Edinburgh Research Explorer

A Novel Core Genome-Encoded Superantigen Contributes to Lethality of Community-Associated MRSA Necrotizing Pneumonia

Citation for published version:

Wilson, GJ, Seo, KS, Cartwright, RA, Connelley, T, Chuang-Smith, ON, Merriman, JA, Guinane, CM, Park, JY, Bohach, GA, Schlievert, PM, Morrison, WI & Fitzgerald, JR 2011, 'A Novel Core Genome-Encoded Superantigen Contributes to Lethality of Community-Associated MRSA Necrotizing Pneumonia', *PLoS Pathogens*, vol. 7, no. 10, pp. e1002271. https://doi.org/10.1371/journal.ppat.1002271

Digital Object Identifier (DOI):

10.1371/journal.ppat.1002271

Link:

Link to publication record in Edinburgh Research Explorer

Document Version:

Publisher's PDF, also known as Version of record

Published In:

PLoS Pathogens

Publisher Rights Statement:

This is an open-access article distributed under the terms of the Creative Commons Attribution License, which

unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

General rights

Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy
The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact openaccess@ed.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.



A Novel Core Genome-Encoded Superantigen Contributes to Lethality of Community-Associated MRSA Necrotizing Pneumonia

Gillian J. Wilson¹, Keun Seok Seo², Robyn A. Cartwright¹, Timothy Connelley¹, Olivia N. Chuang-Smith³, Joseph A. Merriman³, Caitriona M. Guinane¹, Joo Youn Park², Gregory A. Bohach^{2,4}, Patrick M. Schlievert³, W. Ivan Morrison¹, J. Ross Fitzgerald¹*

1 The Roslin Institute and Centre for Infectious Diseases, University of Edinburgh, Easter Bush Campus, Midlothian, United Kingdom, 2 Department of Basic Sciences, Mississippi State University, Mississippi State, Mississippi, United States of America, 3 Department of Microbiology, University of Minnesota Medical School, Minnesota, United States of America, 4 Department of Biochemistry and Molecular Biology, Mississippi State University, Mississippi State, Mississippi, United States of America

Abstract

Bacterial superantigens (SAq) stimulate T-cell hyper-activation resulting in immune modulation and severe systemic illnesses such as Staphylococcus aureus toxic shock syndrome. However, all known S. aureus SAgs are encoded by mobile genetic elements and are made by only a proportion of strains. Here, we report the discovery of a novel SAg staphylococcal enterotoxin-like toxin X (SEIX) encoded in the core genome of 95% of phylogenetically diverse S. aureus strains from human and animal infections, including the epidemic community-associated methicillin-resistant S. aureus (CA-MRSA) USA300 clone. SEIX has a unique predicted structure characterized by a truncated SAq B-domain, but exhibits the characteristic biological activities of a SAg including $V\beta$ -specific T-cell mitogenicity, pyrogenicity and endotoxin enhancement. In addition, SEIX is expressed by clinical isolates in vitro, and during human, bovine, and ovine infections, consistent with a broad role in S. aureus infections of multiple host species. Phylogenetic analysis suggests that the selx gene was acquired horizontally by a progenitor of the S. aureus species, followed by allelic diversification by point mutation and assortative recombination resulting in at least 17 different alleles among the major pathogenic clones. Of note, SEIX variants made by human- or ruminant-specific S. aureus clones demonstrated overlapping but distinct Vβ activation profiles for human and bovine lymphocytes, indicating functional diversification of SEIX in different host species. Importantly, SEIX made by CA-MRSA USA300 contributed to lethality in a rabbit model of necrotizing pneumonia revealing a novel virulence determinant of CA-MRSA disease pathogenesis. Taken together, we report the discovery and characterization of a unique core genomeencoded superantigen, providing new insights into the evolution of pathogenic S. aureus and the molecular basis for severe infections caused by the CA-MRSA USA300 epidemic clone.

Citation: Wilson GJ, Seo KS, Cartwright RA, Connelley T, Chuang-Smith ON, et al. (2011) A Novel Core Genome-Encoded Superantigen Contributes to Lethality of Community-Associated MRSA Necrotizing Pneumonia. PLoS Pathog 7(10): e1002271. doi:10.1371/journal.ppat.1002271

Editor: Frank R. DeLeo, National Institute of Allergy and Infectious Diseases, National Institutes of Health, United States of America

Received December 6, 2010; Accepted August 1, 2011; Published October 13, 2011

Copyright: © 2011 Wilson et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Funding: This project was supported by grant BB/D521222/1 from the Biotechnology and Biological Sciences Research Council (www.bbsrc.ac.uk) (to J.R.F.) and Pfizer Animal Health (www.pfizerah.com) (to W.I.M. and J.R.F.). P.M.S. was supported by National Institutes of Health Grants R01 Al074283 and U54 Al57153. P.M.S. and O.N.C.-S. are members of the Great Lakes Regional Center of Excellence in Biodefense and Emerging Infectious Diseases. G.A.B. was supported by P20-RR16549 and P20-RR15587 (GAB), and by United States Department of Agriculture NIFA AFRI grant 2008-35204-04582 (KSS and GAB). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

* E-mail: Ross.Fitzgerald@ed.ac.uk

Introduction

Staphylococcus aureus is responsible for an array of diseases including life-threatening toxinoses such as toxic shock syndrome (TSS) and necrotizing pneumonia. Many strains of *S. aureus* have been shown to produce members of a family of more than 20 serologically distinct superantigenic (SAg) toxins, which contribute to disease pathogenesis via modulation of the host immune response [1,2]. Previously characterized SAgs are small secreted proteins of 20 to 28 kDa in size, which share similar biochemical, structural, and immunobiological properties [1,2], but can be differentiated into 4 distinct subgroups according to their phylogenetic relatedness [3–5]. They share a compact 2-domain protein structure consisting of domain A which contains a long

central α -helix and a C-terminal β -grasp motif, and the smaller domain B which contains an N-terminal oligonucleotide-oligosaccharide binding (OB) fold [1,2,5,6]. SAgs bypass the conventional antigen processing pathway, by binding simultaneously to MHC class II molecules of antigen-presenting cells, outside of the antigen-binding groove, and the variable region of the T-cell receptor (TCR) β -chain (V β) [7,8]. In so doing, SAgs can react with all T-cells expressing reactive V β -TCR regions (up to 50%), instead of the normal responding frequency to an antigen of approximately 0.01%, resulting in massive inflammatory cytokine release and consequent toxic shock [9]. SAgs also have the capacity to cause immune suppression by inducing T-cell anergy, and may contribute to bacterial persistence during chronic infection [10].

Author Summary

Staphylococcus aureus is a global pathogen, responsible for an array of different illnesses in humans and animals. In particular, community-associated methicillin-resistant S. aureus (CA-MRSA) strains of the pandemic USA300 clone have the capacity to cause lethal human necrotizing pneumonia, but the molecular basis for the enhanced virulence remains unclear. Bacterial superantigens (SAg) stimulate T-cell hyper-activation resulting in severe systemic illnesses such as toxic shock syndrome (TSS). However, all S. aureus SAgs identified to date are encoded by mobile genetic elements found only in a proportion of clinical isolates. Here, we report the discovery of a unique core genome-encoded SAg (SEIX) which was acquired by an ancestor of the S. aureus species and which has undergone genetic and functional diversification in pathogenic clones infecting humans and animals. Importantly, we report that SEIX made by pandemic USA300 contributes to lethality in a rabbit model of human necrotizing pneumonia revealing a novel virulence determinant of severe CA-MRSA infection.

All staphylococcal SAgs identified to date are encoded by mobile genetic elements (MGE) such as plasmids, phages, transposons and S. aureus pathogenicity islands (SaPIs), or the highly variable genomic region $vSa\beta$ [4,11–14]. Accordingly, the distribution of SAg genes is S. aureus strain-dependent. For example, Omoe et al. reported that 80% of human isolates contain at least one SAg gene, including 50% which contain the enterotoxin gene cluster (egc) locus, and Smyth et al. demonstrated that 57% of animal strains examined contained at least one SAg gene with the egc found in 30% of isolates [15,16]. These data collectively suggest that no single SAg is encoded by more than 50% of strains and that some strains may not have superantigenic capacity at all. The USA300 S. aureus clone which is responsible for the current CA-MRSA epidemic affecting the USA and other countries is typically associated with skin and soft tissue infections and has the capacity to cause lethal toxinoses such as necrotizing pneumonia and extreme pyrexia [17-19]. However, USA300 strains do not typically produce TSST-1 or staphylococcal enterotoxins B and C, the SAgs most often associated with toxic shock [20,21]. Here, we report the discovery of a unique core genome-encoded SAg (designated SEIX), encoded by the great majority of S. aureus strains, determine its immunobiological function and examine its molecular evolution. Our data suggest that SEIX contributes to immune modulation in both human and animal disease pathogenesis, and demonstrate a role for SEIX in the development CA-MRSA USA300 necrotizing pneumonia in a rabbit model of infection.

Results

Identification of a novel putative SAg gene encoded by the great majority of S. aureus strains

In order to examine the superantigenic capacity of the epidemic CA-MRSA clone, USA300, we carried out a genome-wide screen of USA300 strain FPR3757 to identify genes encoding proteins with homology to known SAgs [21]. In addition to characterized SAgs, SEIK and SEIQ, we discovered a SAg gene homolog, SAUSA300_0370 which encodes a protein with 24% amino acid identity to the major virulence factor, TSST-1 and 27% identity with the SAg-like protein, SSL7. The gene, designated selx encoding staphylococcal enterotoxin-like toxin X (SEIX), is 612 bp in length and is most closely-related to the phylogenetic group IV of staphylococcal SAgs represented by TSST-1, closely allied to the staphylococcal SAg-like (SSL) protein family (Fig. 1a). The mature form of SEIX is 168 amino acids in length with a predicted molecular weight of 19343 Da, smaller than any known staphylococcal SAg [1,4]. In order to examine the distribution of selx among S. aureus strains, we screened a total of 114 isolates by a combination of bioinformatic interrogation of 53 publicly available S. aureus genome sequences (Table S1), and PCR analysis using selx-specific oligonucleotide primers (Table S2) of a further 61 S. aureus isolates selected to represent the full breadth of species diversity and to include the most important human and animal pathogenic clones (Table S3). Remarkably, 108 of the 114 isolates (95%) contained the selx gene which included all strains examined except isolates of the CC30 clonal complex (Fig. 1b; Table S1 and

Analysis of the location of selx in the genomes of strains representing diverse multi-locus sequence types (ST)s, including MSSA476 (ST1), N315 (ST5), USA300 TCH3757 (ST8), H19 (ST10), A9635 (ST45slv), TCH130 (ST72), JKD6159 (ST93), JH1 (ST105), ED133 (ST133), RF122 (ST151), 04-02891 (ST225), JKD6008 (ST239), COL (ST250) and TCH959 (ST1159), revealed that selx is located at an identical chromosomal site in all strains examined (Fig. S1). Specifically, it is located ~400 kb from the origin of replication in the oriC environ among a cluster of 4 genes specific for the S. aureus species including 2 genes encoding hypothetical proteins of unknown function, and a predicted integrase pseudogene which contains partial homology to an integrase encoded by the S. aureus bacteriophage, PT1028 [22]. The gene cluster is flanked by conserved genes encoding ribosomal proteins and a DNA-binding protein involved in DNA replication, which are located at the same genomic location in other staphylococcal species such as Staphylococcus epidermidis (Fig. S1). The genetic linkage of *selx* with an integrase pseudogene, its wide distribution across the full breadth of S. aureus species diversity, and its absence in the genomes of other staphylococcal species indicates an ancient horizontal acquisition which may have occurred during S. aureus speciation. The existence of a single S. aureus clonal complex (CC30), which does not contain selx but contains the adjacent integrase pseudogene at the same chromosomal site, is consistent with a deletion event in a progenitor of the clonal lineage which contained the selx gene (Fig. S1).

selx exhibits considerable species-wide allelic variation and evidence for assortative recombination

In order to examine the allelic variation of selx among S. aureus strains representing the breadth of species diversity, DNA sequencing of selx was carried out for 11 S. aureus strains representative of ST 9, 12, 15, 25, 45, 97, 121, and 126 and compared to the sequence of selx for 49 strains available in Genbank (Table S1). selx was represented by 17 distinct allelic variants encoded by human and animal strains, designated selx1-14, selxbov1-2, and selxov, respectively. Novel nucleotide sequences for selx8-11 and selxbov2 have been deposited in GenBank under accession numbers HQ850970, HQ850971, HQ850968, HQ850969 and HQ85097, respectively. Of note, the human ST398 strain OS385 contains selx13, which is a 315 bp truncated C-terminal fragment of the gene predicted to be non-functional. With the exception of the truncated selx13 there are 85.9% nonvariable nucleotide sites among the selx alleles identified. This corresponds to a total of 40 (13%) variable amino acid residues among the 204 amino acid sites of SEIX (Fig. S2). Pairwise amino acid alignment reveals 89.5 to 98.5% amino acid identity among

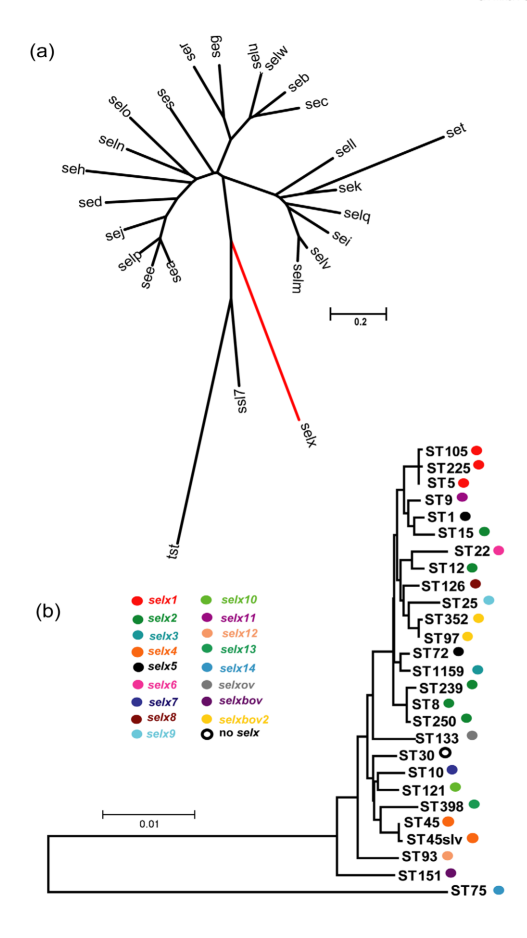
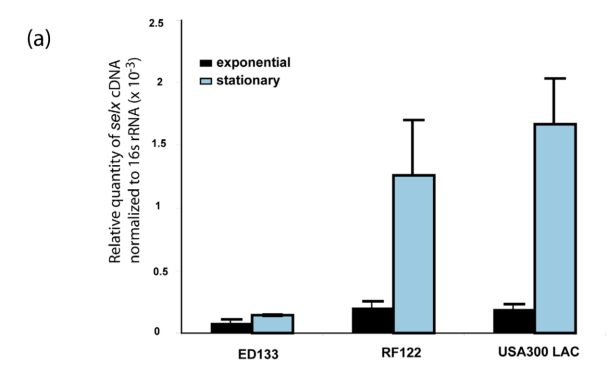


Figure 1. Phylogenetic analysis of *selx* **and its species-wide distribution. a**) Neighbor-joining tree based on the sequences of all known staphylococcal SAgs and the SAg-like protein, SSL7 **b**) Neighbor-joining tree of concatenated MLST sequences of representative *S. aureus* STs representing the breadth of species diversity. Coloured circles denote the presence of specific *selx* alleles. doi:10.1371/journal.ppat.1002271.g001

the different allelic variants made by the major *S. aureus* pathogenic clones (Fig. S2).

In general, each clonal complex (consisting of closely-related STs) contains a unique *selx* allele (Fig. 1b) consistent with point mutation being the major driving force for evolution within

individual lineages. However, allele *selx2* was identified in ST12, ST15 and CC8 strains, and *selx5* was encoded by both ST1 and ST72 strains, respectively, indicating that assortative recombination of the *selx* gene has contributed to its distribution among some lineages which have not shared a recent common ancestor



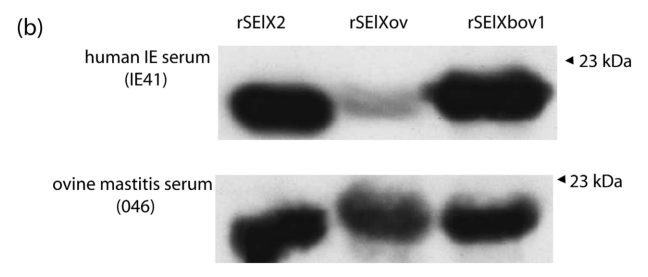


Figure 2. SEIX is expressed by clinical isolates *in vitro* **and during infection. a**) Growth-phase dependent transcription of *selx* determined by qRT-PCR. **b**). Representative Western blot analysis of rSEIX proteins with human infective endocarditis (IE) or ovine mastitis sera samples. doi:10.1371/journal.ppat.1002271.g002

Table 1. Immunoreactivity of recombinant SAg variants with sera from healthy humans, and bovine, ovine and human patients recovering from S. aureus infections.

Serum sample	SEIXbov1 ^a	SEIXov ^a	SEIX2 ^a	SEIO ^a
Human (Normal)				
HV093	n/d	n/d	+	-
HVRC	n/d	n/d	-	+
HVJRF	n/d	n/d	-	-
HV100	n/d	n/d	-	-
HV091	n/d	n/d	+	-
HV126	n/d	n/d	+	-
HV097	n/d	n/d	+	-
HV101	n/d	n/d	+	-
HV118	n/d	n/d	+	-
HV115	n/d	n/d	+	-
HV017	n/d	n/d	+	-
HV105	n/d	n/d	+	
HV003	n/d	n/d	+	-
HV008	n/d	n/d	+	-
HV148	n/d	n/d	+	-
HV139	n/d	n/d	+	-
HV014	n/d	n/d	+	-
HV047	n/d	n/d	+	-
HV020	n/d	n/d	+	-
HV057	n/d	n/d	+	-
HV104	n/d	n/d	+	-
HV002	n/d	n/d	+	-
HV098	n/d	n/d	+	-
Human IE				
E19	+	+	+	-
E37	+	+	+	-
E41	+	+	+	-
Serum sample	SEIXbov1 ^a	SEIXov ^a	SEIX2 ^a	SEIO ^a
E51	+	+	+	-
E54	+	+	+	-
Bovine mastitis				
2480	+	+	+	-
2487	+	+	+	-
2521	+	+	+	-
4227	+	+	+	-
2211	-	-	-	-
Ovine mastitis				
011	+	+	+	-
040	+	+	+	-
046	+	+	+	-
0112	+	+	+	-

^a+, positive reactivity; -, negative reactivity; n/d, not done. doi:10.1371/journal.ppat.1002271.t001

(Fig. 1b). In addition, examination of the contribution of recombination to selx diversity employing the RDP3 suite of programs [23], revealed evidence for at least 3 distinct

recombination events leading to hybrid alleles of selx (Fig. S3). Furthermore, a phylogenetic tree based on selx gene sequences had a topology distinct from the phylogenetic tree derived from

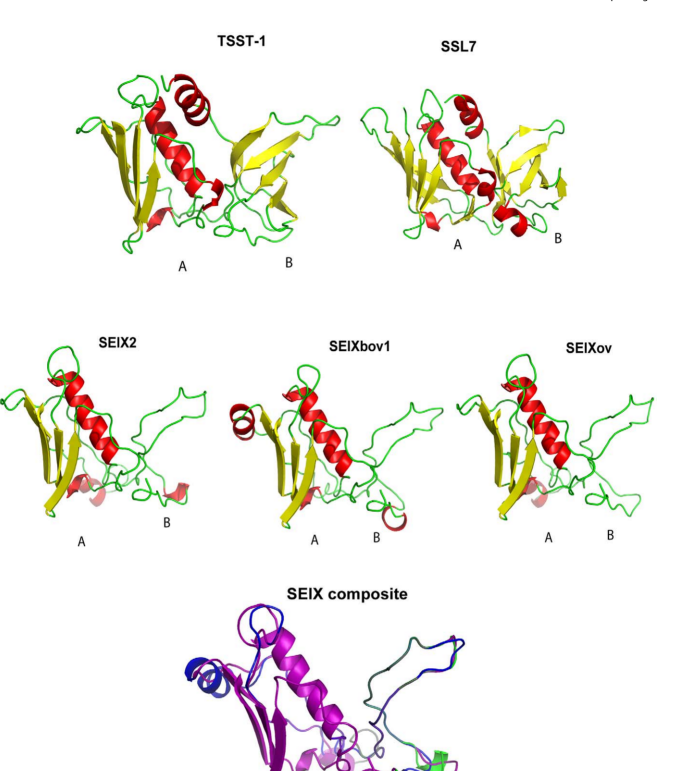


Figure 3. SEIX is predicted to have a truncated B-domain. Schematic diagram of the solved structures of TSST-1, SSL7 and modelled 2-domain structures of human, bovine and ovine variants of SEIX. SEIX composite structure was obtained by superimposition of protein variants in PyMol. doi:10.1371/journal.ppat.1002271.g003

Α

concatenated MLST loci (Fig. S4). Taken together, these data suggest that recombination has contributed to the diversification and distribution of selx among S. aureus clones.

SEIX is expressed by clinical isolates in vitro and during human, bovine and ovine infection

In order to determine whether selx was expressed by clinical isolates we examined by quantitative real-time PCR (qRT-PCR), the transcription of selx by human S. aureus CA-MRSA USA300 strain LAC, bovine strain RF122, and ovine strain ED133 during exponential and stationary phases of growth in vitro. selx was transcribed by each strain in a growth-phase dependent manner, and the highest relative expression level was demonstrated by USA300 strain LAC (Fig. 2). To investigate the in vitro expression of SEIX we carried out Western blot analysis with stationary phase culture supernates of 15 representative clinical isolates of S. aureus of human, bovine, and ovine origin, with rat anti-serum specific for SEIX. Expression of SEIX was detected in 13 of 15 isolates including 5 of 5 human isolates from sepsis, scalded skin syndrome and infective endocarditis patients, 4 of 4 bovine mastitis isolates, 3 of 4 ovine mastitis isolates, and a single caprine mastitis isolate. A single avian isolate ED98 did not express SEIX in detectable quantities. These data indicate that SEIX is made by the majority of clinical isolates at detectable levels in nutrient replete conditions in vitro (Table S3).

To determine whether SEIX is expressed during human and animal colonization or infection, Western immunoblot analysis was carried out using recombinant SEIX human (SEIX2), bovine (SEIXbov1) and ovine (SEIXov) variants, with sera from humans, cows and sheep recovering from S. aureus infections, and from healthy human volunteers. Specific SEIX antisera did not crossreact with TSST-1 or SSL7, its closest homologs (Fig. S5). In addition, we identified individuals who were seroconverted for SEIX but not for SSL7 or TSST-1, and an individual who was seroconverted for SSL7 and TSST-1 but not for SEIX (Fig. S5). Taken together, these data indicate a lack of cross-reactivity of antibodies specific for TSST-1, SSL7 and SEIX. All 5 human, 4 bovine and 5 ovine serum samples from infected individuals, and 20 of 23 serum samples from healthy humans contained antibody that recognized the SEIX recombinant proteins, whereas only one serum sample (from a healthy human) out of the total of 47 contained antibody that recognized recombinant SEIO SAg (Table 1). These data indicate that SEIX is made during S. aureus colonization or infection of humans and animals and stimulates a humoral immune response. Of note, densitometric analysis of SEIX immuno-reactive bands indicated approximately 2-fold lower reactivity of human IgG with recombinant SElXov, in comparison to IgG from S. aureus-infected sheep, suggesting that SEIXov made by ovine S. aureus may vary antigenically (Fig. 2 b). Overall, the expression of SEIX during infection of humans and animals suggests an important general role during S. aureus infection of multiple host species.

SEIX has a unique predicted SAg structure

The SEIX2 variant encoded by S. aureus USA300 strain LAC, contains 27% and 24% amino acid identity with its nearest protein homologs, superantigen-like protein 7 (SSL7) and TSST-1, respectively. In order to investigate the predicted structure of the novel SAg, we carried out structural modeling using the program Phyre [24] with SSL7 and TSST-1 crystal structures from the PDB database as templates. Considering, the low sequence homology of SEIX with its closest homologs SSL7 and TSST-1, used as templates for modeling, the results should be considered to be speculative. The hypothetical predicted structures of SEIX

variants imply a potential to form a characteristic 2 domain SAg structure with some unusual features (Fig. 3). The predicted Adomain containing the C-terminus has a long central α-helix, positioned along one side of a 4-stranded β-sheet, forming a structure typical of the β-grasp motif [25]. However the B-domain which contains the N-terminus, is considerably smaller than the Bdomain of previously characterized SAgs and lacks an OB fold (Fig. 3). The OB fold is the site typically involved in SAg binding to the α-chain of the MHC class II complex on antigen presenting cells [2,26]. The functional implications of this unique predicted B-domain structure are currently unknown.

SEIX is mitogenic and demonstrates Vβ-dependent T-cell proliferation

We examined the function of representative allelic variants of SEIX from human, bovine and ovine isolates. Recombinant human (rSElX2), bovine (rSElXbov1), and ovine (rSElXov) variants were purified and used to stimulate human PBMC in a thymidine incorporation assay. All 3 rSEIX variants were mitogenic for human T-cells at concentrations above 0.1 ng/μl (Fig. 4a). Further, in order to examine the mitogenicity of SEIX expressed in a native S. aureus background, selxbov1 was cloned into the pALC2073 vector and expressed in a SAg-deficient strain RF122-8. The supernatant from cultures of RF122-8 + pALC2073:SEIXbov1 had mitogenic activity for bovine lymphocytes, but supernatant from strain RF122-8 containing the pALC2073 plasmid only had none (data not shown). SEIX was also mitogenic for rabbit lymphocytes and importantly, the superantigenic activity of SEIX for both human and rabbit Tcells is comparable or higher to the activity of TSST-1, the SAg responsible for the toxinoses toxic shock syndrome (TSS) (Fig. 4b,

We recently designed a novel assay for quantifying human $V\beta$ $(humV\beta)$ gene expression including all of the known $V\beta$ subgroups [27]. Previous studies of the bovine $V\beta$ (bov $V\beta$)-dependent expansion capacity of staphylococcal SAgs have been restricted by the number of identified bovine $V\beta$ subfamilies [11,28]. The recent bovine genome sequencing project has facilitated a comprehensive description of the bovine $V\beta$ subgroups and the repertoire of functional Vβ genes [29,30]. In the current study, design of a novel bovV_β-dependent expansion assay has allowed us for the first time to evaluate the response of 23 human and 18 bovine Vβ subfamilies to stimulation with a staphylococcal SAg (Fig. 5a, b). rSEIX2 and rSEIXbov1 activated humVβ subfamiles 1, 6, 18 and 21, whereas rSElXov activated 1, 6 and 18 but not 21 (Table 2). All SEIX variants stimulated bovVβ subfamily X. In addition SElXov activated bovVβ 16, SElXbov1 activated bovVβ 3, 5, 11, 16 and 17, and SEIX2 activated bovVβ 5, 6, 17 and 24. Different human and bovine VB subfamilies are activated in response to stimulation with SEIX. This is in part due to the activation of humV \beta subfamilies 18 and 21, for which there are no bovine orthologs, and bovVB X which has no human ortholog. However, the orthologous humV β 6 and bovV β 6 are both activated by SEIX2. In addition, bovVβ subfamily 16 which is activated by SElXbov1 and SElXov is phylogenetically related to $humV\beta$ 6. Of note, the $humV\beta$ 1 subfamily and the closely related bovV β 5 but not the orthologous bovV β 1 were activated by SEIX variants. These data indicate a unique pattern of $V\beta$ gene activation for SEIX in comparison to other previously characterized SAgs [3,27]. Importantly, the results demonstrate differences in superantigenic activity and distinct VB activation profiles for different SEIX variants made by S. aureus strains associated with different host species.

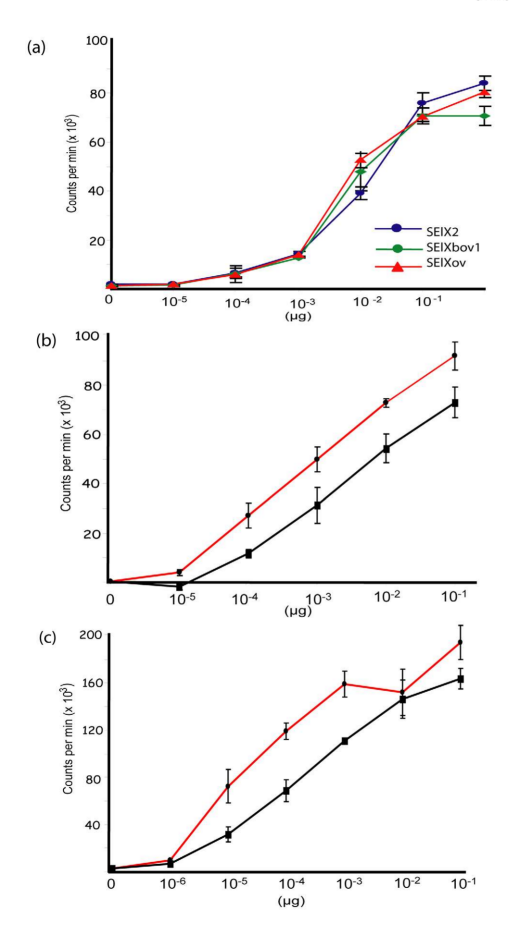


Figure 4. SEIX stimulates proliferation of human and rabbit lymphocytes. a) Proliferation of human PBMC cultures with recombinant SEIX2, SEIXbov1, and SEIXov variants measured by [3H]-thymidine incorporation. Proliferation of human (b) or rabbit (c) PBMC cultures with recombinant SEIX2 (red line) or TSST-1 (black line) measured by [3H]-thymidine incorporation. doi:10.1371/journal.ppat.1002271.g004

SEIX has functional activities characteristic of SAgs

We examined the ability of rSEIX2 to cause TSS in rabbits using a standard mini-osmotic pump model of TSS at a dose of 200 µg/kg. Of the 5 rabbits given rSEIX2, 4 succumbed within 7 d, compared to 0 of 5 treated with PBS (p<0.05) (Fig. 6a). The positive control TSST-1 was lethal for 3 of 3 animals in the 7 d test period (Fig. 6a). Changes in temperature of the rabbits were recorded before and 24 h post-SAg challenge. Rabbits treated with rSEIX2 developed fever with a 2.0°C increase in body temperature over the 24 h test period, compared to 0.4°C for PBS treated animals (p<0.001) (Fig. 6a).

Rabbits were also used to evaluate fever responses over 4 h after intravenous injection, followed by the ability to enhance lethality of LPS over a 48 h test period. The capacity to cause fever that peaks 4 h after injection, and to amplify the lethal effects of LPS by up to 10⁶-fold are defining activities of SAgs. rSEIX2 caused dosedependent fever responses in rabbits that peaked 4 h after injection and the 4 h fever response of animals receiving 10 µg/ kg of rSEIX2, compared to PBS, was significantly different (p value of 0.001) (Fig. 6b). The minimum pyrogenic dose of SAg is defined as the dose per kg required to cause an average 0.5°C rise in rabbit body temperature in 3 animals; the minimum pyrogenic dose of rSEIX2 was calculated to be 1.0 µg/kg. rSEIX2 also enhanced rabbit susceptibility to lethal shock by LPS (Fig. 6b). Taken together, rSEIX2 demonstrated the functional characteristics required for classification as a SAg, including mitogenicity, pyrogenicity, enhancement of endotoxin shock, and lethality when administered by mini-osmotic pump.

SEIX contributes to lethality of CA MRSA USA300 in a rabbit model of necrotizing pneumonia

CA-MRSA is notorious for causing skin and soft tissue infections, and severe necrotizing pneumonia [20,31]. We found that CA-MRSA USA300 strains expressed elevated levels of selx in vitro relative to other clinical isolates (Fig. 2a). In order to investigate the hypothesis that SEIX contributes to the pathogenesis of necrotizing pneumonia, we constructed an SEIX-deficient mutant of CA-MRSA USA300 strain LAC (LAC Δselx), and a repaired derivative with an intact functional selx gene (LAC $\Delta selx$ rep). To rule out the possibility of spurious mutations acquired during the construction of LAC $\Delta selx$ in loci which could influence virulence, such as the accessory gene regulator (agr) [32], we compared the isogenic LAC wild type, LAC $\Delta selx$ and LAC $\Delta selx$ rep strains for hemolytic titre, secreted and cell wall-associated protein profiles, and in vitro growth rate. In each case we found that the strains were indistinguishable (Fig. S6 and data not shown). In addition, we compared LAC wild type and LAC $\Delta selx$ for α -toxin and PVL expression levels by ELISA and found identical levels of expression (data not shown). We then compared the ability of wild type, mutant and repaired strains to cause lethal necrotizing pneumonia in a rabbit model [33]. For animals receiving wild type LAC, all 11 rabbits succumbed within 4 d, compared to 2 of 11 receiving LAC Δselx (p<0.002) (Fig. 6c). Importantly, 4 of 4 rabbits infected with the repaired strain LAC $\Delta selx$ rep succumbed within 4 d (Fig. 6c). Body temperatures were recorded in the first 24 h of the experiment, and animals receiving strain LAC demonstrated significantly higher body temperatures than animals receiving the SEIX-deficient strain LAC $\Delta selx$ (p<0.002) and LAC $\Delta selx$ rep had wild type levels of pyrogenicity (Fig. 6d). On gross examination, regionally extensive to lobar areas of lung from rabbits infected with wild type USA300 LAC and the repaired strain USA300 LAC Δselx rep were dark red to purple, heavy and oozed blood on the cut surface (Fig. 6e). In contrast, lungs from rabbits infected with the LAC $\Delta selx$ mutant strain were reddened with well demarcated, focal white areas (abscess), but did not show signs of haemorrhage (Fig. 6e). Histological examination of tissues from both the wild type USA300 LAC and the repaired strain USA300 LAC Δselx rep revealed severe haemorrhagic and necrotising pneumonia (Fig. 6f). Tissue sections from rabbits infected with the LAC $\Delta selx$ mutant strain were only moderately affected (Fig. 6f). Taken together, these data collectively indicate that SEIX made by USA300 LAC causes high fever and contributes to severity of infection and lethality in a rabbit model of necrotizing pneumonia.

Discussion

We have identified a novel SAg which is encoded by the great majority of strains, and which causes lethality in a model of severe S. aureus human disease. The distribution of selx across the full breadth of S. aureus diversity, its absence among other staphylococcal species, and its genetic linkage with an integrase pseudogene suggests an ancient horizontal acquisition event which happened prior to the most recent common ancestor of the S. aureus species. The gene is absent from only a single clone examined (CC30) but the associated integrase pseudogene is retained suggesting that a deletion event has resulted in the loss of selx during the evolution of the CC30 lineage. Of note, previous studies have discovered that the majority of CC30 isolates contain a SaPI which encodes TSST-1, the most closely-related SAg to SEIX [34,35]. These data imply that virtually all S. aureus isolates have the capacity to produce either TSST-1 or SEIX suggesting an important role for this sub-group of toxins in S. aureus pathogenesis.

The selx gene has undergone diversification leading to at least 17 different allelic variants identified among the major clonal lineages (Fig. 1 b). Although SEIX is the first SAg identified which is encoded in the core genome of S. aureus, the SAgs, SMEZ and SPEG, are made by the majority of Group A Streptococcal (GAS) isolates [36]. Extensive allelic variation of SMEZ contributes to antigenic variation, but does not influence Vβ-specificity or mitogenicity [36]. Whereas GAS is specific for human hosts, S. aureus is also represented by strains which are specialized for ruminant or avian host species [37,38]. In contrast to SMEZ, SEIX bovine- and ovine-specific variants displayed distinct Vβ subgroup proliferation profiles for bovine lymphocytes in comparison to a human-specific SEIX variant suggesting that they have undergone adaptive diversification leading to enhanced activity in ruminants. Previously, it was shown that the proliferation of human and bovine lymphocytes varied in response to stimulation with allelic variants of the SAg staphylococcal enterotoxin C (SEC). The SEC bovine variant (SECbov) requires between 10- and 1000-fold more toxin than SEC1 and the SEC ovine variant (SECov) to induce proliferation of human or bovine PBMCs, and results in a distinct Vβ-dependent T-cell activation profile [28,39].

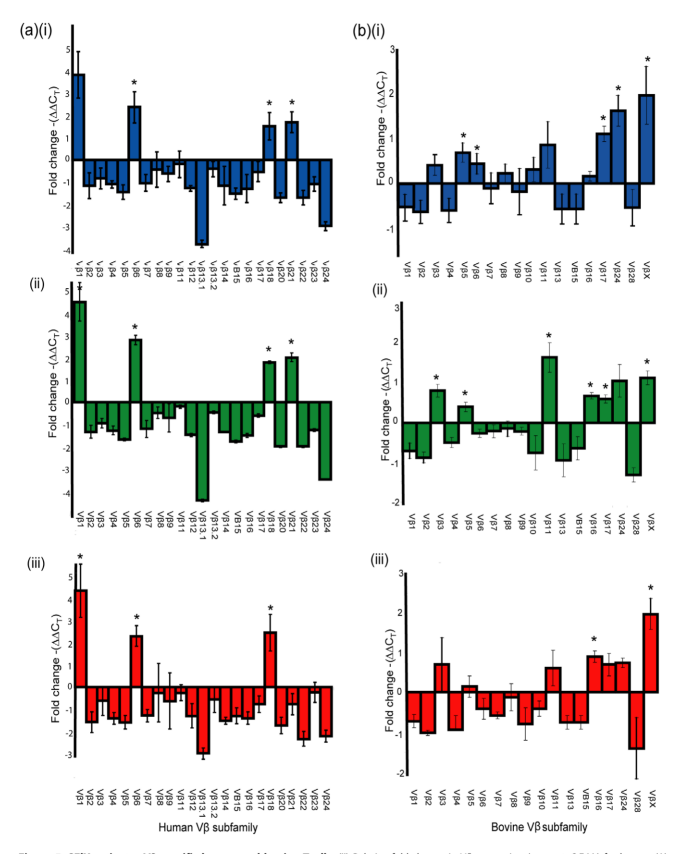


Figure 5. SEIX activates Vβ-specific human and bovine Tcells. (ii) Relative fold-change in Vβ expression (mean \pm S.E.M.) for human (A) or bovine (B) T cells after stimulation with (i) SEIX2, (ii) SEIXbov1, and (iii) SEIXov. *indicates statistical significance (p<0.05). doi:10.1371/journal.ppat.1002271.g005

Table 2. Human and bovine $V\beta$ subfamilies activated in response to rSEIX variants.

SEIX variant	Human Vβ ^{a, b}	Bovine Vβ ^{a, b}
SEIX2	1, 6, 18, 21	3, 5, 6 , 11, 17 , 24 , X
SEIXbov1	1, 6, 18, 21	3, 5, 11, 16, 17, 24, X
SEIXov	1, 6, 18	16 , 17, 24, X

 a Vβ subfamily nomenclature followed the classification of Arden *et al* [65]. b Bold type indicates subfamilies activated with a significance of p<0.05, and normal type indicates subfamilies with a trend (p<0.1) towards significant activation above baseline control (unstimulated). doi:10.1371/journal.ppat.1002271.t002

Although most of the allelic variation in *selx* was clonal and due to point mutation, we found evidence of a role for recombination in the diversification and distribution of SEIX, including whole gene transfer of the *selx* allele between pathogenic clones. Recombination rates are predicted to be low in *S. aureus* in general [40], but some recombination, particularly among genes involved in virulence has been observed [41,42]. Such rapid evolution may facilitate antigenic or functional diversification of proteins which are critical for bacterial survival during infection.

Analysis of *selx* transcription levels revealed a growth phase-dependent expression analogous to numerous other staphylococcal virulence proteins which are under the control of *agr*. Of note, the human CA-MRSA USA300 strain LAC had relatively high levels of expression of SEIX which correlates with the previously reported up-regulation of RNAIII and secreted virulence proteins by the USA300 epidemic clone [43]. In addition to demonstrating the *in vitro* expression of SEIX by *S. aureus* strains, we also identified SEIX-specific antibodies in human and ruminant convalescent serum samples, and in healthy individuals. The high frequency of sero-conversion to SEIX among human and livestock populations is consistent with the expression of SEIX by most commensal and disease-associated isolates of *S. aureus*, and imply a broad role for SEIX in *S. aureus* colonization or infection of multiple host species.

Of particular importance, the USA300 epidemic clone is currently the primary infectious cause of human mortality in the USA, and unlike most strains of hospital-associated MRSA can cause disease in otherwise healthy individuals [44]. The molecular basis for the increased virulence of USA300 strains has been the subject of intensive research efforts. Several reports have highlighted the high levels of expression of secreted proteins such as Panton-Valentine leukocidin, α-toxin, and phenol-soluble modulins and evidence exists for a role for each of these toxins in the pathogenesis of CA-MRSA severe infection [43,45-50]. Recently, Assimacopoulos et al. described an S. aureus extreme pyrexia syndrome that is associated with CA-MRSA USA300 strains [51]. Our studies suggest that SEIX contributes to the pathogenesis of pulmonary illness caused by these strains, and its induction of high fever during infection is consistent with a role in extreme pyrexia syndrome. While most studies on the pathogenesis of severe S. aureus pulmonary infections have focused on the role of cytolysins [43,45,49,50], our data suggest that the novel SAg SEIX may influence the outcome of severe infection caused by CA-MRSA USA300. This is in agreement with a recent study that shows that TSST-1, SEB and SEC contribute to lethal pneumonia of rabbits caused by CA-MRSA USA200 and USA400 strains [33]. Of note, rabbits are much more similar to humans than mice in their sensitivity to both SAgs and cytolytic toxins [52]. Importantly, we have shown that SEIX is mitogenic for both rabbit and human T cells. As the majority of previous

studies into the virulence of *S. aureus* infections have been carried out in mice, the importance of SAgs in the pathogenesis of severe infections may have been underestimated.

Because of the role of SAgs in *S. aureus* pathogenesis and their potential as an agent of biological warfare, therapeutic strategies which target these toxins have been the focus of considerable research efforts [6]. Structural modeling revealed the potential of SEIX to form the characteristic 2 domain SAg structure joined by a central α -helix. However domain B of SEIX is predicted to be much smaller than that of other SAgs identified to date (Fig. 4). Ongoing crystallography analyses should result in important insights into the molecular interaction of the uniquely structured SEIX with its ligands which may facilitate the design of molecules to inhibit its function.

CA-MRSA USA300 strains are characterized by high expression of a number of secreted virulence factors which through direct activity or immunopathology result in the severe symptoms associated with infections such as necrotizing pneumonia [43,45–50]. Here, we report the discovery of a novel secreted virulence determinant made by most strains of *S. aureus* that may contribute to the severity of some human diseases caused by highly-virulent clones such as CA-MRSA USA300. Furthermore, the phylogenetic and immunobiological characterization of a unique *S. aureus* core genome-encoded SAg provides new insights into the evolution of pathogenic *S. aureus* and its capacity to cause disease in multiple host species.

Materials and Methods

Ethics statement

All animals were handled in strict accordance with good animal practice as defined by the relevant national and/or local animal welfare bodies. Animal experimentation was performed under a University of Minnesota approved Institutional Animal Care and Use Committee (IACUC) protocol (0908A71722). University of Minnesota is accredited by the Association for Assessment and Accreditation of Laboratory Animal Care International (AALAC). Animals are maintained in accordance with the applicable portions of the Animal Welfare Act and the DHHS "Guide for the Care and Use of Laboratory Animals". In agreement with the University of Minnesota IACUC, animals that failed to exhibit escape behaviour and at the same time could not right themselves were prematurely euthanized. Animals were euthanized with intravenous injection of 1 ml of Beuthanasia D, whether prematurely or at the end of experimentation. Human peripheral blood mononuclear cells were isolated from heparinized venous blood of four different healthy donors in accordance with a human subject protocol approved by the University of Idaho Institutional Review Board for Human Subjects (approval number 05–056), in accordance with the principles of the Declaration of Helsinki. Donors were informed the procedure risks and provided a written consent prior to participation. The University of Idaho is fully accredited by the AALAC.

S. aureus strains

S. aureus strains were selected to represent clonal genotypes which represent the breadth of species diversity, different host-associations and geographic origins (Table S3). S. aureus strains were grown in tryptone soya broth (TSB) or brain heart infusion (BHI) broth with shaking at 200 rpm, or on tryptone soya agar (TSA) plates. For use in animal studies, organisms were cultured in Todd Hewitt broth or on Todd Hewitt agar. Media was supplemented where appropriate with chloramphenicol (Sigma-Aldrich) at 12.5 μ g/ml and 150 μ g/ml X-gal (5-bromo-4-chloro-3-indolyl-beta-D-galactopyranoside) (Melford).

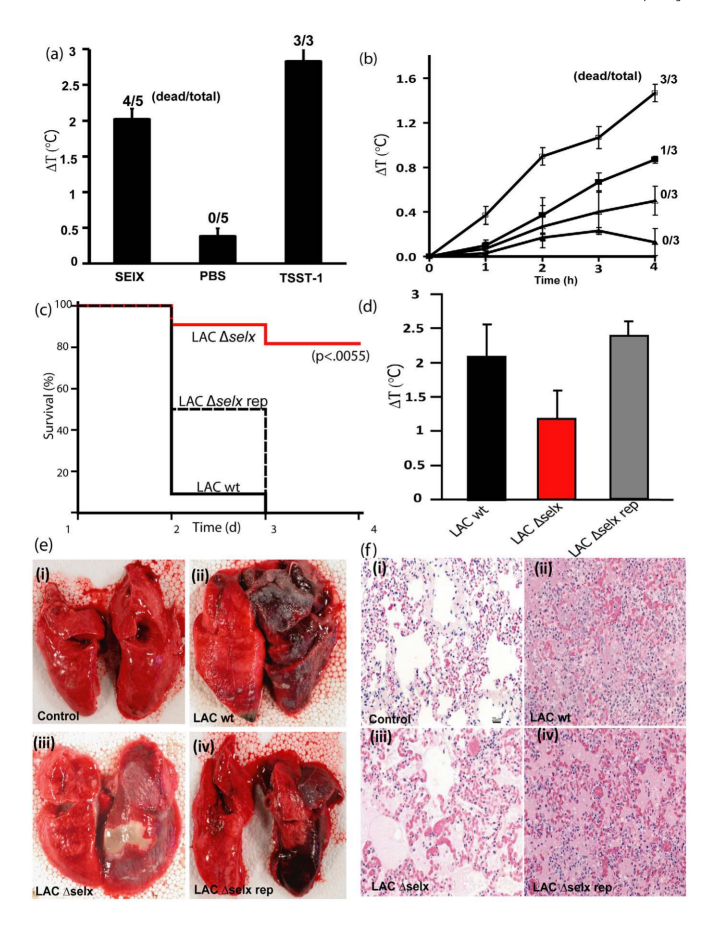


Figure 6. SEIX causes TSS and contributes to the severity and lethality of necrotizing pneumonia caused by CA-MRSA USA300. A) The ability of rSEIX2, administered at a dose of 200 μg/kg, to cause TSS was measured using a standard miniosmotic pump model of TSS. TSST-1 administration was used as the positive control and PBS the negative control. B) Fever in response to intravenous injection of rSEIX2 at concentrations of 10 μg/kg (\square), 0.1 μg/kg (\square), 0.1 μg/kg (\square), 0.6 body weight per ml, or PBS (\triangle), was evaluated over a 4 h test period, followed by the ability to enhance the lethality of LPS over 48 h. C) Kaplan-Meier Curves of % survival of rabbits infected with *S. aureus* LAC, *S. aureus* LACΔ*selx*, and LACΔ*selx* rep. D). Increase in rabbit core temperature (\triangle T °C) 2 d after pulmonary infection with LAC wt, LACΔ*selx*, and LACΔ*selx* rep. E). Gross pathology of lungs from uninfected rabbit (i) or from rabbits infected with LAC wt (iii), LACΔ*selx* (iii), and LACΔ*selx* rep (iv). F) Haematoxylin Eosinstrains.

Bioinformatic analyses

The novel putative SAg gene, selx was first identified by BLAST analysis of the genome of USA300 FPR3757 with the gene sequence specific for the SAg TSST-1. The sequences of distinct selx alleles were identified by BLASTn analysis of S. aureus whole genome sequences deposited in GenBank (Table S1). Additional selx alleles were sequenced with selxseq primers (Table S2) by Genepool Sequencing Service (University of Edinburgh, UK), aligned by ClustalW using MEGA 4.0.1 software [53] and Neighbor-Joining phylogenetic trees constructed using the Nucleotide Maximum Composite Likelihood model. At least 500 bootstrap trees were generated to examine the robustness of the inferred phylogenetic relationships. Recombination detection program RDP v3.44 was used to identify putative end points of recombination events [23]. Unique recombination events were detected by at least 3 of the programs employed by the RDP suite, p<0.05. Protein structural modeling of the derived amino acid sequences of selx2, selxbov1 and selxov was carried out. Predicted 3D structures were obtained for SEIX2, SEIXbov1 and SEIXov with the program Phyre using the crystal structure of TSST-1 as a template, (PDB file c5tssA) [24]. PDB files were generated for each predicted structure and analyzed using Pymol software [54]. Amino acid sequence alignment was carried out using ClustalW and viewed with Jalview [55].

Transcriptional analysis of selx

Total RNA was extracted from S. aureus strains LAC, RF122, and ED133 exponential (OD₆₀₀ = 0.6) and stationary phase (16 h) cultures using the RNeasy miniprep kit (QIAgen) as described in the manufacturer's instructions except for re-suspension in TE buffer with 100 µg/ml Lysostaphin and incubation at 37°C for 20 min. RNA was treated with Turbo DNase (Ambion Inc) and. 0.5 µg mRNA from at least 3 independent total RNA extractions were reverse-transcribed to cDNA with the Power SYBR Green RNA-to-CT 2-Step Kit (Applied Biosystems). To quantify cDNA generated by reverse transcription from target RNA, qRT-PCR reactions were carried out in 25 µl reactions containing 50 ng of cDNA, 300 nM selxq or 16S rRNA primers (Table S2), and SYBR Green I dye master mix (AB), using a Mx3000P light cycler (Strategene). 16S rRNA primers have been described elsewhere [56]. Relative values of transcription of selx were determined by comparative quantification to the internal control 16SrRNA. The thermal conditions were: 10 min at 95°C for 1 cycle, 20 s at 95°C, 20~s at $60^{\circ}\mathrm{C}$ and 20~s at $72^{\circ}\mathrm{C}$ for 40 cycles. RNA samples were processed in triplicate with no template (NTC), no Reverse Transcriptase (no RT) and positive genomic DNA controls. Fluorescence was measured at the end of the annealing phase of each cycle and a threshold value for the fluorescence set by the MxPro qPCR software version 4.1.

Cloning and purification of recombinant SAgs

5' primers for cloning selx2, selxbov1, and selxov into the pET15b plasmid (Novagen), were designed to anneal immediately after the signal peptide coding region, as predicted by Signal P 3.0 Server

(http://www.cbs.dtu.dk/services/SignalP/), and 3' primers were designed to include the stop codon of the gene (Table S2). PCR reactions were carried out using Vent polymerase (NEB) and contained 100 nM selxpET forward and reverse primers designed to incorporate restriction sites NdeI and BamHI (Table S2) and 10 ng of template DNA. PCR products were restriction digested with NdeI and BamHI (NEB), purified, and ligated with T4 DNA ligase (NEB), and transformed into E. coli DH5α cells. pET constructs were isolated from DH5\alpha using the OIAprep Spin Miniprep kit (OIAgen), and transformed into E. coli BL21. BL21 cells containing the pET plasmid constructs were cultured in Luria-broth containing 50 µg/ml ampicillin (Sigma-Aldrich) and induced in mid-exponential phase of growth ($OD_{600} = 0.6$), with 1 mM isopropyl β-D-1-thiogalactopyranoside (IPTG) (ForMedium Ltd.) for 4 h. Cells were recovered by centrifugation at 8000 xg, disrupted using a French Press, and His-tagged recombinant proteins were purified by affinity chromatography on a Ni-NTA nickel affinity column (Invitrogen). Proteins were dialysed using Spectra/Por Float-A-Lyzer tubing with a 8000 to 10000 molecular weight cut off (MWCO) (Spectrum Laboratories), and LPS was removed using ProteoSpin endotoxin removal kit (Norgen Biotek).

Western immunoblot analysis

Supernates from stationary phase cultures of S. aureus strains were concentrated with Amicon Ultra-15 Centrifugal Filter units (10000 MWCO) (Millipore). Recombinant proteins and concentrated secreted proteins were separated by sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE) and transferred to nitrocellulose membranes (Amersham Hybond ECL, GE Healthcare) in Towbin transfer buffer. The membrane was incubated in 1 x PBS (pH 7.3) containing 8% powdered milk (Fluka), at 4°C overnight, washed 3 times with washing buffer, 1 x PBS (pH 7.3) containing 1% powdered milk and 0.05% Tween 20 (Sigma-Aldrich). The membrane was incubated for 1 h with primary antibody, which was either a 1:2500 (bovine and ovine) or 1:5000 (human) dilution of serum, or 2 h with 1:1500 dilution of rat anti-sera raised against rSElXbov1. The membrane was then incubated with horse radish perixodase conjugated (HRP) secondary antibodies for 1 h, which was 1:2500 goat anti-bovine IgG, 1:2500 rabbit anti-sheep IgG, (Santa Cruz Biotechnology), 1:5000 rabbit anti-human IgG (Dako), or 1:1500 goat anti- rat IgG (Abcam), depending on the experiment. Human sera samples were obtained from infectious endocarditis patients [57], ovine serum samples were obtained from experimentally-infected sheep and provided by E. Vautor [58] and from cows with bovine mastitis by C. Smyth. Semi-quantitative spot densitometry was carried out with the ChemiImager 4000i.V4 program, using a MultiImager light cabinet (Alpha Innotech).

Construction of *selx*-deficient and *selx*-repaired derivatives of *S. aureus* USA300 LAC

An selx-deficient strain of USA300 LAC was obtained by allele replacement using the pMAD-CM plasmid [59]. PCR products of



578 bp and 592 bp, flanking the left (AB) and right (CD) regions of selx, respectively were amplified with primers listed in Table S2. PCR amplification with primers A and D was carried out with 100 ng of purified AB and CD template DNA to produce a single spliced AD fragment by overlapping PCR, facilitated by sequence complementary to primer B incorporated into the CD fragment by primer C. AD products were then purified and cloned into the Strataclone pSC-B plasmid (Stratagene). pSC-B plasmid containing the AD insert was digested with EcoRI for 3 h at 37°C. The AD fragment was then purified, ligated into the dephosphorylated gene replacement plasmid, pMAD-CM, to create pMAD::selx, which was transformed by electroporation into LAC. LAC containing pMAD::selx was shaken overnight at 30°C in TSB containing chloramphenicol. To select for integration of the plasmid into the chromosome through homologous recombination, 10-fold dilutions were incubated on TSA containing chloramphenical at 44°C, a temperature nonpermissive for plasmid replication. Double cross-over excisants were selected for by growing integrants in TSB without chloramphenicol at 30°C for 24 h. 10-fold dilutions were plated onto TSA containing Xgal and incubated on TSA at 30°C overnight. White colonies were screened for antibiotic sensitivity on chloramphenicol plates (inferring loss of the plasmid) and screened for WT or mutated forms of the gene by PCR with primers upstream (E) and downstream (Z) of selx (Table S2). The resulting selx-deficient strain, LAC Δ selx was sequenced using E and Z primers in order to confirm the in-frame deletion event. The selx deletion in LAC Δ selx was repaired using an identical allelic exchange approach employing pMAD-CM. Oligonucleotide primers (Table S2) designed to incorporate a single synonymous substitution into the selx gene were used to amplify 2 PCR products specific for left and right regions of the selx gene (rep AB and rep CD primers). The resulting PCR products were spliced to produce an intact selx gene, and cloned into pMAD as previously described resulting in pMAD::selx rep. Allelic exchange was carried out as described for selx mutant construction and the repaired selx gene sequenced. Western blot analysis confirmed the restoration of SEIX expression (Fig. S6a).

Pyrogenicity, endotoxin enhancement and miniosmotic pump lethality studies

American Dutch Belted rabbits were injected with rSEIX2 at doses of 10, 1, and 0.1 µg/kg of body weight per ml intravenously. Three rabbits were injected with each dose and temperature was measured hourly for 4 h. After 4 h, each rabbit was injected intravenously with 1 µg of lipopolysaccharide (LPS) from Salmonella enterica serovar typhimurium (1/500 of the 50% lethal dose of endotoxin alone). SAgs characteristically amplify the lethal effects of LPS by up to 10⁶-fold. Lethality was assessed over a 48 h period [60]. In agreement with the University of Minnesota IACUC, animals that failed to exhibit escape behaviour and at the same time could not right themselves were prematurely euthanized. It is our experience in over 30 years of similar experimentation that this point is 100% predictive of SAg lethality. Animals were euthanized with intravenous injection of 1 ml of Beuthanasia D, whether prematurely or at the end of experimentation. Miniosmotic pumps, containing 200 µg of rSEIX2 or TSST-1, or PBS, were implanted subcutaneously into 5 American Dutch Belted rabbits per group (3 for TSST-1) [61]. Fever was assessed after 24 h, and lethality of the toxins over a period of 7 d. The same criteria as above were used for assessing need for premature euthanasia.

Necrotizing pneumonia rabbit model

Wild-type LAC and the selx knock-out strain were cultured in Todd Hewitt broth for 16 h and washed once in Todd Hewitt broth to remove exoproteins. The bacteria were re-suspended in Todd Hewitt broth at 1×10¹⁰ colony-forming units (CFU)/ml for use in injections. American Dutch Belted rabbits (11 per group for LAC wt and LAC $\Delta selx$, and 4 for LAC $\Delta selx$ rep) were anesthetized with ketamine and xylazine. Their tracheas were exposed and 2×10⁹ USA300 CA-MRSA strain LAC, the isogenic selx-deficient LAC strain or its repaired derivative were administered intra-tracheally through catheters in 0.2 ml volumes. The animals were closed and monitored for 4 d for development of fatal necrotizing pneumonia.

Preparation and stimulation of lymphocytes

Blood was obtained from 2 Holstein-Friesian cattle aged 18-36 m via jugular vein puncture. Animals were reared indoors and maintained on a ration of hay and concentrates. Peripheral blood mononuclear cells (PBMC) were isolated from blood by density gradient centrifugation using Ficoll Paque Plus (GE Healthcare) as described previously [27,62]. Human PBMC were isolated from venous blood of 3 healthy donors as described previously [27]. Splenocytes were obtained from American Dutch Belted rabbits as previously described [63]. Bovine PBMC were adjusted to a concentration of 1×10^6 cells/ml in complete cell culture medium (RPMI 1640, Gibco) supplemented with 10% heat-inactivated FCS, 100 U/ml penicillin, 100 µg/ml streptomycin, 292 µg/ml Lglutamine (PSG) and 50 µM 2-Mercaptoethanol (Sigma-Aldrich), and cultured at 37°C, 5% CO₂. Human PBMC (1 \times 10⁶) were suspended in RPMI 1640 medium (Life technologies) supplemented with 2% FBS, 100 U/ml penicillin G, and 100 µg/ml streptomycin. The cultures were co-incubated with 10-fold dilutions of rSEIX proteins (1 to $10^{-6} \mu g/ml$) for 72 h at 37°C, 5% CO₂. After adding [³H] thymidine (1 μCi), Cells were cultured for a further 18 h and cellular DNA was harvested on glass fiber filters. [3H]-thymidine incorporation was quantified by liquid scintillation counting as described previously [27].

Analysis of Vβ-dependent T-cell activation

Total RNA was extracted from human PBMC prior to and after stimulation with rSEIX proteins (1 µg/ml) for 96 h. Expansion of cells expressing different humV β gene subfamilies was determined using qRT-PCR as described previously [27]. Total RNA was extracted from bovine PBMC before and after stimulation with rSEIX variants (1 µg/ml) using Tri-reagent (Sigma-Aldrich). Firststrand cDNA was generated from 0.5 µg of RNA using Power SYBR Green RNA-to-CT 2-Step Kit (AB). BovVβ subfamilyspecific qRT-PCR primers were designed with Primer3 [64], based on an alignment of bovV β sequences. cDNA sequences for bovine TRBV genes were derived from the bovine genome and cDNA analyses described by Connelley et al [29] (Table S2). To quantify bovV β subfamily gene expression, qRT-PCR reactions were carried out in 25 µl reactions containing 100ng cDNA, 100 nM primers (Table S2), and SYBR Green I dye master mix (AB) using a Stratagene Mx3000P light cycler. The thermal conditions were 1 cycle at 50°C for 10 min, 1 cycle at 95°C for 10 min, 15 s at 95°C and 1 min at 60°C for 40 cycles. RNA samples were processed in triplicate with NTC and noRT controls. The threshold cycle (C_T) was determined using MxPro software version 4.1 and normalized to internal controls, β -actin and Constant β (C β), by calculating ΔC_T [C_T target - C_T $_{\beta ext{-actin}}$ - C_T $_{(\beta ext{-actin}}$ - $_{C}\beta$)]. Normalized ΔC_T data were then compared by calculating - $\Delta\Delta C_{\mathrm{T}}$ = -(ΔC_{T} stimulated - ΔC_{T} of unstimulated). Values >0 indicate expansion of particular subset in response to stimulation.

Statistical analysis

Student's t-test analysis was used to assess differences in fever responses in rabbits and differences in humV β and bovV β gene



expression. Fisher's exact test was used to assess differences in rabbit survival.

Supporting Information

Figure S1 selx is located in the core genome of S. aureus. Schematic representation of the genomic context of selx. selx is depicted in red, the integrase pseudogene is depicted with a blue arrow, white arrows represent hypothetical proteins of unknown function and conserved staphylococcal genes are indicated by black arrows.

(PDF)

Figure S2 Amino-acid sequence alignment of 16 SEIX allelic variants (all alleles except the truncated SEIX13). (PDF)

Figure S3 Identification of predicted recombination events among selx alleles. Coordinates of beginning breakpoints (BB) and end breakpoints (EB) detected by at least 3 different programmes are indicated. Differing filled patterns indicate gene fragments of distinct evolutionary origin. Colored outlines and letters indicate different selx alleles. (PDF)

Figure S4 Phylogenetic tree of *selx.* A neighbour joining tree based on *selx* gene sequences has a distinct topology to a concatenated multilocus sequence-based tree (Fig. 1b). Bootstrap values greater than 40 are indicated. (PDF)

Figure S5 Western blot analysis indicates a lack of cross-reactivity of SEIX antibodies for TSST-1 or SSL7. Western blot analysis of recombinant SEIX TSST-1, SSL7 with SEIX-specific antisera raised in rats and serum samples from human and ovine infections. (PDF)

Figure S6 Phenotypic analysis of the LAC $\Delta selx$ mutant and LAC $\Delta selx$ rep. a) Western blot analysis of LAC wt, LAC $\Delta selx$, and LAC $\Delta selx$ repaired, with SEIX-specific antibody. b) Hemolytic titration of LAC wt, LAC $\Delta selx$ mutant and LAC $\Delta selx$ repaired supernatants incubated with washed rabbit erythrocytes.

References

- Thomas D, Chou S, Dauwalder O, Lina G (2007) Diversity in Staphylococcus aureus enterotoxins. Chem Immunol Allergy 93: 24–41.
- Dinges MM, Orwin PM, Schlievert PM (2000) Exotoxins of Staphylococcus aureus. Clin Microbiol Rev 13: 16–34.
- Thomas D, Dauwalder O, Brun V, Badiou C, Ferry T, et al. (2009) Staphylococcus aureus superantigens elicit redundant and extensive human Vbeta patterns. Infect Immun 77: 2043–2050.
- Ono HK, Omoe K, Imanishi Ki, Iwakabe Y, Hu D-L, et al. (2008) Identification and Characterization of Two Novel Staphylococcal Enterotoxins, Types S and T. Infect Immun 76: 4999–5005.
- Seo KS, Bohach GA (2007) Staphylococcus aureus; MP. Doyle LB, TJ. Montville, eds. Washington D.C: ASM Press.
- Fraser JD, Proft T (2008) The bacterial superantigen and superantigen-like proteins. Immunol Rev 225: 226–243.
- Choi YW, Kotzin B, Herron L, Callahan J, Marrack P, et al. (1989) Interaction of Staphylococcus aureus toxin "superantigens" with human T cells. Proc Natl Acad Sci U S A 86: 8941–8945.
- Dellabona P, Peccoud J, Kappler J, Marrack P, Benoist C, et al. (1990) Superantigens interact with MHC class II molecules outside of the antigen groove. Cell 62: 1115–1121.
- Drake CG, Kotzin BL (1992) Superantigens: biology, immunology, and potential role in disease. J Clin Immunol 12: 149–162.
- Kawabe Y, Ochi A (1990) Selective anergy of V beta 8+,CD4+ T cells in Staphylococcus enterotoxin B-primed mice. J Exp Med 172: 1065–1070.
- Fitzgerald JR, Monday SR, Foster TJ, Bohach GA, Hartigan PJ, et al. (2001) Characterization of a putative pathogenicity island from bovine Staphylococcus aureus encoding multiple superantigens. J Bacteriol 183: 63–70.

Hemolytic titre was determined to be the reciprocal of the dilution which resulted in $\sim 50\%$ hemolysis (circled in black). c) SDS PAGE analysis of concentrated supernatant protein fractions resulted in indistinguishable profiles. In addition, quantification of α -toxin and PVL levels in LAC wt and LAC Δ selx supernatants revealed identical toxin levels at 3 h, 6 h, 8 h, and 24 h time-points during growth in CCY medium by specific enzyme-linked immunosorbent assays (data not shown) (α -toxin, bioMerieux, Nabi Biopharmaceuticals, and PVL; Besseyre des Horts et al, Infect Immun. 2010 78:260-4). (PDF)

Table S1 Distribution and coordinates of selx in sequenced S. aureus genomes. (DOC)

Table S2 Oligonucleotide primers used in this study. (DOC)

Table S3 S. aureus strains employed and selx genotype and phenotype. (DOC)

Acknowledgments

The authors are grateful to F. DeLeo for providing USA300 strain LAC, A. Cheung for pMAD-CM, C. Smyth, D. Smyth and J. Lindsay for clinical isolates, and to C. Smyth, E. Vautor, and S. Daga for patient sera samples. In addition, we thank J Etienne, G Lina and C. Badiou for ELISA determination of PVL and α -toxin levels, K. Taylor and A. Fattom for providing α -toxin-specific antibody and the ELISA protocol. In addition, we thank P. Beard for assistance with histopathology, I. Poxton, N. MacHugh and Danielle Potter for technical assistance, and S. Tong for providing the sequence of selx from an ST75 strain. Finally, we thank 3 anonymous reviewers for constructive comments which helped to improve the manuscript markedly.

Author Contributions

Conceived and designed the experiments: GJW KSS GAB PMS WIM JRF. Performed the experiments: GJW KSS RAC ONCS JAM JYP PMS JRF. Analyzed the data: GJW KSS RAC TC ONCS JAM CMG JYP GAB PMS WIM JRF. Wrote the paper: GJW JRF.

- Jarraud S, Peyrat MA, Lim A, Tristan A, Bes M, et al. (2001) egc, a highly prevalent operon of enterotoxin gene, forms a putative nursery of superantigens in Staphylococcus aureus. J Immunol 166: 669–677.
- Johns MB, Jr., Khan SA (1988) Staphylococcal enterotoxin B gene is associated with a discrete genetic element. J Bacteriol 170: 4033–4039.
- Ben Zakour NL, Guinane CM, Fitzgerald JR (2008) Pathogenomics of the staphylococci: insights into niche adaptation and the emergence of new virulent strains. FEMS Microbiol Lett 289: 1–12.
- 15. Smyth DS, Hartigan PJ, Meaney WJ, Fitzgerald JR, Deobald CF, et al. (2005) Superantigen genes encoded by the egc cluster and SaPIbov are predominant among Staphylococcus aureus isolates from cows, goats, sheep, rabbits and poultry. J Med Microbiol 54: 401–411.
- Omoe K, Hu DL, Takahashi-Omoe H, Nakane A, Shinagawa K (2005) Comprehensive analysis of classical and newly described staphylococcal superantigenic toxin genes in Staphylococcus aureus isolates. FEMS Microbiol Lett 246: 191–198.
- Chambers HF (2005) Community-associated MRSA--resistance and virulence converge. N Engl J Med 352: 1485–1487.
- Daum RS, Ito T, Hiramatsu K, Hussain F, Mongkolrattanothai K, et al. (2002)
 A novel methicillin-resistance cassette in community-acquired methicillin-resistant Staphylococcus aureus isolates of diverse genetic backgrounds. J Infect Dis 186: 1344–1347.
- Klevens RM, Morrison MA, Nadle J, Petit S, Gershman K, et al. (2007) Invasive methicillin-resistant Staphylococcus aureus infections in the United States. JAMA 298: 1763–1771.
- 20. Francis JS, Doherty MC, Lopatin U, Johnston CP, Sinha G, et al. (2005) Severe community-onset pneumonia in healthy adults caused by methicillin-resistant



- Staphylococcus aureus carrying the Panton-Valentine leukocidin genes. Clin Infect Dis 40: 100–107.
- Diep BA, Gill SR, Chang RF, Phan TH, Chen JH, et al. (2006) Complete genome sequence of USA300, an epidemic clone of community-acquired meticillin-resistant Staphylococcus aureus. Lancet 367: 731–739.
- Kwan T, Liu J, DuBow M, Gros P, Pelletier J (2005) The complete genomes and proteomes of 27 Staphylococcus aureus bacteriophages. Proc Natl Acad Sci U S A 102: 5174–5179.
- Heath L, van der Walt E, Varsani A, Martin DP (2006) Recombination Patterns in Aphthoviruses Mirror Those Found in Other Picornaviruses. J Virol 80: 11827–11832.
- Kelley LA, Sternberg MJ (2009) Protein structure prediction on the Web: a case study using the Phyre server. Nat Protoc 4: 363–371.
- Burroughs AM, Balaji S, Iyer LM, Aravind L (2007) Small but versatile: the extraordinary functional and structural diversity of the beta-grasp fold. Biol Direct 2: 18.
- McCormick JK, Tripp TJ, Llera AS, Sundberg EJ, Dinges MM, et al. (2003) Functional Analysis of the TCR Binding Domain of Toxic Shock Syndrome Toxin-1 Predicts Further Diversity in MHC Class II/Superantigen/TCR Ternary Complexes. J Immunol 171: 1385–1392.
- Seo KS, Park JY, Terman DS, Bohach GA (2010) A quantitative real time PCR method to analyze T cell receptor Vbeta subgroup expansion by staphylococcal superantigens. J Transl Med 8: 2.
- Deringer JR, Ely RJ, Monday SR, Stauffacher CV, Bohach GA (1997) Vbetadependent stimulation of bovine and human T cells by host-specific staphylococcal enterotoxins. Infect Immun 65: 4048–4054.
- Connelley T, Aerts J, Law A, Morrison WI (2009) Genomic analysis reveals extensive gene duplication within the bovine TRB locus. BMC Genomics 10: 192.
- Elsik CG, Tellam RL, Worley KC, Gibbs RA, Muzny DM, et al. (2009) The genome sequence of taurine cattle: a window to ruminant biology and evolution. Science 324: 522–528.
- Herold BC, Immergluck LC, Maranan MC, Lauderdale DS, Gaskin RE, et al. (1998) Community-acquired methicillin-resistant Staphylococcus aureus in children with no identified predisposing risk. JAMA 279: 593–598.
- Somerville GA, Beres SB, Fitzgerald JR, DeLeo FR, Cole RL, et al. (2002) In Vitro Serial Passage of Staphylococcus aureus: Changes in Physiology, Virulence Factor Production, and agr Nucleotide Sequence. J Bacteriol 184: 1430–1437.
- Strandberg KL, Rotschafer JH, Vetter SM, Buonpane RA, Kranz DM, et al. (2010) Staphylococcal superantigens cause lethal pulmonary disease in rabbits. J Infect Dis 202: 1690–1697.
- Lindsay JA, Moore CE, Day NP, Peacock SJ, Witney AA, et al. (2006) Microarrays reveal that each of the ten dominant lineages of Staphylococcus aureus has a unique combination of surface-associated and regulatory genes. J Bacteriol 188: 669–676.
- Holden MT, Feil EJ, Lindsay JA, Peacock SJ, Day NP, et al. (2004) Complete genomes of two clinical Staphylococcus aureus strains: evidence for the rapid evolution of virulence and drug resistance. Proc Natl Acad Sci U S A 101: 9786–9791.
- Proft T, Moffatt SL, Weller KD, Paterson A, Martin D, et al. (2000) The streptococcal superantigen SMEZ exhibits wide allelic variation, mosaic structure, and significant antigenic variation. J Exp Med 191: 1765–1776.
- Lowder BV, Guinane CM, Ben Zakour NL, Weinert LA, Conway-Morris A, et al. (2009) Recent human-to-poultry host jump, adaptation, and pandemic spread of Staphylococcus aureus. Proc Natl Acad Sci U S A 106: 19545–19550.
- Guinane CM, Ben Zakour NL, Tormo-Mas MA, Weinert LA, Lowder BV, et al. (2010) Evolutionary Genomics of Staphylococcus aureus Reveals Insights into the Origin and Molecular Basis of Ruminant Host Adaptation. Genome Biol Evol 2: 454–466.
- Marr JC, Lyon JD, Roberson JR, Lupher M, Davis WC, et al. (1993) Characterization of novel type C staphylococcal enterotoxins: biological and evolutionary implications. Infect Immun 61: 4254–4262.
- Feil EJ, Cooper JE, Grundmann H, Robinson DA, Enright MC, et al. (2003) How clonal is Staphylococcus aureus? J Bacteriol 185: 3307–3316.
- Robinson DA, Monk AB, Cooper JE, Feil EJ, Enright MC (2005) Evolutionary Genetics of the Accessory Gene Regulator (agr) Locus in Staphylococcus aureus. J Bacteriol 187: 8312–8321.
- Watanabe S, Ito T, Sasaki T, Li S, Uchiyama I, et al. (2009) Genetic diversity of staphylocoagulase genes (coa): insight into the evolution of variable chromosomal virulence factors in Staphylococcus aureus. PLoS One 4: e5714.

- Li M, Diep BA, Villaruz AE, Braughton KR, Jiang X, et al. (2009) Evolution of virulence in epidemic community-associated methicillin-resistant Staphylococcus aureus. Proc Natl Acad Sci U S A 106: 5883–5888.
- Otto M (2010) Basis of virulence in community-associated methicillin-resistant Staphylococcus aureus. Annu Rev Microbiol 64: 143–162.
- Li M, Cheung GY, Hu J, Wang D, Joo HS, et al. (2010) Comparative analysis of virulence and toxin expression of global community-associated methicillinresistant Staphylococcus aureus strains. J Infect Dis 202: 1866–1876.
- Schlievert PM (2009) Cytolysins, superantigens, and pneumonia due to community-associated methicillin-resistant Staphylococcus aureus. J Infect Dis 200: 676–678.
- Montgomery CP, Boyle-Vavra S, Adem PV, Lee JC, Husain AN, et al. (2008) Comparison of virulence in community-associated methicillin-resistant Staphylococcus aureus pulsotypes USA300 and USA400 in a rat model of pneumonia. J Infect Dis 198: 561–570.
- Labandeira-Rey M, Couzon F, Boisset S, Brown EL, Bes M, et al. (2007) Staphylococcus aureus Panton-Valentine leukocidin causes necrotizing pneumonia. Science 315: 1130–1133.
- Diep BA, Chan L, Tattevin P, Kajikawa O, Martin TR, et al. (2010) Polymorphonuclear leukocytes mediate Staphylococcus aureus Panton-Valentine leukocidin-induced lung inflammation and injury. Proc Natl Acad Sci U S A 107: 5587–5592.
- Bubeck Wardenburg J, Bae T, Otto M, Deleo FR, Schneewind O (2007) Poring over pores: alpha-hemolysin and Panton-Valentine leukocidin in Staphylococcus aureus pneumonia. Nat Med 13: 1405–1406.
- Assimacopoulos AP, Strandberg KL, Rotschafer JH, Schlievert PM (2009)
 Extreme pyrexia and rapid death due to Staphylococcus aureus infection: analysis of 2 cases. Clin Infect Dis 48: 612–614.
- Dinges MM, Schlievert PM (2001) Comparative analysis of lipopolysaccharideinduced tumor necrosis factor alpha activity in serum and lethality in mice and rabbits pretreated with the staphylococcal superantigen toxic shock syndrome toxin 1. Infect Immun 69: 7169–7172.
- Tamura K, Dudley J, Nei M, Kumar S (2007) MEGA4: Molecular Evolutionary Genetics Analysis (MEGA) Software Version 4.0. Mol Biol Evol 24: 1596–1599.
- DeLano WL (2002) The PyMOL Molecular Graphics System. In: Scientific D, ed. San CarlosCA, , USA, .
- Waterhouse AM, Procter JB, Martin DM, Clamp M, Barton GJ (2009) Jalview Version 2--a multiple sequence alignment editor and analysis workbench. Bioinformatics 25: 1189–1191.
- Ster C, Gilbert FB, Cochard T, Poutrel B (2005) Transcriptional profiles of regulatory and virulence factors of Staphylococcus aureus of bovine origin: oxygen impact and strain-to-strain variations. Mol Cell Probes 19: 227–235.
- Daga S, Shepherd J, Callaghan G, Hung R, Dawson D, et al. (2011) Platelet receptor polymorphisms do not influence Staphylococcus aureus-platelet interactions or infective endocarditis. Microbes Infect 13: 216–225.
- Le Maréchal C, Jan G, Even S, McCulloch JA, Azevedo V, et al. (2009) Development of serological proteome analysis of mastitis by Staphylococcus aureus in ewes. J Microbiol Methods 79: 131–136.
- Memmi G, Filipe SR, Pinho MG, Fu Z, Cheung A (2008) Staphylococcus aureus PBP4 is essential for beta-lactam resistance in community-acquired methicillin-resistant strains. Antimicrob Agents Chemother 52: 3955–3966.
- Schlievert PM (1982) Enhancement of host susceptibility to lethal endotoxin shock by staphylococcal pyrogenic exotoxin type C. Infect Immun 36: 123–128.
- Parsonnet J, Gillis ZA, Richter AG, Pier GB (1987) A rabbit model of toxic shock syndrome that uses a constant, subcutaneous infusion of toxic shock syndrome toxin 1. Infect Immun 55: 1070–1076.
- Goddeeris BM, Morrison WI (1988) Techniques for the generation, cloning, and characterization of bovine cytotoxic T cells specific for the protozoan Theileria parva. Methods Cell Sci 11: 101–110.
- Barsumian EL, Schlievert PM, Watson DW (1978) Nonspecific and specific immunological mitogenicity by group A streptococcal pyrogenic exotoxins. Infect Immun 22: 681–688.
- 64. Rozen S (2000) Primer3 on the WWW for general users and for biologist programmers. In: Krawetz S MSe, ed. Bioinformatics Methods and Protocols: Methods in Molecular Biology. TotowaNJ: Humana Press. pp 365–386.
- Arden B, Clark SP, Kabelitz D, Mak TW (1995) Human T-cell receptor variable gene segment families. Immunogenetics 42: 455–500.

