

Staphylococcal Enterotoxin-Like Toxins U2 and V, Two New Staphylococcal Superantigens Arising from Recombination within the Enterotoxin Gene Cluster

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To test the hypothesis that the *Staphylococcus aureus* enterotoxin gene cluster (*egc*) can generate new enterotoxin genes by recombination, we analyzed the *egc* locus in a broad panel of 666 clinical isolates of *S. aureus*. *egc* was present in 63% of isolates, confirming its high prevalence. The archetypal organization of the *egc* locus, consisting of five enterotoxin genes plus two pseudogenes, was found in 409 of 421 *egc*-positive strains. The *egc* locus was incomplete in a few strains and occasionally harbored an insertion sequence and transposase genes. These strains may represent evolutionary intermediates of the *egc* locus. One strain with an atypical *egc* locus produced two new enterotoxins, designated SEV and SEU2, generated by (i) recombination between *selm* and *sei*, producing *selv*, and (ii) a limited deletion in the φ ent1- φ ent2 pseudogenes, producing *selu2*. Recombinant SEV and SEU2 had superantigen activity, as they specifically activated the T-cell families V β 6, V β 18, and V β 21 (SEV) and V β 13.2 and V β 14 (SEU2). Immunoscope analysis showed a Gaussian CDR3 size distribution of T-cell receptor V β chain junctional transcripts of expanded V β subsets in toxin-stimulated cultures, reflecting a high level of polyclonality. These data show that *egc* is indeed capable of generating new superantigen genes through recombination.

Staphylococcus aureus produces a large variety of exotoxins, including staphylococcal enterotoxins A to E (SEA to SEE), SEG to SER, and SEU; staphylococcal enterotoxin-like toxins (SEIs); and toxic shock syndrome (TSS) toxin-1 (5, 23). These toxins are responsible for specific acute clinical syndromes such as TSS (due to both TSS toxin-1 SEs and SEIs), food poisoning (due to SEs), and staphylococcal scarlet fever (considered a mild form of TSS) (10, 26, 34).

All these toxins share certain structural and biological properties, suggesting that they derive from a common ancestor (16, 21). They exhibit superantigen activity, stimulating polyclonal T-cell proliferation through coligation between major histocompatibility complex class II molecules on antigen-presenting cells (APC) and the variable portion of the T-cell antigen receptor β chain or α chain (TCR V β and TCR V α , respectively), with no need for prior APC processing (4, 13, 21, 22, 37, 39). The pattern of V β /V α activation is specific to each superantigen (4, 12). T-cell/APC activation by these toxins leads to the release of various cytokines/lymphokines and interferon, enhances endotoxic shock, and causes T- and B-cell immunosuppression, all of which may undermine the immune response against bacterial infection (5, 10, 25).

All the genes encoding these toxins are harbored by mobile elements, including bacteriophages, pathogenicity islands, genomic islands, and plasmids (10, 20, 28, 36). Only the en-

terotoxin gene cluster (*egc*) is organized as an operon, consisting of two enterotoxin genes (*seg* and *sei*), three enterotoxin-like genes with proven superantigenic activity but not emetic properties (*selo*, *selm* and *seln*), and two pseudogenes (φ ent1 and -2). This organization suggests that *egc* arose through gene duplication and variation from an ancestral gene and that gene recombination created variant toxins with different biological activities (7, 12). SEs and SEIs can be divided into three phylogroups, each of which contains one or more *egc*-encoded toxins, suggesting that all SEs and SEIs potentially derive from the *egc* locus. Several allelic variants of *egc* toxin in clinical, animal, and food isolates of *S. aureus* have been described. At least four SEG variants and three SEI variants have been described, and most bear key amino acids involved in TCR and major histocompatibility complex interactions (3, 6, 12, 15, 27). Letertre et al. have described a new SE-like toxin, designated SEU, that apparently arises from fusion between φ ent1 and φ ent2, which itself results from a 15-nucleotide (nt) deletion in φ ent1 (15). Note that SEU should be designated “SEIU,” as its emetic properties have not yet been demonstrated.

The aim of the present study was to further examine the possibility that *egc* can generate new enterotoxin genes by recombination and that the new enterotoxins thereby have superantigen activity. For this purpose we analyzed the *egc* locus in a broad collection of clinical isolates for signs of evolutionary intermediates and new *egc* toxins. The activities of the new recombinant enterotoxins were investigated.

MATERIALS AND METHODS

Bacteria and culture. A panel of 666 *S. aureus* clinical isolates were selected from the collection of the French National Reference Center for Staphylococci

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(Lyon, France) between 2001 and 2003. They had been isolated from patients with suppurative infections (arthritis, skin infection, pneumonia, or infective endocarditis), acute toxemia (TSS, staphylococcal scalded skin syndrome, or staphylococcal scarlet fever), or asymptomatic nasal carriage. All isolates were identified as *S. aureus* by their ability to coagulate citrated rabbit plasma (Pasteurex-Staph-Plus; Sanofi Diagnostics Pasteur GmbH, Freiburg, Germany) and their catalase and DNase activities (Toluidine; Bio-Rad, Marnes-la-Coquette, France). *S. aureus* strain A900322 was used as the *egc*⁺ reference strain (*sea*⁻, *seb*⁻, *sec*⁻, *sed*⁻, *see*⁻, *seg*⁺, *seh*⁻, *sei*⁺, *selm*⁺, *seln*⁺, *selo*⁺, *φent1*⁺, *φent2*⁺, *selp*⁺, *lukD/E*⁺) (12). *S. aureus* strain RN4220 was used as a negative control for SE genes.

DNA extraction and purification. All strains were grown in brain heart infusion at 37°C overnight, and DNA was extracted with the standard phenol-chloroform procedure (35).

PCR detection of *egc* toxin genes. Sequences specific for *egc* toxin genes (*selo*, *selm*, *sei*, *seln*, and *seg*) were detected by PCR as previously described (12). Amplification of *gyrA* was used to confirm the quality of each DNA extract and the absence of PCR inhibitors. The PCR products were resolved by electrophoresis through 0.8% agar gel (Sigma, St. Louis, Missouri). The location of *egc* in the chromosome was determined by PCR with one primer (*seloY*) located in *egc* and another primer (*hemY*) located 2.3 kb upstream of the *egc* locus (11).

Cloning and sequencing of *egc* variants. The Clontech Genome Walker kit (Ozyme; Montigny-Le-Bretonneux, France) was used to identify the flanking regions in selected *egc* variants, following the supplier's instructions, with enzymes and specific primers for each strain and target gene (Table 1).

Production and purification of recombinant enterotoxins. Primers were designed following the identification of suitable hybridization sites in *selv* and *selu2* of strain A900624. DNA was extracted and used as a template for PCR amplification and recombinant protein production.

The 5' primer was placed in the coding region of *selm* for *selv* (R-SEIV 1) and of *φent1* for *selu2* (R-SEIU2 1), omitting the region predicted to encode the signal peptide (<http://www.cbs.dtu.dk/services/SignalP/>). The 3' primer (PstI restriction site) was chosen to overlap the stop codon of *seln* for *selv* (R-SEIV 2) and of *φent2* for *selu2* (R-SEIU2 2) (Table 1; see Fig. 2). PCR products were codigested with appropriate restriction enzymes (Promega, Madison, Wisconsin), purified with the High Pure PCR product purification kit (Boehringer Mannheim, Meylan, France), and ligated (T4 DNA ligase; Boehringer Mannheim, Meylan, France) in the pQE-30 expression vector (QIAGEN) digested with the same restriction enzymes. The resulting plasmid was transformed into *Escherichia coli* strain M15. Open reading frame (ORF) integrity was verified by sequencing the junction between pQE-30 and the insert. *E. coli* was grown overnight in Luria-Bertani (LB) broth with appropriate antibiotics (ampicillin, 100 µg/ml; kanamycin, 25 µg/ml). Overnight cultures were transferred (1/10) into 1 liter of LB broth containing antibiotics and were incubated until the optical density at 540 nm reached 0.5 to 0.6. Protein expression was induced by adding 1 mM isopropylthio-β-D-galactoside. After 4 hours, bacterial cells were harvested by centrifugation at 13,000 rpm for 20 min at +4°C. The cell pellets were resuspended in lysis buffer and kept at -20°C overnight. Cell debris were removed by centrifugation at 14,000 rpm for 15 min at +4°C. The recombinant protein (His₆ tag) was purified from cell lysates on affinity chromatography columns according to the supplier's instructions (New England Biolabs). The protein content was determined by the Bradford method (12) as modified by Bio-Rad Laboratories (Richmond, California), using bovine serum albumin (Sigma, Steinheim, Germany) as a standard. Protein purity was verified by sodium dodecyl sulfate-polyacrylamide gel electrophoresis.

T-cell activation assay. Specific activation of T cells by SEIV and SEIU2 toxins was measured as previously described (13). Briefly, 50 µl of fresh blood from healthy donors was incubated with 50 µl of RPMI medium, either alone (negative control) or supplemented with each toxin (from 1 µg/ml to 10 ng/ml) or with 10 µg/ml phytohemagglutinin (PHA) (positive control) for 18 h at 37°C in humidified 5% CO₂-air. The cell suspension was then treated with ammonium chloride lysis buffer and centrifuged to eliminate erythrocytes. The remaining peripheral blood mononuclear cells (PBMC) were incubated for 20 min at 4°C with anti-CD3, anti-CD4, and anti-CD69 monoclonal antibodies. Activated cells were detected by CD69 staining. The cells were treated with lysis buffer, resuspended in 300 µl of sterile phosphate-buffered saline (Gibco, Invitrogen, Paisley, United Kingdom), and analyzed with a flow cytometer (FACScan; Becton Dickinson, Mountain View, CA). Data were analyzed with LYSIS II software.

Flow cytometric analysis of Vβ TCR-triggered PBMC proliferation. PBMC were isolated from heparinized venous blood of healthy donors by Ficoll density gradient sedimentation (PANCOLL; PAN Biotech GmbH, Aidenbach, Germany). The cells were washed three times in Hanks balanced salt solution (Sigma-Aldrich) and suspended in RPMI 1640 culture medium (Gibco, Invitrogen

TABLE 1. Primers used for detection of *egc* genes and for SEIU2 and SEIV cloning and sequencing

Primer	Location within <i>egc</i> ^a	Oligonucleotide sequence (5'→3') ^b
primer5'	1-19	GTC CCG TTA GGA GTC ATA C
seo1	481-506	AGT TTG TGT AAG AAG TCA AGT GTA GA
seo2	630-660	ATC TTT AAA TTC AGC AGA TAT TCC ATC TAA C
seo3	427-449	GCA TTG TTT ACA CTA CAT ATT GC
seo4	244-270	CTG TTT GTT CAA TAG TAA GTA GGA TTG
seo5	557-581	GTT GAT ACA ATT GAT TTT ACT GTC G
invseo2	630-660	GTT AGA TGG AAT ATC TGC TGA ATT TAA AGA T
invsem1	1785-1811	GTT CTC CAT TAA CCC AAA GAT TAA TAG
sem1	1785-1811	CTA TTA ATC TTT GGG TTA ATG GAG AAC
sem2	2085-2110	TTC AGT TTC GAC AGT TTT GTT GTC AT
invsei1	2260-2281	CCT ACA CCA ATA TCA CCT TGA G
sei1	2260-2281	CTC AAG GTG ATA TTG GTG TAG G
sei2	2886-2915	GTT ACT ATC TAC ATA TGA TAT TTCGAC ATC
invsei2	2886-2915	GAT GTC GAA ATA TCA TAT GTA GAT AGT AAC
yent1	3352-3375	ACG TAG ATT TGT TTG GGA CAA ACT
yent2	3502-3529	GTG CTG TTA TGT TTT TCT TAT TAG TAG G
invsen1	3969-3988	GAC TCG TCT AAT TGC CAC GT
sen1	3969-3988	ACG TGG CAA TTA GAC GAG TC
sen2	4415-4444	GAT TGA TCT TGA TTA TGA GAA TGA AAG
invsen2	4415-4444	CTT TCA TTC TCA TAA TCA TCA AGA TCA ATC
invseg1	4979-5003	CAT TAC ATT ACC CAT AGT TCC CTT A
seg1	4979-5003	TAA GGG AAC TAT GGG TAA TGT AAT G
seg2	5514-5541	GAA CAA AAG GTA CTA GTT CTT TTT TAG G
invseg2	5514-5541	CCT AAA AAA GAA CTA GTA CCT TTT GTT C
primer3'	6163-6189	CTT TAA CTT CAT AAA TTA GCA GTA GTC
pQE-F ^c		CCC GAA AAG TGC CAC CTG
PQE-R ^c		GTT CTG AGG TCA TTA CTG G
R-SEIU2-1		CA <u>GGATCC</u> ATG TTA AAT GGC AAT CCT AAA C CA
R-SEIU2-2		GC <u>CTGCAG</u> TTA TTT TTT GGT TAA ATG AAC TTC TAC ATT AAT AGA TTT A
R-SEIV 1		GCA <u>GGATCC</u> GAT GTC GGA GTT TTG AAT CTT AGG
R-SEIV 2		TAA <u>CTGCAG</u> TTA GTT ACT ATC TAC ATA TGA TAT TTC GAC ATC

^a Location based on the *S. aureus egc* sequence (GenBank accession number AF9285760).

^b Restriction sites for BamHI and PstI are underlined.

^c Primer used for sequencing the inserted gene in the pQE-30 vector.

Corporation) supplemented with 10% heat-inactivated fetal bovine serum (Gibco, Invitrogen Corporation), 20 mM HEPES buffer, 2 mM L-glutamine (Sigma-Aldrich), 100 µg/ml streptomycin, and 100 µg/ml penicillin (Gibco-Invitrogen Corporation). Cells (2 × 10⁶ to 5 × 10⁶/ml) were cultured for 3 days with each toxin (500 ng/ml) in 24-well plates (Falcon Becton-Dickinson) and then washed in Hanks balanced salt solution and suspended in culture medium con-

taining increasing concentrations (20 to 100 U/ml) of human interleukin-2 (Eurobio, France) for 12 to 14 days. The V β profile was then determined by flow cytometry using the IOTest Beta Mark (Immunotech, Marseille, France), according to the supplier's instructions. The multiparameter data files were analyzed with the Cellquest program (Becton Dickinson). Cells incubated in culture medium or stimulated with PHA (10 μ g/ml) were used as negative and positive controls, respectively. For kinetic experiments, cells were cultured for 10 days and the V β profile was determined on days 0, 1, 3, 4, 6, and 10 by using the same procedure.

Quantitative Immunoscope analysis. Total RNA was extracted with the GenElute Total RNA Miniprep kit (Sigma-Aldrich, St. Quentin Fallavier, France) as recommended by the manufacturer. RNA (50 μ l) was reverse transcribed with oligo(dT)₁₇ and 400 U of SuperScript II RNase H reverse transcriptase (Invitrogen, Cergy Pontoise, France). An aliquot of cDNA synthesis reaction mix was amplified with each of the 24 TCR V β family-specific primers, together with a TCRC β primer and a fluorochrome-labeled nested oligonucleotide TaqMan probe for TCRC V β . Real-time PCR was carried out in an ABI7300 system (Applied Biosystems, Courtaboeuf, France). All quantitative PCRs took place in a total volume of 25 μ l with 1 \times TaqMan buffer (Applied Biosystems), 25 mM of AmpliTaq, 5 mM MgCl₂, 0.2 mM deoxynucleoside triphosphates, 400 nM each primer, and 200 nM probe. The thermal cycling conditions consisted of 95°C for 10 min followed by 95°C for 15 seconds and 60°C for 1 min for 40 cycles. The relative usage of each TCR V β segment was computed as described elsewhere (17). Two microliters of each amplification product was used as template in a runoff reaction initiated with a nested TCRC β fluorescent primer in a total volume of 10 μ l with 1 \times polymerase buffer (Promega, Charbonnières, France), 25 mM of Taq DNA polymerase, 3 mM MgCl₂, 0.2 mM deoxynucleoside triphosphates, and 0.1 mM nested TCRC β fluorescent primer. The fluorescent products were then separated and run on an automated 3730 DNA sequencer (Applied Biosystems) and analyzed with Immunoscope software (32) in order to profile the TCR V β repertoire.

Detection of bacterial RNA by RT-PCR. Total RNA was extracted from staphylococcal cultures by using RNeasy spin columns (QIAGEN, Courtaboeuf, France). cDNA was synthesized using Ready-To-Go reverse transcriptase PCR (RT-PCR) beads (Pharmacia Biotech, Orsay, France) by incubating 0.1 μ g of total RNA with the following pairs of primers: primer5' and seo3, seo4 and seo5, seo1 and seo2, invseo2 and invsem1, sem1 and invsei1, sei1 and sei2, invsei2 and yent2, yent1 and invsen1, sen1 and sen2, invsen2 and invseg1, seg1 and seg2, and invseg2 and primer3' (Table 1). The reaction mixtures were incubated with each primer pair described above at 42°C for 30 min for reverse transcription, followed by 30 cycles of amplification (1 min of denaturation at 94°C, 1 min of annealing at 55°C, and 1 min of extension at 72°C). The RT-PCR products were then analyzed by electrophoresis through a 1% agarose gel. RNA extracts were tested for DNA contamination by preincubating the reaction mixtures at 95°C for 10 min to inactivate reverse transcriptase prior to the RT-PCRs.

RESULTS

Detection of incomplete *egc*. Among the 666 isolates tested by PCR for the presence of the *egc* toxin genes, 421 isolates were positive for at least one gene. The *egc* was complete (including *selo*, *selm*, *sei*, *seln* *seg*, and the two pseudogenes ϕ ent1 and ϕ ent2) in 409 isolates and was atypical in 12 isolates. Seven of these 12 atypical isolates harbored five *egc* toxin genes, one had four, one had three, one had two, and two had one (Table 2). In order to analyze the most common *egc* variations, we selected for further investigation 6 of these 12 isolates (A950227, A980341, A940440, A900624, LY19991222, and LY1991287), representing each of the above *egc* variations. The chromosomal location of *egc* in these six strains was investigated by using a PCR primer annealing to the *hemY* gene located 2.3 kb upstream of the *egc* locus and another primer located within *seo*. PCR was positive with all six strains and with the *egc*⁺ control strain, indicating that the *egc* loci were all located in the same chromosomal region. The complete sequence of each of the genes and their arrangement within the *egc* locus were determined, and the results were compared with those for the *egc* locus of the reference strains

TABLE 2. Distribution of the *egc* genes in the 666 *S. aureus* isolates

Category and strain	<i>egc</i> gene content ^a						Additional genes ^b	n (%)
	<i>selo</i>	<i>selm</i>	<i>sei</i>	<i>ent1-ent2</i>	<i>seln</i>	<i>seg</i>		
Strains with complete <i>egc</i> cluster	+	+	+	+	+	+		409 (61.4)
Strains with 5 <i>egc</i> genes or pseudogenes								7 (1.05)
HT20011243	+	+	+	+	+	-	ND	
HT2000480	+	+	+	+	+	-	ND	
LY1991222	+	+	+	+	+	-	TC	
A950227	+	+	+	+	+	-	TC	
LY1990182	+	+	+	+	+	-	ND	
A980341	+	+	-	+	+	+	TC	
LY1991287	+	+	+	+	-	+	TC	
Strain with 4 <i>egc</i> genes or pseudogenes, A950062	+	-	+	+	+	-	TC	1 (0.15)
Strain with 3 <i>egc</i> genes or pseudogenes, A900624	+	-	-	-	+	+	<i>selu2</i> , <i>selv</i>	1 (0.15)
Strain with 2 <i>egc</i> genes, A940440	+	+	-	-	-	-	TC	1 (0.15)
Strains with 1 <i>egc</i> gene								2 (0.3)
HT2000461	+	-	-	-	-	-	ND	
HT2000458	+	-	-	-	-	-	ND	
Total strains with at least 1 <i>egc</i> gene								421 (63.2)
Total strains with no <i>egc</i> gene or pseudogene								245 (36.7)

^a The *egc* genes are represented in their order of transcription (from left to right). *seo* was previously designated *sel*; *sen* was previously designated *sek*. The nomenclature for the other genes is unchanged (12).

^b ND, not determined; TC, transposase cassette.

A900322 and Mu50 (1). Schematic representation of the *egc* loci in these variants (Fig. 1) showed that the overall organization and sequence of the archetypal *egc* locus were conserved. However, we observed two types of genetic events that generated variations in the *egc* locus: (i) two types of insertion sequence (IS) and (ii) new enterotoxin genes generated by homologous recombination between *egc* genes or pseudogenes.

Insertion sequences. The first IS was detected in four strains (Fig. 1) and in three different genes (*sei*, *seln*, and *seg*). It was composed of 0.757 to 1.222 kb of DNA containing two very short terminal inverted repeats and two additional open reading frames (ORF 1 and ORF 2, of 282 and 807 bp, respectively). ORF 1 encoded 93 amino acids (aa) showing 96% sequence identity (91/93 aa) to transposase 8 (Tnase 8) (from the methicillin-resistant *S. aureus* [MRSA] 252 genome) and is referred to below as transposase 8-like. ORF 2 encoded a 268-aa peptide that had 95% sequence identity (257/268 aa) with MRSA 252 transposase, 96% (182/188 aa) with the truncated transposase from strain Mu50, and 88% (198/224 aa) with the truncated transposase from strain MW2. This protein belonged to the Rve family, which corresponds to the catalytic domains of numerous transposases and is referred to below as Rve-like protein. This cassette (Tnase 8-like/Rve-like) was in the same orientation as the *egc* operon in strains A950227,

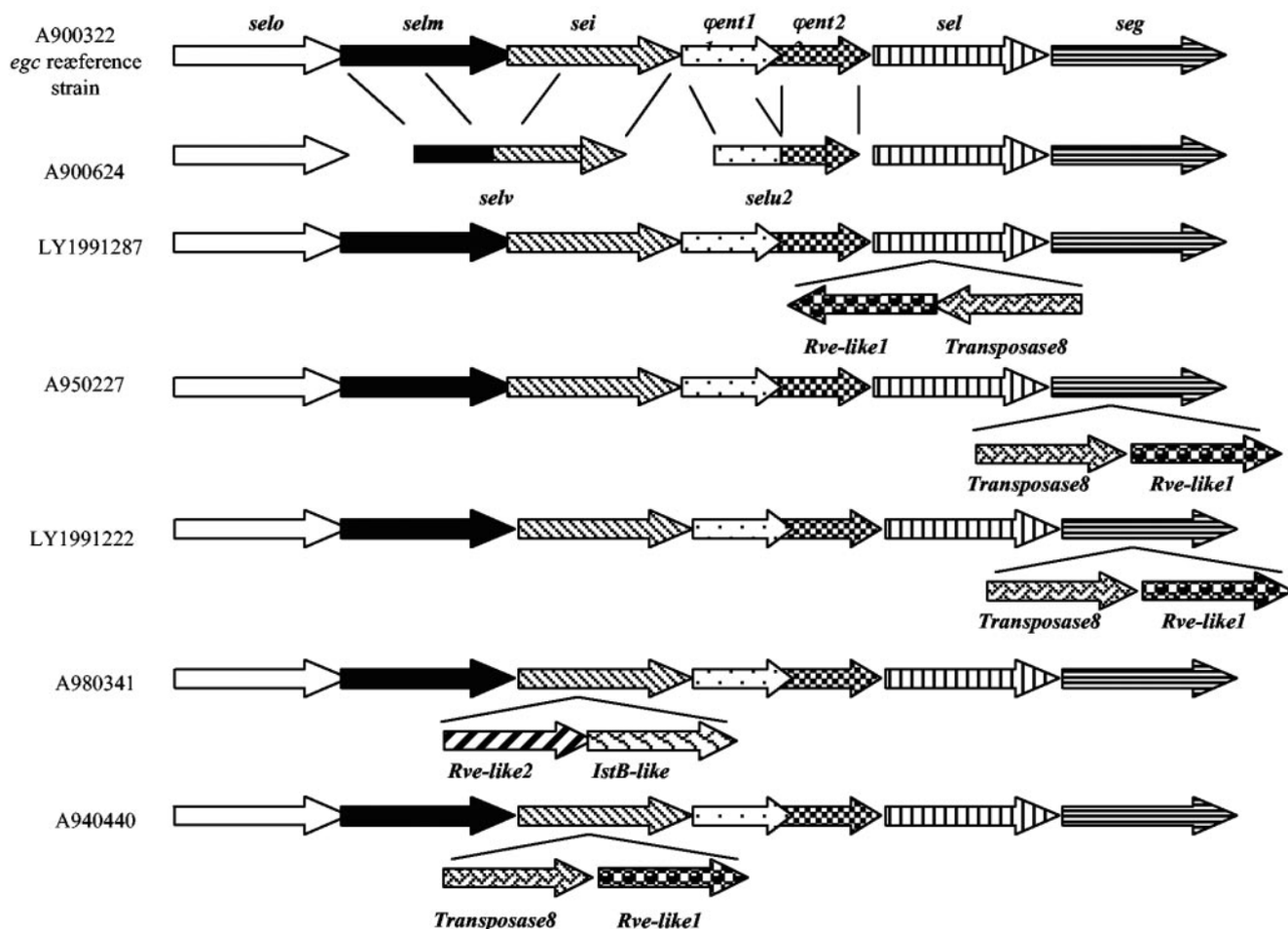


FIG. 1. Organization of the *egc* locus in reference and atypical strains. A900322 (GenBank accession number AFP285760) is the *egc* reference strain. A900624 harbors two new enterotoxin genes, *selu2* and *selv*. LY1991287, A950227, LY1991222, A980341, and A940440 have an insertion sequence (Rve-like 1–transposase 8 or Rve-like 2–IstB-like). These sequences are shown schematically below the genes into which they are inserted.

LY991222, and A940440 and in the opposite orientation to that in strain LY991287 (Fig. 1). The intergenic sequence of 41 nucleotides between the Tnase 8-like and Rve-like genes was strictly identical in the four strains, while the stretch between the Tnase 8-like gene and the next *egc* gene varied in sequence and length (11 to 110 nt), as did the stretch between the Rve-like gene and the next *egc* gene (16 to 17 nt).

A putative –10 and –35 promoter sequence (GATTTT-N3-TATTGT-N27-ATG) was found upstream of the Tnase 8-like start codon in three of the four strains. In addition, the Tnase 8-like and Rve-like genes had appropriate Shine-Dalgarno sequences. In three strains the direct repeats corresponding to the insertion site of the Tnase 8-like/Rve-like cassette were ATTT, AAGG, and CATGAT. These nucleotide motifs were detected 102, 19, and 5 times, respectively, in the *egc* sequence of the prototype strain (GenBank accession number AFP 285760).

The second type of IS was detected only in strain A980341 (Fig. 1). It was composed of a 2.122-kb DNA sequence containing two terminal inverted repeats of 8 bp and 14 bp and two ORFs (ORF 3 and ORF 4) of 1,197 and 768 bp, respectively.

These ORFs encoded 398 aa and 255 aa, respectively. Comparison of the deduced amino acid sequences of these ORFs with translated sequences from GenBank showed that the putative translation product of ORF 3 had 55% identity (220/393 aa) to the putative integrase core domain (Rve) of *Enterococcus faecium* and to the transposase conserved domain and 54% identity (213/393 aa) to the transposase-like protein of *Lactobacillus collinoides*. This protein was designated Rve-like 2. The peptide sequence encoded by ORF 4 was homologous to the putative IstB-like ATP-binding protein and the DNA replication protein conserved domain and had 52% identity (127/242) to a transposition helper protein of *E. faecium*. It was designated IstB-like protein. A putative –10 and –35 promoter sequence (TATTTT-N5-ATGACA-N26-ATG) was found upstream of the Rve-like 2 start codon. In addition, Rve-like 2 and IstB-like had an appropriate Shine-Dalgarno sequence. The direct repeat sequence CTTCA, corresponding to the insertion site of the Rve-like 2/IstB-like cassette, was detected only once in the *egc* sequence of the prototype strain.

Two new enterotoxin genes generated by *egc* gene recombination and/or deletion. Two additional ORFs (ORF 5 and

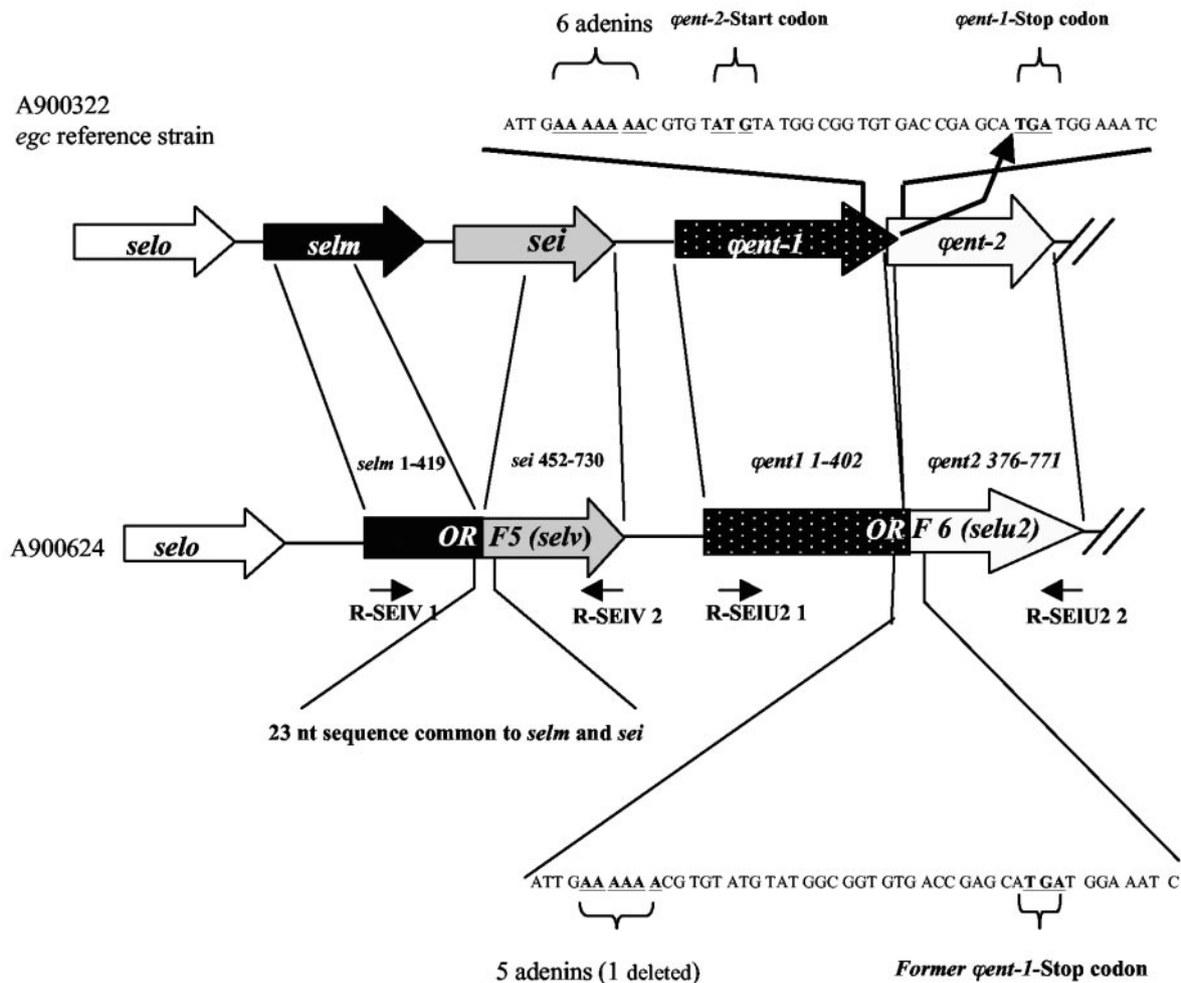


FIG. 2. Genetic organization of the strain A900624 *egc* locus in comparison with the archetypal *egc* locus of reference strain A900322. Black arrows indicate the locations of primers used for the SEIV (R-SEIV 1 and R-SEIV 2) and SEU2 (R-SEIU2 1 and R-SEIU2 2) recombinant strategy.

ORF 6) of 720 and 771 bp, respectively, both related to staphylococcal enterotoxins, were found in strain A900624 (Fig. 2). ORF 5 had 99% identity (424/428) to the 5' end of the *selm* gene (nt 1 to 419) and almost 100% identity (300/301) to the 3' end of the *sei* gene (nt 452 to 730) of the reference strain. Thus, ORF 5 appears to result from a recombination event between *selm* and *sei*. Interestingly, a 23-nucleotide sequence (TAGTAACAGCTCAAGAAATTGAT; nt 418 to 441) overlapping the junction region was shared by the *selm* and *sei* genes.

Analysis of the deduced amino acid sequence of ORF 5 showed the same features, with 97.2% identity (139/143) of the first 143 aa with the N-terminal part of SE/M and 100% identity of aa 144 to 239 with the C-terminal part of SEI. Globally, however, ORF 5 shared only 87.5% amino acid identity with the SEM and less with other SEs/SEIs. Based on the recently described nomenclature for staphylococcal superantigens (18), ORF 5 was designated "selv" and the corresponding protein "SE-like V" (SEIV).

ORF 6 appears to result from another type of molecular reorganization. Analysis of the DNA sequence showed that the first part of this ORF (nt 1 to 402) had 99.3% identity (399/402) to the ϕ_{ent1} pseudogene of strain Mu50 and that the

second part (nt 376 to 771) had 100% identity to the ϕ_{ent2} pseudogene of the same strain. As with ORF 5, the junction region of ORF 6 corresponded to a 25-nt conserved sequence between the 3' end of ϕ_{ent1} and the 5' end of ϕ_{ent2} (Fig. 2). ORF 6 contained a single adenine deletion (A 365) very close to the end of ϕ_{ent1} , which abolished the ochre stop codon present in the end of ϕ_{ent1} . This generated a 256-aa translation product corresponding to fusion of ϕ_{ent1} and ϕ_{ent2} . Analysis of the deduced amino acid sequence showed strong similarity with SE/U (94% identity) and moderate similarity with SEC1 and SEC3 (53% identity). The gene was designated *selu2* and its product SE/U2. The putative Shine-Dalgarno sequences of *selu2* and *selv* are, respectively, TGGAGT and GGAGAA. To verify that ORFs 5 and 6 correspond to transcribed genes in strain A900624, RT-PCR analyses using primers specific for these two genes were performed. In both cases a positive RT-PCR signal was obtained and the signal intensity was comparable to that of *sen* (Fig. 3).

Superantigen activities of recombinant SEIV and SEU2. Purified recombinant SEIV and SEU2 expressed in *E. coli* were first studied for their ability to induce T-cell activation in the CD69 assay (19). When incubated overnight with PBMC,

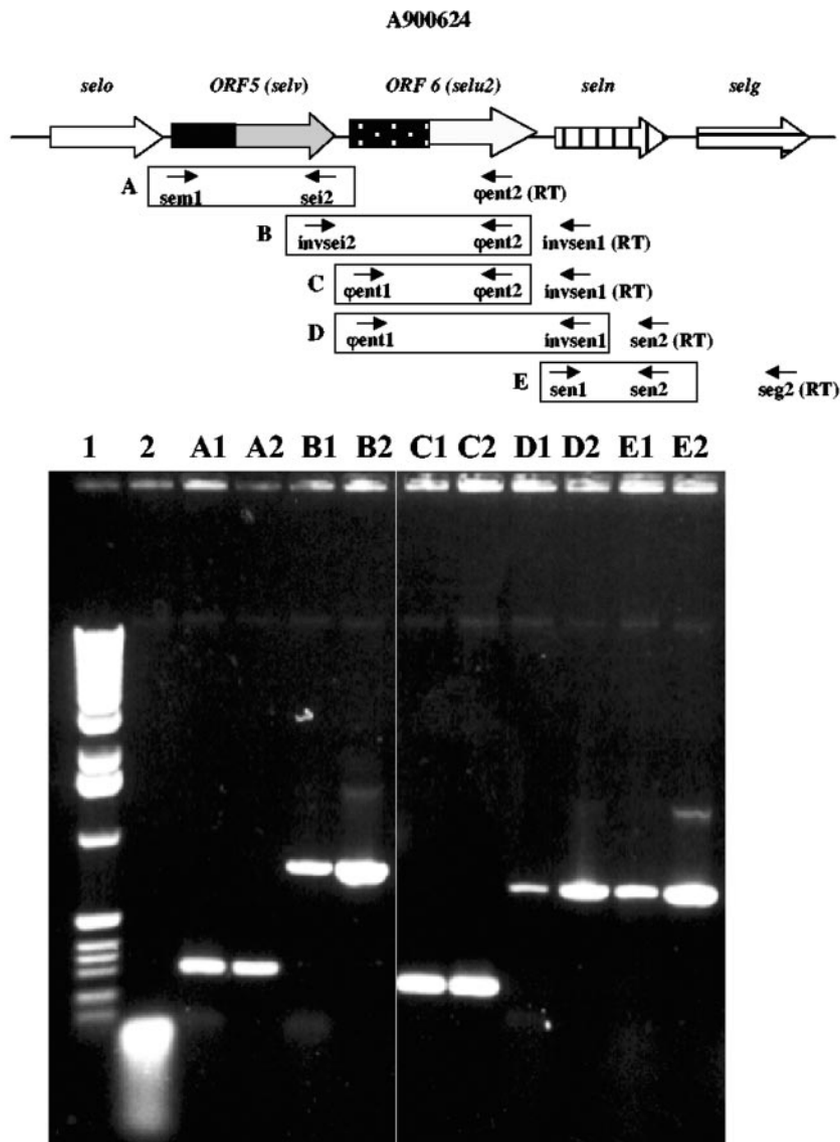


FIG. 3. Analysis of *selv* and *selu2* transcripts by RT-PCR. cDNA was prepared from *S. aureus* A900624 total RNA using the RT primers shown (RT), followed by PCR using primer pairs A to E (boxed). Lanes A to E, results obtained using the corresponding primer pairs. Lane 1, 1-kb molecular size marker; lane 2, RT-PCR negative control (heat inactivation of reverse transcriptase); lanes A1 to E1, RT-PCR from extract of A900624; lanes A2 to E2, PCR positive control (A900624 DNA as template).

both toxins induced concentration-dependent CD69 expression on CD3⁺ T cells, with the proportions of CD3⁺ CD69⁺ cells ranging from 1.5% to 11.8% (not shown) (PHA [48.6%] and RPMI medium [0.5%] were used as positive and negative controls, respectively).

The toxins were then examined for their ability to induce TCR Vβ-selective expansion of T cells in 12-day PBMC culture. As shown in Table 3, SE/U2 induced selective expansion of Vβ 14 and Vβ 13.2 T-cell subsets, while SE/V induced Vβ 18 and Vβ 21.3 expansion. However, the extent of Vβ 18 expansion varied from one donor to another.

To document the TCR Vβ composition of superantigen-stimulated T cells and the repertoire diversity of the expanded TCR Vβ subsets, the size distribution of PCR-amplified TCR

Vβ chain junctional products was studied by using the Immunoscope method (14–16). This method enables the quantification of each of the TCR Vβ usage frequencies within total T cells and the visualization of the TCR third complementary region (CDR3) length distribution for each Vβ–Jβ rearrangement. Therefore, it gives a representation of the diversity of the TCR repertoire. Polyclonal populations show a typical Gaussian distribution of the CDR3 lengths, whereas oligoclonal populations are characterized by the presence of prominent peaks. TCR Vβ quantitative Immunoscope analysis, shown in Fig. 4, confirmed that SE/V triggered the expansion of the Vβ 18 and Vβ 21 T-cell subsets. In addition, it showed the expansion of Vβ 6b cells, a subset which is not covered by the IOTest Beta Mark method. As shown in Table 3, TCR Vβ

TABLE 3. FACS and RT-PCR analysis of PBMC from three donors cultured for 12 days in the presence of SE/U2 or SE/V

Sample	TCR Vβ ^a	% of total CD3 ⁺ cells in culture ^b for donor:					
		A		B		C	
		FACS	RT-PCR	FACS	RT-PCR	FACS	RT-PCR
Day 0	13	4.52	4.5	1.33	8.8	1.35	3.9
	14	5	1.1	1.6	0.2	0.71	0.2
	18	3.1	4.6	1.6	4.9	1.3	3.6
	21	3.22	9.4	3.34	1.6	3.28	7.6
PHA ^c	13	3.12	10.1	1.03	4.1	1.13	11.3
	14	4.68	1.1	3.57	1.2	2.7	0.6
	18	0.42	6.9	0.5	9.2	0.64	4.6
	21	2.22	8.6	1.4	9.8	1.84	8.1
Se/U2	13	16.2	4.7	7.46	11.6	3.9	8.9
	14	54.28	56.4	76.37	68.3	71.39	48.8
	18	0.46	0.9	0.11	0.9	0.03	0.8
	21	1.28	2	2,13	0.2	1.47	1
Se/V	13	1.96	7	1.76	9	0.67	4.3
	14	3.9	0.1	1.23	0.6	1.9	0.3
	18	5.04	14.8	1.81	14.8	1.3	28.3
	21	5.76	30.5	7.22	37.8	5.4	40.8

^a 13, Vβ 13.2 in FACS assay and Vβ 13a in RT-PCR quantification assay; 21, Vβ 21.3 in FACS assay and Vβ 21f in RT-PCR quantification assay.

^b Values in boldface indicate specific expansions.

^c Four-day PHA stimulation.

quantitative Immunoscope analysis showed, for all the donors, a constant and significant expansion of Vβ 18-expressing T cells following stimulation with SE/V. This was confirmed by fluorescence-activated cell sorter (FACS) analysis for donor A but hardly for donors B and C. Based on the clear-cut RT-PCR data, we believe that this apparent discrepancy is linked to the poor reactivity of the Vβ 18-specific antibody included in the IOTest. Regarding SE/U2, Immunoscope analysis confirmed the expansion of Vβ 14 cells but not of Vβ 13 cells. To investigate this apparent discrepancy, we performed early kinetic studies (from day 1 to day 10 of stimulation with SE/U2), in which FACS analyses were carried out on both small lymphocytes and blasts (gated according to forward scatter/side scatter criteria). As shown in Fig. 5, SE/U2 triggered the expansion of Vβ 13.2 cells, as Vβ 13.2 blasts were detected from day 3 to day 10, with a peak at day 4, while Vβ 13.2 small lymphocytes were detected later (day 6 to at least day 10). These kinetic studies also confirmed that Vβ 14 cells were triggered by SE/U2, as BLAST expansion was detected after as little as 1 day of stimulation, while small-lymphocyte expansion was detected only on day 8. It is noteworthy that both the kinetics and the proportions of Vβ 14 and Vβ 13 expansion triggered by SE/U2 were markedly different, with the proportion of the Vβ 13 BLAST cells being almost nine times smaller than the proportion of the Vβ 14 cells (9% versus 80% in this experiment).

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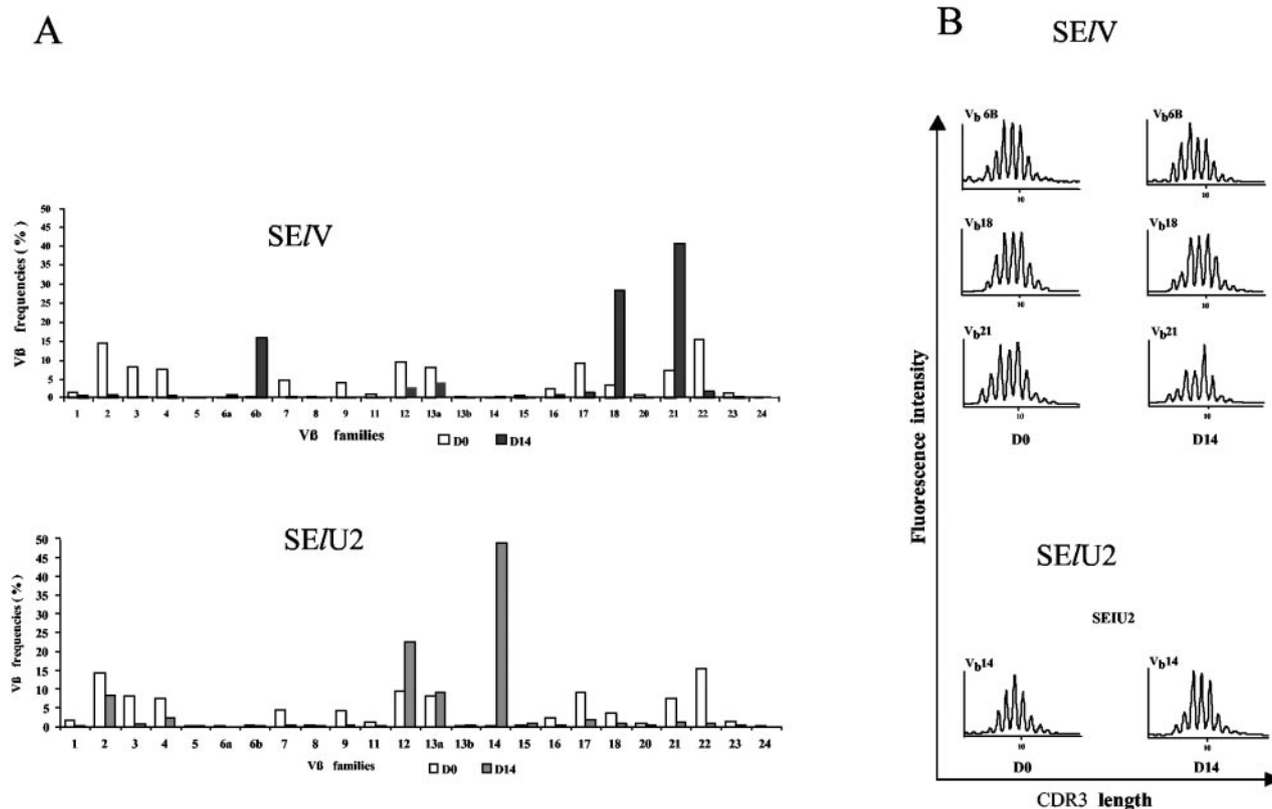


FIG. 4. TCR Vβ repertoire analysis of superantigen-stimulated PBMC by the Immunoscope approach. (A) Quantitative Vβ repertoire determined by real-time PCR analysis on day 0 (D0) and day 14 (D14) following stimulation with SE/V or SE/U2. The x axis indicates Vβ families and the y axis their relative frequency of usage. Selective Vβ expansion was considered to occur when the D14/D0 Vβ frequency ratio was >3. (B) Immunoscope profiles of the fluorescent Vβ-Cβ runoff products obtained with 14-day-stimulated mononuclear cells. The x axis indicates the CDR3 length. Only profiles corresponding to Vβ expansion are shown.

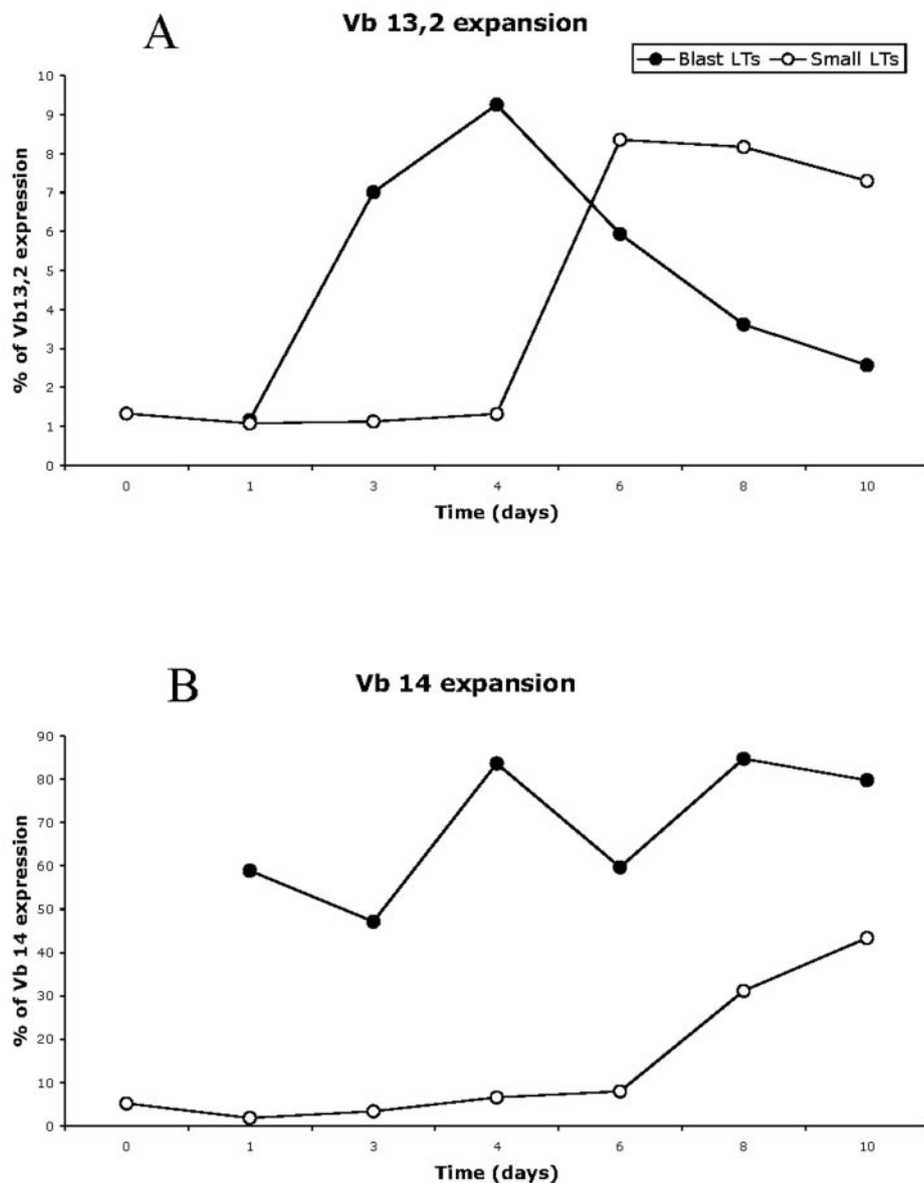


FIG. 5. Flow cytometric kinetic analysis of T cell stimulation by SE/U2. Time courses of V β 13.2 (A) and V β 14 (B) expression in blast cells (closed circles) and small cells (open circles), identified with forward scatter/side scatter criteria, after stimulation of PBMC with 500 ng/ml SE/U2 are shown. V β expansion was determined by flow cytometry. Data are percentages of CD3 T cells expressing the corresponding V β chains.

Similar kinetic experiments were performed for TCR V β quantitative Immunoscope analysis (Fig. 6). V β 14 expansion was detected from day 1 (6.1%, versus 0.5% at day 0) and was still high on day 6 (8.8%). In contrast, no specific V β 13a or V β 13b expansion was detected.

Immunoscope analysis of the CDR3 size distribution of TCR V β chain junctional transcripts within expanded V β subsets showed Gaussian profiles in all toxin-stimulated cultures, reflecting a high level of polyclonality (Fig. 4). This was further confirmed by sequence analysis of TCR V β junctional transcripts derived from some expanded TCR V β subsets (not shown). Taken together, these TCR V β repertoire studies confirmed the superantigenic nature of the two new toxins described here.

DISCUSSION

The aim of this study was to test our hypothesis that the *S. aureus* *egc* locus is a "superantigen nursery." We obtained two major lines of evidence in support of this hypothesis: (i) some strains harbored an incomplete *egc* and possibly correspond to evolutionary intermediates, and (ii) we identified two new enterotoxins that apparently arise from *egc* gene recombination events.

Analysis of the *egc* locus in a panel of 666 strains confirmed that *egc* is highly prevalent (63%) in *S. aureus* strains from nasal colonization, suppurative infections, and acute toxemia (2, 8, 12, 24, 29). As shown in other studies, the classical organization of the *egc* locus, i.e., the five enterotoxin genes

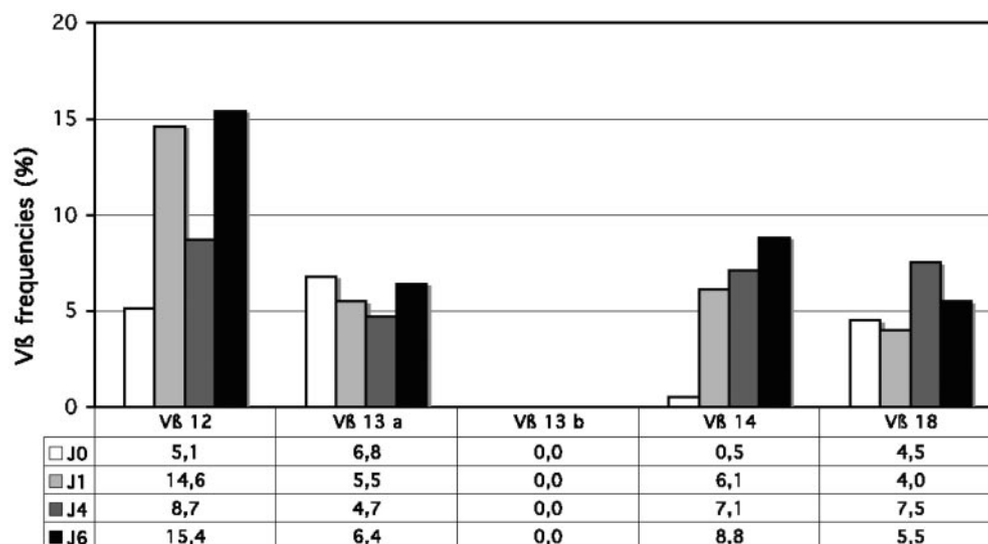


FIG. 6. Immunoscope analysis of the time course of T-cell stimulation by SE/U2. The quantitative V β repertoire determined with real-time PCR analysis on days 0, 1, 4, and 6 following stimulation with SE/U2 is shown. The x axis indicates V β families and the y axis their relative frequency of usage.

plus two pseudogenes, was most frequent (409 of 421 *egc*-positive strains) (Table 2), suggesting that this organization is currently the most successful. However, the *egc* locus of a few strains contained the *egc* promoter and the first gene (*selo*) but lacked one or more of the downstream genes of the operon, suggesting that these strains might correspond to evolutionary intermediates (Table 2). Unlike Omoe et al., we found no strains with a nonarchetypal *egc* operon that lacked the *selo* gene (29).

How might this tandem duplication and recombination occur? In five strains, two different insertion sequences were always inserted in the middle of an *egc* gene (Fig. 1). These ISs were typical, being composed of two ORFs encoding a transposase and a transposase helper protein, flanked by short terminal inverted repeat sequences. The transposases observed in our strains are related to Tnase 8-Rve or to IstB-Rve. Tnase 8 is a member of the IS3 family, which consists of various *E. coli* insertion elements and other bacterial transposases. Tnase 8 is required for efficient DNA transposition and could be involved in DNA replication, recombination, and repair. Proteins of the Rve family are described as integrase core domains, in that they have a catalytic integrase domain function. Insertion sequences have been found in the MRSA 252 genome (9) and in the staphylococcal cassette chromosome SCCmec type IVc of *S. aureus* strain MR 108 (11), in which their putative function is to facilitate mutations by transpositional mutagenesis. Thus, the five strains carrying an IS might represent evolutionary intermediates of the contemporary *egc* operon. In the same way, these insertion sequences might be involved in *egc* toxin gene export. Indeed, during transposition, they could integrate within an adjacent gene and transfer an *egc* toxin gene to a region outside the *egc* locus; this is the case for *seg2*, which is encoded by phage ϕ SA3 in strains MW2 and MSSA 476 (1).

A similar model of genetic variation involving tandem duplication and recombination, associated with the IS/transposase cassette, has been proposed for the DR13 locus. DR13

possesses various combinations of genes encoding staphylococcal exotoxin-like proteins (*set*). As in *egc*, all the *set* genes are similar to one another, suggesting that they were generated by successive duplication and variation of an ancestral gene. By sequence analysis of the DR13 region, Fitzgerald et al. showed that the DR13 locus could contain from 5 to 11 *set* genes (7). Interestingly, those authors identified a transposase gene downstream of the DR13 locus. Genetic analyses confirmed that all the *set* genes had a common ancestor and that multiple events of gene acquisition, duplication, recombination, and/or loss had led to the diversification of the DR13 locus.

A major result of our study is the identification, in one strain, of two new enterotoxins encoded by the *egc* locus and generated by (i) recombination between *selm* and *sei*, producing *selv*, and (ii) a limited deletion in the ϕ ent1- ϕ ent2 pseudogenes, producing *selu2*. We reconstructed the *selv* gene in silico by concatenation of the first 442 nucleotides of *selm* with nucleotides 451 to 730 of *sei*. Interestingly, we identified a 23-nucleotide sequence present in both the *selm* and *sei* genes, representing a potential template for recombination. This indicates that recombination between two toxin genes may lead to the emergence of new toxins in *S. aureus*. RT-PCR analysis showed that these two new toxin genes are transcribed at a level similar to that of other *egc* genes such as *sen* (Fig. 3). The biological activity of SE/V is consistent with that of a superantigen, specifically triggering T-cell subsets of the V β 6, V β 18, and V β 21 families. This pattern of activation is different from that reported for SE/M (6, 8, 9, 18 and 21.3) and SEI (1, 5.1, 5.3, and 23), in keeping with reports that slight amino acid sequence differences between members of the pyrogenic family can result in markedly altered biological properties (38). The *selu2* gene likely results from a simple deletion at the 3' end of the ϕ ent1 gene. The deletion abolishes the stop codon of ϕ ent1 and restores an open reading frame extending from the beginning of ϕ ent1 to the end of ϕ ent2. Letertre et al. described a similar mechanism generating the *selu* gene (15). However,

selu and *selu2* differ slightly in their nucleotide sequences, notably by the presence of an additional 15-nucleotide sequence at position 206 of the *selu* gene. As these 15 nucleotides are present in the *φent1* gene of the archetypal *egc* locus of strain A900322 (12) but are missing from the *φent1* gene of strain MU50 (14), *selu* would appear to derive from *φent1-φent2* in strain A900322, while *selu2* would derive from *φent1-φent2* in strain MU50. The superantigenic activity of SE/U2 was demonstrated by specific activation of Vβ 13.2 and Vβ 14 T cell subsets. In contrast to SE/V, for which both flow cytometric analysis and the TCR Vβ quantitative Immunoscope approach showed Vβ 6, 18, and 21 T-cell expansion (with the exception of Vβ 6, which is not covered by the IOTest Beta-Mark), no SE/U2-dependent triggering of Vβ 13 T cells was detected by the Immunoscope method. This could have been due to the broad specificity of the primer used to detect the Vβ 13a family (it amplifies Vβ 13.1, 13.2, 13.3, 13.4, 13.6, 13.7, 13.8, and 13.9), whereas the antibody used for flow cytometry detected only the Vβ 13.2 subset. Therefore, unless a Vβ 13.2-specific primer is used, the mild Vβ 13.2 T-cell expansion (fewer than 10% of CD3 T cells) cannot be detected by RT-PCR among total Vβ 13 cells. Selective toxin stimulation of a Vβ subset within a Vβ family has previously been reported with recombinant SE/P, which stimulates Vβ 5.1 and no other Vβ 5 subset (Vβ 5.2 or 5.3) (30), and with recombinant SE/R, which selectively stimulates Vβ 13.2 cells within the Vβ 13 family (31). In addition to the selective expansion of TCR Vβ subsets observed with the different toxins, Immunoscope analysis revealed that the CDR3 size distribution of TCR β-chain junctional transcripts within expanded Vβ subsets was Gaussian, reflecting a high level of polyclonality (Fig. 4).

The widespread distribution of the archetypal *egc* organization in strains from various genetic backgrounds (12, 33) and its high prevalence in *S. aureus* strains of various clinical origins suggest that this organization confers an optimal selective advantage. The single strain that was shown here to produce two new superantigens, SE/V (Vβ 6, 18, and 21.3) and SE/U2 (Vβ 13.2 and 14) but that lacked SE/M (Vβ 6, 8, 9, 18, and 21.3) and SEI (Vβ 1, 5.1, 5.2, and 5.3), stimulated different and, in theory, fewer Vβ subsets than archetypal *egc* strains; it is thus tempting to speculate that this strain might be less efficient at stimulating polyclonal T-cell proliferation. This *egc* variant might be counterselective, possibly explaining its rarity. Conversely, it is possible that ongoing variations within the *egc* locus, illustrated by strains carrying the insertion sequence, may produce new superantigen variants that could extend the stimulated Vβ subset and thereby supersede strains possessing the classical *egc* locus.

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AUTHOR'S CORRECTION

Staphylococcal Enterotoxin-Like Toxins U2 and V, Two New Staphylococcal Superantigens Arising from Recombination within the Enterotoxin Gene Cluster

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Volume 74, no. 8, p. 4724–4734, 2006. Page 4726: The paragraph below should have been included at the end of the Materials and Methods section.

Nucleotide sequence accession numbers. The nucleotide and amino acid sequences for *selu2* and *selv* have been deposited in GenBank under accession numbers EF030428 and EF030427, respectively.