
<td>50</td>
<td>1.0</td>
<td>$-8.2$</td>
<td>$-0.5$</td>
</tr>
<tr>
<td>G16V</td>
<td>$&gt;5,000,000$</td>
<td>$&gt;5,000,000$</td>
<td>0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Y115A</td>
<td>500</td>
<td>50–500</td>
<td>13.1</td>
<td>$-6.7$</td>
<td>1.0</td>
</tr>
<tr>
<td>W116A</td>
<td>50,000</td>
<td>50–500</td>
<td>NB$^*$</td>
<td>$&gt;-5.5$</td>
<td>$&gt;2.2$</td>
</tr>
<tr>
<td>E132K</td>
<td>$&gt;5,000,000$</td>
<td>$&gt;5,000,000$</td>
<td>NB</td>
<td>$&gt;-5.5$</td>
<td>$&gt;2.2$</td>
</tr>
<tr>
<td>H135A</td>
<td>5,000,000</td>
<td>5,000,000</td>
<td>NB</td>
<td>$&gt;-5.5$</td>
<td>$&gt;2.2$</td>
</tr>
<tr>
<td>Q136A</td>
<td>$&gt;5,000,000$</td>
<td>500,000</td>
<td>NB</td>
<td>$&gt;-5.5$</td>
<td>$&gt;2.2$</td>
</tr>
<tr>
<td>L137A</td>
<td>500</td>
<td>50–500</td>
<td>1.1</td>
<td>$-8.1$</td>
<td>$-0.4$</td>
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<tr>
<td>Q139A</td>
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<td>5,000,000</td>
<td>NB</td>
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<td>H141A</td>
<td>50–500</td>
<td>500</td>
<td>1.9</td>
<td>$-7.8$</td>
<td>$-0.1$</td>
</tr>
<tr>
<td>Y144A</td>
<td>500</td>
<td>500</td>
<td>1.4</td>
<td>$-8.0$</td>
<td>$-0.3$</td>
</tr>
<tr>
<td>G31S/S32P</td>
<td>$&gt;5,000,000$</td>
<td>$&gt;5,000,000$</td>
<td>1.4</td>
<td>$-8.0$</td>
<td>$-0.3$</td>
</tr>
</tbody>
</table>

$^*$ Mitogenic capacity of mutant TSST-1 molecules was estimated by the dose (picograms per milliliter) required to induce 50% of mitogenic activity compared with maximal activity of wild-type TSST-1.

$^\ddagger$ Affinity measurements were determined by surface plasmon resonance as described in Materials and Methods. The values for the individual ΔG's were calculated from the $K_d$ obtained from surface plasmon resonance measurements according to the equation $\Delta G = -RT \ln (K_d)$, where R is the universal gas constant (~0.001986 kCal/mo).K), and T is the absolute temperature in Kelvin (298.15 K).

$^a$ NB, binding not detected up to 100 μM human Vβ2.1C.
capacity was defined as the SAG dose required to induce T cell proliferation that was 50% or greater than maximal proliferation induced by wild-type TSST-1 for each assay. Mitogenicity assays typically exhibited a clear dose response, where wild-type mitogenic capacity was achieved with SAG concentrations at 50 pg/mL, similar to previous data (30, 31, 39). For some protein preparations, the highest dose tested (5 µg/mL) occasionally had reduced activity or failed to stimulate proliferation. In this case, the lower dilutions that gave higher, dose-dependent mitogenic activities, were used for mitogenic capacity determination. Using these criteria, the mitogenic capacity of mutants Y13A and S15A was similar to that of wild-type TSST-1, while the mutants Y115A, L137A, H141A, and Y144A each displayed a slight loss of activity (>10-fold increase in mitogenic capacity) for both human and rabbit cells (Table II). The single-site mutants G16V, E132K, H135A, Q136A, and Q139A and the double-site mutant Y115A/W116G each had drastically reduced mitogenicity activity for human cells (Table II). The H135A, Q139A, and Y115A/W116G mutants each failed to produce significant proliferation for human lymphocytes above background levels at any concentration tested. Similar results were obtained for the control MHC class II binding mutant G31S/S32P. The W116A mutant displayed a strongly reduced ability to stimulate human cells (~1000-fold), yet maintained near wild-type activity for rabbit cells. Despite the Q136A and E132K mutants being inactive for human lymphocytes, both stimulated rabbit lymphocytes, but with reduced activity (Table II).

Direct binding of TSST-1 mutants to hVb2.1
Since TSST-1 stimulates TCR Vβ2 (40), the binding affinity of each TSST-1 mutant to the hVb2.1 chain was determined by surface plasmon resonance (BIAcore; Table II). This method has been successfully used to evaluate the stabilizing contribution of the TCR α-chain to the trimeric TCR/SAG/MHC complex (6, 41) and to map the free energy of the interactions between SEC3 and mouse Vβ8.2 by mutating both SEC3 (5) and Vβ8.2 TCR (42). Each mutated TSST-1 protein was coupled directly to the dextran matrix of sensor chips through the primary amino groups. Unglycosylated hVb2.1 was injected, and concentration-dependent surface plasmon resonance profiles were recorded. A representative profile for wild-type TSST-1 and the SEB control, with a corresponding Scatchard plot, are shown in Fig. 1.

The Y13A, S15A, L137A, H141A, and Y144A mutants with similar or slightly reduced mitogenic profiles compared with that of wild-type TSST-1 also bound hVb2.1 with affinities comparable to that of wild-type TSST-1. The Y115A mutant had a slightly reduced binding affinity (Table II), indicating a possible minor stabilizing interaction with hVb2.1. Although L137V has been reported to cause a major decrease in mitogenic activity (30), the data presented here showed that L137A maintained both binding to hVb2.1 and near wild-type mitogenic capacity. Therefore, we believe that Leu137 is not critical for interaction with the TCR. We also demonstrate that G16V, W116A, E132K, H135A, Q136A, and Q139A did not detectably bind to the hVb2.1, providing direct evidence that the corresponding residues are critical for this interaction. Consistent with the G31S/S32P mutant being located within the MHC class II binding interface, this mutant maintained binding to hVb2.1.

Induction of fever and toxicity of various TSST-1 mutants
Mutant proteins were examined in vivo for their ability to invoke fever responses and to induce lethal shock in two rabbit models of TSS (Table III). The presence of fever was taken as an average increase of ≥±0.5°C after 2 days of exposure to toxin by the miniosmotic pump model. Using these criteria, wild-type TSST-1 and mutants Y13A, S15A, Y115A, and L137A were all pyrogenic. These toxins as well as the H141A mutant were also lethal in both the miniosmotic pump and endotoxin enhancement models, indicating that individual residues were not critical for this activity despite slight decreases in mitogenic capacity or hVb2.1 binding for Y115A, W116A, L137A, and H141A (Table II). Although the H141A mutant toxin did not cause significant fever, temperatures appeared to be elevated, yet did not reach the threshold criteria to be indicative of a fever response. This may indicate that the H141A mutant retained weak pyrogenic activity. The Y115A/W116G, Q139A, and Y144A mutants were nontoxic in the miniosmotic pump model of TSS, but only Y115A/W116G and Q139A mutants were nontoxic in the endotoxin enhancement model of TSS (Table III). Surprisingly, the Y144A mutant remained lethal in the endotoxin enhancement model of TSS. This result was remarkable because both models have always directly correlated for TSST-1 mutants and also for experiments with other staphylococcal and streptococcal SAGs. The inability of Y144A to cause death in the miniosmotic pump model initially led us to believe that residue Tyr144 was important for lethal activity. The miniosmotic pump model typically takes up to 5 days or more for the animal to develop shock, and we hypothesize that this protein is structurally unstable and has degraded to nontoxic levels by this time. The short time course leading to death in the endotoxin enhancement model (<24 h) and the relatively high doses of TSST-1 that the animals received for this model probably allowed for activity before toxin degradation. This may also be true for the lack of pyrogenic activity for this mutant. Alternatively, the endotoxin enhancement model may be more sensitive to TSST-1-mediated lethality than is the continuous infusion model. Finally, evaluation of the Q139A mutant, as assessed by the inability to proliferate T cells, induce fever, bind hVb2.1, or demonstrate toxicity in two

![Image](http://www.jimmunol.org/Download/323x501.png)
TSS Models (lethal/total)

<table>
<thead>
<tr>
<th>Protein</th>
<th>Baseline ± SD</th>
<th>48 hr (± SD)</th>
<th>Δ°C</th>
<th>Miniosmotic pump</th>
<th>Endotoxin enhancement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild type</td>
<td>38.46 ± 0.18</td>
<td>39.46 ± 0.48</td>
<td>1.00</td>
<td>8/9</td>
<td>6/6</td>
</tr>
<tr>
<td>Y13A</td>
<td>38.87 ± 0.15</td>
<td>39.50 ± 0.20</td>
<td>0.63</td>
<td>3/3</td>
<td>3/3</td>
</tr>
<tr>
<td>S15A</td>
<td>38.63 ± 0.12</td>
<td>39.60 ± 0.36</td>
<td>0.97</td>
<td>3/3</td>
<td>3/3</td>
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<tr>
<td>G16V</td>
<td>39.18 ± 0.26</td>
<td>38.82 ± 0.36</td>
<td>−0.36</td>
<td>ND</td>
<td>ND</td>
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<tr>
<td>Y115A</td>
<td>37.90 ± 0.35</td>
<td>39.00 ± 0.72</td>
<td>0.10</td>
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<td>3/3</td>
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<tr>
<td>W116A</td>
<td>38.30 ± 0.15</td>
<td>39.30 ± 0.72</td>
<td>0.51</td>
<td>3/3</td>
<td>3/3</td>
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<tr>
<td>Y115A/W116G</td>
<td>38.37 ± 0.45</td>
<td>38.27 ± 0.06</td>
<td>−0.10</td>
<td>0/3</td>
<td>0/3</td>
</tr>
<tr>
<td>L137A</td>
<td>38.53 ± 0.12</td>
<td>39.77 ± 0.78</td>
<td>1.24</td>
<td>3/3</td>
<td>3/3</td>
</tr>
<tr>
<td>Q139A</td>
<td>38.73 ± 0.21</td>
<td>38.53 ± 0.40</td>
<td>−0.20</td>
<td>3/3</td>
<td>0/3</td>
</tr>
<tr>
<td>H141A</td>
<td>38.83 ± 0.38</td>
<td>39.10 ± 0.92</td>
<td>0.27</td>
<td>3/3</td>
<td>3/3</td>
</tr>
<tr>
<td>Y144A</td>
<td>38.75 ± 0.26</td>
<td>38.82 ± 0.31</td>
<td>0.07</td>
<td>0/6</td>
<td>0/6</td>
</tr>
<tr>
<td>G31S/S32P</td>
<td>38.73 ± 0.35</td>
<td>39.23 ± 0.45</td>
<td>0.5</td>
<td>0/3</td>
<td>0/3</td>
</tr>
</tbody>
</table>

*Temperatures were taken prior to administration of miniosmotic pumps containing 200 μg of each toxin. Three or more rabbits were used for each protein. The induction of fever is taken as an increase in 0.5°C over a period of 4 h compared with baseline temperatures.

**Two hundred micrograms of each protein was administered in miniosmotic pumps.

Rabbits were treated with 5 μg/ml/kg of TSST-1 or various mutant proteins administered i.v., and mortality was recorded over 10 days. After 4 h, 10 μg/ml/kg of endotoxin was given i.v., and mortality was recorded over 48 h.

Discussion

The three-dimensional structures of several TCR β-chain/SAG complexes have now been reported, revealing the mechanism by which SAGs circumvent the normal mechanism for T cell activation (2). Although the interaction between TSST-1 and the human MHC class II molecule HLA-DR1 has been defined by co-crystal complex analysis (16), the interaction of TSST-1 with the TCR has not yet been fully characterized. Because TSST-1 is associated with most cases of S. aureus TSS, and this SAG represents a distinct subgroup among the pyrogenic toxin SAGs (3), a complete understanding of the interaction between TSST-1 and the TCR is highly desirable. Furthermore, the location of the predicted TCR binding domain of TSST-1 appears to be distinct from the now well characterized TCR binding domains of SEB, SEC, SPE A, and SPE C.

Two of the mutated residues in TSST-1 (Leu<sup>137</sup> and Tyr<sup>144</sup>) appear to be universally conserved among all the characterized pyrogenic toxin SAGs, and both these residues are buried in the TSST-1 crystal structure. The failure of L137A and Y144A to affect functional activity was not unexpected, yet this may imply that these and possibly other universally conserved residues are not important for ligand formation. In conjunction with our current data for Tyr<sup>144</sup>, this finding is complementary to a previous double mutant (H141A/Y144A) that was found to have no mitogenic activity or toxicity in an in vivo rabbit model (27). The H141A/Y144A mutant also lost the ability to be recognized by a mAb, suggesting conformational changes or degradation. We believe that the combined effects of mutating these two residues resulted in a net loss of binding energetics sufficient to cause the loss of function. This may be similar to the double-mutant Y115A/W116G, constructed in this study. The failure of the L137A mutation to largely inhibit mitogenicity is, however, in contradiction to earlier results (30). It was previously determined that L137V had dramatically decreased T cell stimulatory activity, but maintained binding to HLA-DR. The majority of residue Leu<sup>137</sup> is not surface exposed, and it is possible that the Leu→Val mutation imposed a localized steric alteration within the TCR binding region. Indeed, Hurley et al. (30) originally hypothesized that this may explain the inactivity of this mutant. The G31S/S32P mutant behaved as expected in all experiments, including the ability to bind hVβ2.1, although this protein was not mitogenic or pyrogenic, and these results further demonstrate that TSST-1 interaction with the MHC class II molecule is critical for each of these functional activities, including the development of lethal shock.

The slight decrease in hVβ2.1 binding seen for mutations at positions Tyr<sup>115</sup>, Tyr<sup>115</sup>, and His<sup>141</sup> may indicate minor involvement in the TCR/TSST-1 complex, probably through stabilizing interactions. Based on the co-crystal structure of SEC3 in contact with the mouse 14.3.d β-chain (21), the mutation of all SEC3 residues involved in contact with the β-chain revealed that only two of 12 residues (N23A, Q210A) completely lost detectable binding using their criteria for stimulatory capacity, while a third mutation (F176A) had barely detectable mitogenic activity (5). Most other SEC3 mutations did show more intermediate decreases in stimulatory capacity, which we did not observe for the TSST-1 mutants, although three SEC3 mutants showed similar minor decreases in mitogenic activity (5). These residues (Gly<sup>102</sup>, Lys<sup>103</sup>, and Gly<sup>106</sup> in SEC3) are located on the flexible disulphide loop and do not form hydrogen bonds with the 14.3.d TCR (21). It is likely that Tyr<sup>113</sup>, Tyr<sup>115</sup>, and His<sup>141</sup> may participate in similar weak bonding arrangements that are not critical for functional interaction, but contribute to overall binding energetics.

SAG activity leads to massive T cell stimulation and proliferation, and it is generally believed that cytokine release as a result of this activity is responsible for the most severe consequences of TSS. Alternatively, the role of SAGs in autoimmune disorders is probably due to the proliferation of autoreactive T cell subsets. Although mitogenicity and lethality are clearly affiliated, some evidence indicates that superantigenic activity and lethality are separable biological properties. The Q136A mutation of TSST-1 was previously reported to maintain superantigenicity (~60%) for rabbit T cells (but not human T cells), although lethal activity in rabbits was not exhibited at any dose tested (31). The three-dimensional structure of the Q136A mutant has been determined (32). This mutation caused a dramatic alteration in the β7-β9 loop, which covers the central α helix. Because this mutation retained some activity for rabbit T cells, but lost lethal activity in rabbits, the authors hypothesized that the β7-β9 loop may mediate the ability of the toxin to induce lethality (32). Significantly, residues Tyr<sup>115</sup> and Trp<sup>116</sup> reside in this loop. However, using our current criteria, the Q136A mutation would result in a mitogenic capacity.
of 500,000 pg/ml for rabbit splenocytes, indicating a >1,000-fold reduction in activity. This mitogenic capacity may predict a threshold of SAG activity required for lethal activity induced through cytokine release. This would predict high levels (>2 mg/rabbit) of Q136A to induce T cell-mediated lethal shock. The ability of W116A, E132K, and Q136A to have much stronger mitogenic activity for rabbit cells compared with human cells may also indicate a separate Vβ-chain that is stimulated in the rabbit or may reflect differences in the rabbit equivalent to hVβ2.1. This hypothesis remains a possibility due to the close location of residues Glu^{132} and Glu^{136} in TSST-1 and probably explains the residual activity for rabbit cells seen with these two mutants.

Since many SAGs have affinities for both TCR and MHC in the micromolar range (2), it can be predicted that most SAG trimeric complexes would be unbound at physiological concentrations (43). Andersen et al. (6) have elegantly shown that the TCR α-chain interacts with the MHC β-chain to stabilize the trimeric complex involving SEB. Although, the TCR/SPE C/MHC class II model predicts that in this interaction SAG acts as a bridge without any direct interactions between TCR and MHC class II, SPE C contains a high affinity ($K_d = 4 \times 10^{-8}$ M), zinc-dependent, MHC class II binding domain (19), and small changes in MHC affinity can overcome larger increases in TCR affinity (5). Thus, it appears that the high affinity MHC binding site can circumvent the requirement for stabilizing interactions between the TCR and MHC class II (19). Mutagenesis of SEC3 showed a direct correlation between the affinity of SEC3 for the TCR β-chain and mitogenicity (5). Recently, phage display was used to engineer SEC3 variants in the disulfide loop with increased affinities to the α/β TCR revealing that increasing affinity of the SEC3/TCR complex above that normally seen with SAG/TCR interactions resulted in increased T cell activation (8). These studies each indicate a direct correlation between ligand affinity and potency of bacterial SAG/TCR complexes. This correlation also exists for TSST-1 with hVβ2.1.

For each of the cocrystal structures involving bacterial SAGs (SEB, SEC, SPE A, and SPE C) and their corresponding TCR β-chains (21–23), each SAG binds the TCR in the cleft between the large and small domains, although SPE C engages the TCR in a different orientation compared with SEB, SEC, and SPE A. To illustrate the hVβ2.1 binding domain of TSST-1, Fig. 2 shows ribbon (Fig. 2A) and space-filling (Fig. 2B) diagrams, with residues important for hVβ2.1 interaction highlighted. The six residues of TSST-1 that are individually critical for functional interaction are shown in red. The residue that partially inhibited binding is highlighted in blue, and residues believed to play insignificant or minor stabilizing roles are shown in green. Our mutagenesis data presented here show the location of the hVβ2.1 binding domain of TSST-1 to exist in a groove formed between the central and N-terminal α helices, adjacent to the β7-β8 loop. The location of the hVβ2.1 binding domain of TSST-1 is in sharp contrast to known TCR β-chain binding regions of SEB (Fig. 2C), SPE A (Fig. 2D), and SPE C (Fig. 2E) (22, 23).

**FIGURE 2.** Mapping the functionality of TSST-1 residues to its three-dimensional structure and comparison with other SAG TCR β-chain contact residues. A, Ribbon diagram of TSST-1, highlighting the location and functional effect of mutations. Functionality is color-coded in the following way: TSST-1 mutations that result in abrogation of its complex with hVβ2.1 ($\Delta \Delta G > 2$ kcal/mol) are red, TSST-1 mutations that partially inhibit binding to hVβ2.1 ($1 < \Delta \Delta G < 2$ kcal/mol) are blue, and TSST-1 mutations that do not affect interaction with hVβ2.1 ($\Delta \Delta G < 1$ kcal/mol) are green. B, Mutations mapped to the molecular surface of TSST-1 reveal that residues important for complex formation with hVβ2.1 are predominantly clustered together within the groove formed between two α helices. Color-coding of functionality is the same as in A. Molecular surfaces of SAGs structurally aligned with TSST-1 to indicate the SEB/mVβ8.2 (C), SPE A/mVβ8.2 (D), and SPE C/hVβ2.1 (E) interactions, with contact residues colored red (22, 23). The TCR β-chain binding surface of SEC3 is highly similar to SEB and is not shown (21).
From our data and based on the previous cocystal structure of TSST-1 complexed with HLA-DR1 (16), we have constructed a hypothetical model of the ternary complex (Fig. 3). In general, the TSST-1 ternary model fits the standard wedge model for TCR binding (Fig. 3) (21), yet there are important differences. For the model the most unique region of hVb2.1 (the CDR2 loop) was aligned with the TSST-1 region that has the greatest effect on binding according to the mutagenesis data. The CDR2 loop fits nicely into the pocket outlined in red on the TSST-1 molecular surface (Fig. 2B), and the CDR3 loop wraps around underneath the second α helix and fits into the depression bordered by the N-terminal helix. In this position, CDR2 of the TCR α-chain and the MHC β subunit may result in a small interface similar to that seen in the SEB/SEC3 ternary complexes. Because TSST-1 does not contain the high affinity, zinc-dependent, MHC class II binding domain, it is likely that the TCR/MHC interaction in the TSST-1 complex is necessary for complex stabilization. It is also possible that TSST-1 represents a hybrid ternary complex where (like SEB/SEC3 and unlike SpeC) TSST-1 acts as a wedge between MHC and TCR, allowing for direct TCR α-chain/MHC β-chain contacts, yet unlike the SEB/SEC3 trimeric complexes and similar to the SPE C trimeric complex, the CDR3 loop of hVb2.1 is probably involved in direct SAG-TCR molecular interactions that could contribute to the TCR Vβ specificity of TSST-1. Cocryrstallization of TSST-1 with the TCR β-chain will be necessary to confirm our current results and help further our overall understanding of this important interaction.

**References**


**FIGURE 3.** Hypothetical model of the TSST-1-dependent T cell signaling complex. The model was created by aligning the TSST-1 molecular surface important for hVb2.1 binding with the most unique region of the hVb2.1 structure, the CDR2 loop, a likely scenario considering that TSST-1 activates hVb2.1 T cells exclusively. Colors are as follows: TSST-1, yellow; hVb2.1, red; TCR α-chain, orange; MHC α subunit, green; MHC β subunit, blue; antigenic peptide, gray. This model predicts a possible direct interaction between the TCR α-chain and the MHC β-subunit (intersection of blue and orange domains), much like in the SEB- and SEC3-dependent T cell signaling complexes (6, 22, 44).


