Hyperexpression of α-hemolysin explains enhanced virulence of sequence type 93 community-associated methicillin-resistant Staphylococcus aureus

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Abstract

Background: The community-associated methicillin-resistant S. aureus (CA-MRSA) ST93 clone is becoming dominant in Australia and is clinically highly virulent. In addition, sepsis and skin infection models demonstrate that ST93 CA-MRSA is the most virulent global clone of S. aureus tested to date. While the determinants of virulence have been studied in other clones of CA-MRSA, the basis for hypervirulence in ST93 CA-MRSA has not been defined.

Results: Here, using a geographically and temporally dispersed collection of ST93 isolates we demonstrate that the ST93 population hyperexpresses key CA-MRSA exotoxins, in particular α-hemolysin, in comparison to other global clones. Gene deletion and complementation studies, and virulence comparisons in a murine skin infection model, showed unequivocally that increased expression of α-hemolysin is the key staphylococcal virulence determinant for this clone. Genome sequencing and comparative genomics of strains with divergent exotoxin profiles demonstrated that, like other S. aureus clones, the quorum sensing agr system is the master regulator of toxin expression and virulence in ST93 CA-MRSA. However, we also identified a previously uncharacterized AraC/XylS family regulator (AryK) that potentiates toxin expression and virulence in S. aureus.

Conclusions: These data demonstrate that hyperexpression of α-hemolysin mediates enhanced virulence in ST93 CA-MRSA, and additional control of exotoxin production, in particular α-hemolysin, mediated by regulatory systems other than agr have the potential to fine-tune virulence in CA-MRSA.

Keywords: Staphylococcus aureus, CA-MRSA, Pathogenesis, Alpha-hemolysin

Background

Community-associated methicillin-resistant Staphylococcus aureus (CA-MRSA) is an emerging global problem with very similar clinical presentations across different clones, despite significant genetic diversity [1]. Many CA-MRSA strains carry lukSF-PV in the accessory genome, which encodes the Panton-Valentine leukocidin (PVL), an exotoxin that causes neutrophil lysis [1]. Although there has been considerable controversy as to the role of this toxin in CA-MRSA pathogenesis, some of this may be explained by a variable, species dependent susceptibility to PVL – human and rabbit neutrophils are lysed by PVL at very low concentrations whilst mouse and monkey neutrophils are less susceptible, making the interpretation of animal model data difficult in some cases [2]. Additionally, the importance of PVL is also likely to be dependent on the site of infection. In the rabbit pneumonia model, PVL has been demonstrated to have a clear role in mediating severe lung necrosis and inflammation [3]. In contrast, in skin infection, even in the rabbit model, its role remains less clear [4,5].

Notwithstanding PVL, the increased expression of other core genome virulence determinants also contributes significantly to the increased virulence of CA-MRSA strains.
[6,7]. These include α-hemolysin (Hla) and α-type phenol soluble modulins (PSMs). Hla is a pore-forming exotoxin that lyses many cells including red cells, platelets, monocytes and endothelial cells [8]. Hla has been demonstrated to be an important mediator of virulence in skin infection and pneumonia [9,10]. The α-type PSMs have been recently characterized and they lyse neutrophils and red cells [11,12]. The α-type PSMs also mediate virulence in skin infection and septicemia and of these, PSMα3 is the most potent [11].

The study of unique, distantly related CA-MRSA clones that also demonstrate enhanced virulence, may provide insights into the emergence of the global CA-MRSA phenomenon, and also help define the genomic determinants of enhanced virulence. In Australia, the singleton ST93 CA-MRSA clone ST93 (“Queensland clone”) has become dominant in the community [13], and we and others have demonstrated that our reference ST93 strain JKD6159 was highly virulent and caused severe skin infection in a mouse model compared to other CA-MRSA strains including USA300 [14,15].

In this study we used exotoxin analysis, functional genomics and a murine infection model to investigate the relative contribution of α-hemolysin, α-type phenol soluble modulins and Panton-Valentine leukocidin to the enhanced virulence of ST93 CA-MRSA. We show that increased virulence in the BALB/c mouse skin infection model is less dependent on α-type phenol soluble modulin or Panton-Valentine leukocidin production but is instead due to high-level expression of α-hemolysin in this clone, controlled predominantly by the agr system.

**Results and discussion**

The emergence of CA-MRSA is a major public health issue, and there is a clear need to understand the basis for both virulence and transmission of global clones of CA-MRSA. The genetically distinct CA-MRSA clone ST93-IV [28] has rapidly become the dominant clone in Australia and its rise accounts for the increase in incidence of CA-MRSA as a whole in this country [13]. We, and others have previously shown that ST93 strain JKD6159 is the most virulent global clone of *S. aureus* in murine models [14,15]. To determine the mediators of virulence in this clone we initially studied exotoxin expression in a large collection of ST93 *S. aureus* from around Australia, and compared representative high and low expressing strains to an international selection of clones.

**Exotoxin expression in ST93 CA-MRSA strains**

*Staphylococcus aureus* expresses a wide range of exotoxins that may contribute to virulence. Because Hla, PVL and α-type PSMs have been found by others to be important virulence factors in CA-MRSA strains [9,11,16], we measured *in vitro* expression of these exotoxins by the wildtype ST93 strains and non-ST93 comparator strains. The main isolates used in this study are described in Table 1, while the collection of ST93 isolates from around Australia used for comparative exotoxin expression is from a study by Coombs et al. [17] and summarized in Additional file 1. The comparison of expression of international clones to the ST93 reference strain JKD6159 and three additional ST93 strains selected for genome sequencing (see below) are shown in Figure 1, while the results for all 59 ST93 isolates compared to USA300 are shown in Additional file 2 (α-type PSMs) and Additional file 3 (Hla). The results of PVL analysis for the ST93 collection has been previously reported [17]. Because PVL is a 2-component exotoxin and both LukS-PV and LukF-PV are required for activity, we chose to measure LukF-PV expression by quantitative Western blot. LukF-PV was chosen over LukS-PV to obtain anti-LukF-PV antibody with increased specificity of binding as there was more sequence divergence between *lukF-PV* and the orthologous 2-component *S. aureus* exotoxins compared to *lukS-PV*. Although there are four α-type PSMs, PSMα3 causes the most significant neutrophil lysis [11] and we measured deformylated and N-formylated PSMα3 expression by high performance liquid chromatography (HPLC). The proportion of deformylated and formylated forms of PSMs depends on the growth conditions, the activity of *agr* system.

**PVL**

As previously reported [17], PVL expression was consistent across most ST93 strains. We found that there was no significant difference in the LukF-PV expression in the PVL positive strains JKD6159, TPS3104, USA300 and JKD6177. Although USA300 appeared to produce less LukF-PV than JKD6159, the difference was not statistically significant (p = 0.0943, Figure 1A).

**PSMα3**

We found that the deformylated form of PSMα3 was almost always more abundant than the N-formylated form (Figure 1B and Additional file 2). The ST30 CA-MRSA strain JKD6177 did not produce any PSMα3. There was no significant difference in PSMα3 expression between JKD6159 compared to USA300, however JKD6159 produced more PSMα3 compared to JKD6272 (p = 0.0003) and JKD6009 (p = 0.0003). Compared to the other ST93 MRSA strains, JKD6159 produced more PSMα3 compared to TPS3105 (p < 0.0001), and TPS3106 (p = 0.01) but less than TPS3104 (p = 0.0029) (Figure 1B). Expression levels across the whole ST93 collection were variable, although many isolates produced levels at least equivalent to USA300 (Additional file 2).
| S. aureus strain | Molecular type | Date of isolation | Place of isolation | Site of isolation | Relevant characteristics | $\textit{lukSF-PV}$ | reference |
|-----------------|----------------|-------------------|-------------------|-------------------|--------------------------|----------------|----------|
| **Clinical isolates** |
| JKD6159         | ST93-IV [2B]   | 2004              | Victoria, Australia | Blood             | Dominant Australian CA-MRSA clone | +             | [14]     |
| TPS3105         | ST93-IV [2B]   | 2009              | Western Australia, Australia | Nasal cavity | Dominant Australian CA-MRSA clone | +             | This study |
| TPS3106         | ST93-IV [SC2&5] | 2008              | Western Australia, Australia | Blood | Australian CA-MRSA clone | -             | This study |
| JKD6272         | ST1-IV [2B]    | 2002              | Victoria, Australia | Blood             | Australian CA-MRSA clone | -             | [14]     |
| JKD6260         | ST1-IV [2B]    | 2008              | Western Australia, Australia | Skin | Australian CA-MRSA clone | +             | [14]     |
| JKD6177         | ST30-IV [2B]   | 2003              | Melbourne, Australia | Blood             | Australian CA-MRSA clone | +             | [14]     |
| FPR3757 USA300  | ST8-IV [2B]    | NA                | San Francisco, USA | Wrist abscess    | Dominant North American CA-MRSA clone | +             | [19]     |
| JKD6009         | ST239-III [3A] | 2002              | New Zealand        | Wound             | Dominant Australian hospital-associated MRSA clone, AUS2/3 | -             | [20]     |
| **Mutant strains** |
| JKD6159ΔlukSF-PV | ST93-IV [2B]   |                  |                   |                   | Isogenic unmarked $\textit{lukSF-PV}$ KO of JKD6159 | -             | This study |
| JKD6159Δhla     | ST93-IV [2B]   |                  |                   |                   | Isogenic unmarked $\textit{hla}$ KO of JKD6159. Deletion encompassed genome coordinates 1121291–1120441 | +             | This study |
| JKD6159Δhla r   | ST93-IV [2B]   |                  |                   |                   | Isogenic unmarked $\textit{hla}$ KO repaired in JKD6159Δhla. Introduction of a novel PstI site within $\textit{hla}$ | +             | This study |
| JKD6159ΔpsmA    | ST93-IV [2B]   |                  |                   |                   | Isogenic unmarked $\textit{psm-\alpha}$ KO in JKD6159. Deletion encompassed genome coordinates 453364–45378 | +             | This study |
| JKD6159ΔpsmA r  | ST93-IV [2B]   |                  |                   |                   | Isogenic unmarked $\textit{psm-\alpha}$ KO repaired in JKD6159ΔpsmA. Introduction of a novel Sall site within $\textit{psm-\alpha}$ | +             | This study |
| JKD6159Δ00043   | ST93-IV [2B]   |                  |                   |                   | Isogenic unmarked SAA6159_00043 KO of JKD6159. Deletion encompassed genome coordinates S3156 - 54561 | +             | This study |
| JKD6159_AraCr   | ST93-IV [2B]   |                  |                   |                   | Isogenic AraC/XylS regulator repaired in JKD6159 | +             | This study |
| TPS3105r        | ST93-IV [2B]   |                  |                   |                   | Isogenic agrA repair of TPS3105 | -             | This study |

KO: knockout, NA: not available.
Hla expression appeared high for the majority of ST93 isolates, with the exception of four strains where expression was low (Additional file 3). JKD6159 produced greater levels of Hla than all the wildtype strains, including USA300 (p < 0.0001 for all strains except JKD6177, p = 0.0107, Figure 1C). There was no difference in Hla expression between JKD6159 and TPS3104.

Here we have demonstrated that the majority of ST93 strains consistently produce higher levels of Hla compared to other clones, including USA300, while production of PVL and α-type PSM is similar, suggesting that enhanced expression of Hla may be responsible for increased virulence of ST93 CA-MRSA.

Comparative virulence of ST93 isolates with differential exotoxin expression

To further examine the role of these selected exotoxins in our mouse skin infection assay, we compared the virulence of four ST93 isolates selected based on their exotoxin expression profiles (high exotoxin expression, JKD6159 and TPS3104; low exotoxin expression, TPS3105 and TPS3106). TPS3104 was as virulent as JKD6159 in the mouse model in all outcome measures (Figure 2). In contrast, the strains with reduced exotoxin expression TPS3105 and TPS3106 were significantly less virulent compared to JKD6159, with less weight loss at day 5 of infection (p < 0.0001), smaller lesion size (p < 0.0001) and
less CFU recovery from lesions (TPS3105, p = 0.0177; TPS3106, p = 0.0328) in the model (Figure 2).

Impact of exotoxin expression on virulence of ST93 CA-MRSA in the murine skin infection model
To further characterize the contribution of each of the exotoxins to disease in the murine model, genetic deletion and complementation experiments were performed for each of the selected toxins.

Hla
Given the increased in vitro expression of Hla by JKD6159 and TPS3104 and the apparent correlation of this increased expression with increased virulence in the mouse skin infection model, we generated JKD6159Δhla and assessed this mutant in the mouse skin infection assay (Figure 3). There was a marked attenuation in virulence in all outcome measures with significantly decreased weight loss (p < 0.0001), lesion size (p < 0.0001) and CFU recovery (p = 0.0177). To confirm that an unintentional mutation introduced during the procedure to knock-out hla was not responsible for the reduced virulence in this strain, complete genome sequencing of the strain using Ion Torrent sequencing was performed. Mapping of sequence reads from JKD6159Δhla against JKD6159 (40× genome coverage) demonstrated no additional differences between JKD6159 and JKD6159Δhla. We also restored Hla function in this strain by using allelic replacement to repair the deletion in hla and create strain JKD6159Δhla r. As expected, Hla expression was absent in JKD6159Δhla and expression was restored in JKD6159Δhla r when tested by Western Blot (Additional file 4A). JKD6159Δhla r also reverted to high virulence in the mouse skin infection assay (Figure 3). The apparent slight reduction in virulence of this hla repaired strain compared to wild type JKD6159 is explained by incomplete penetration of the restored hla allele in JKD6159Δhla r, resulting in mixed bacterial populations and reversion to JKD6159Δhla for some of the mice (Additional file 4B and C).

α-type PSMs
In order to determine the contribution of α-type PSMs to virulence of JKD6159, we generated JKD6159Δpsma (deletion of the whole α-type PSM locus) and assessed this mutant in the mouse skin infection assay (Figure 3). There was no significant difference in virulence in all outcome measures; weight loss (p = 0.06), lesion size (p = 0.8174) and CFU recovery (p = 0.1925). The absence of PSMa3 production was verified by HPLC and the integrity of the JKD6159Δpsma was confirmed by complementation (Additional file 5). It is noteworthy that the level of expression of PSMa3 by JKD6159 was similar to USA300 (Figure 1), a strain that produces high levels of PSMs and where a contribution to virulence has been demonstrated [7,11]. Despite this, the deletion mutant (JKD6159Δpsma) demonstrated no attenuation of virulence compared to JKD6159 (Figure 3). The significantly divergent genetic background of ST93 compared with USA300 may account for this difference in the importance of α-type PSMs to the virulence of JKD6159 [6].

PVL
We constructed an isogenic PVL negative mutant in JKD6159 by deleting lukSF-PV. Western Blot analysis confirmed the absence of LukF-PV in the mutant (Additional file 6). Assessment of the JKD6159ΔlukSF-PV mutant in the mouse skin infection model showed no decrease in virulence (Figure 3). Therefore PVL was not contributing to the increased virulence in JKD6159 in this
murine model. Murine neutrophils, unlike rabbit and human neutrophils are relatively resistant to the effects of PVL so it is difficult to draw firm conclusions as to the human importance of this result [2]. However, the aim of this study was to uncover the mechanisms for the observed increased virulence of ST93 previously demonstrated using this mouse model [14]. Our results reinforce the results of others who have examined different S. aureus clones which indicate that Hla, rather than PVL is the main mediator of virulence in CA-MRSA in a mouse skin infection model [9,10,21,22]. It should be noted that other authors have concluded that the rabbit skin infection model gave very similar results to the mouse model for infection at the same site [4]. Nonetheless, testing of our PVL deletion mutant in a rabbit model may be warranted in future.

Genome sequencing of three additional ST93 isolates
We have previously fully sequenced and annotated the genome of ST93 strain JKD6159 [14,23]. The differential virulence and exotoxin expression of some ST93 isolates compared to JKD6159 was then exploited by using whole genome sequencing and comparative genomics to determine the genetic basis for exotoxin expression in this clone. We selected the high expression strain TPS3104 and the low virulence and expression strains TPS3105 and TPS3106 to compare to JKD6159. De novo assembly of each of these strains resulted in ~700 contigs per isolate, with a genome length of 2.8 Mbp. The de novo assembly metrics are summarized in Additional file 7. The contigs were aligned to JKD6159 using BLASTN, with some important differences demonstrated between the strains (Figure 4A). TPS3104 contained SCCmecIV and φSA2 with lukSF-PV; TPS3105 contained SCCmecIV but lacked φSA2 and lukSF-PV; TPS3106 contained SCCmecV, and φSA2 without lukSF-PV. In addition, read-mapping against the complete JKD6159 genome (chromosome and plasmid: 2,832,164 bp) was then employed to define a 2,720,685 bp core genome among these four isolates (96% of the JKD6159 genome), revealing 253 polymorphic nucleotide positions, some of which were common to two strains. A phylogeny was inferred that confirmed the close relationship among all isolates, with TPS3106 more distantly related to the others (Figure 4B). The unmapped reads from each isolate were also subjected to de novo assembly to identify DNA not present in JKD6159. TPS3104 and TPS3105 contained no new sequences, while TPS3106 contained 34 kb of additional DNA, predominantly spanning the SCCmecV region.

Comparative genomics of ST93 and the importance of agr in the virulence of ST93 CA-MRSA
We next explored the contribution of specific mutations to the differential virulence of the ST93 strains. Using our read mapping approach described above, we compared the genome sequences of TPS3104, TPS3105 and TPS3106
with each other and with JKD6159. There were a number of single nucleotide polymorphisms (SNPs) and insertions and deletions (indels) differentiating the strains from JKD6159 (Additional files 8, 9, 10). We searched for mutations in regulatory genes that could potentially explain the different virulence phenotypes of the strains. Notably, both avirulent ST93 strains, TPS3105 and TPS3106 contained mutations within the \(agr\) locus. We have since completed whole genome sequencing of TPS3151 and TPS3161 and found they contain predicted amino acid substitutions in AgrC that might disrupt \(agr\) function (Stinear et al., submitted). These isolates demonstrated low expression of Hla (Additional file 3).

Additionally, TPS3106 also contained a mutation in a gene encoding a previously uncharacterized AraC/XylS family regulatory protein. This was also of particular interest as members of this class have been shown to contribute to the regulation of exotoxin expression [24,25].

TPS3105 contained a frame-shift mutation within \(agrA\) (Sa_JKD6159 nucleotide 2096502) and a further substitution (G to A) within \(agrA\) at nucleotide 2096569), while TPS3106 contained an \(\sim\)356 bp deletion spanning the \(agr\) effector molecule, RNAIII (deletion spanning nucleotides 2093372 to 2093728). These mutations suggested these isolates were \(agr\) deficient. To demonstrate that these changes explained the reduced exotoxin expression and loss of virulence in TPS3105 we repaired \(agrA\) using allelic exchange to create TPS3105\(r\) (the parental strain TPS3105 now containing an intact \(agrA\)). Quantitative real time RT-PCR for RNAIII demonstrated that TPS3105\(r\) produced 325-fold more RNAIII than TPS3105. Virulence was also restored and TPS3105\(r\) caused greater weight loss, skin lesion area and CFU recovery from lesions compared to the parental strain TPS3105 (\(p < 0.0001\), Figure 5). There was no significant difference between JKD6159 and TPS3105\(r\) in all outcome measures in the mouse skin infection model (Figure 5). These experiments show that intact \(agr\) is essential for the virulence of ST93 CA-MRSA. The \(agrA\) repaired mutant of TPS3105, TPS3105\(r\) expressed significantly greater amounts of PSM\(\alpha3\) (\(p < 0.0001\)) and Hla (\(p = 0.0019\)), consistent with \(agr\) control of these virulence determinants (Figure 6). Thus, despite the genetic divergence of ST93 from other \(S. aureus\) [14], the molecular foundation of virulence for this CA-MRSA clone is similar in this respect to USA300 [9,26,27] and other \(S. aureus\) strains [28,29], where the importance of \(agr\) has been very well established.

The AraC/XylS regulator (AryK) enhanced Hla expression and virulence in ST93 CA-MRSA

The SNP at position 92551 in SAA6159_00084 introduced a premature stop codon and created a pseudogene within SAA6159_00084 in JDK6159, however the gene was intact in TPS3106. The intact version of this gene, which was also
intact in 19 other publically available S. aureus genome sequences we examined, encodes a previously uncharacterized AraC/XylS family regulatory protein. While the virulence attenuation in TPS3106 was likely a direct result of the agr deficiency, we also wanted to determine if the novel regulator mutation in SAA6159_00084 impacted the virulence in ST93 S. aureus.

To test the hypothesis that SAA6159_00084 encoded a regulator of virulence, we repaired the premature stop codon in SAA6159_00084 in JKD6159 using allelic exchange to generate strain JKD6159_AraC. To confirm we had not introduced additional DNA changes during allelic exchange we sequenced the whole genome of JKD6159_AraC and found no additional mutations (35× coverage). JKD6159_AraC encoding an intact copy of SAA6159_00084 demonstrated a modest, but significant increase in virulence as indicated by lesion size (p < 0.0001) and weight loss in the mouse skin infection assay (p = 0.0311, Figure 5), suggesting that this protein is a positive regulator of virulence in CA-MRSA strains. JKD6159_AraC expressed more PSMα3 (p = 0.0325) and Hla (p = 0.0473) than its parental strain JKD6159 that was consistent with an increase mouse skin lesion size (Figure 6). We propose the name aryK for SAA6159_00084 (AraC family-like gene).

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The major pathways where differential transcription occurred were down regulation of genes linked to purine metabolism (purK, purS, purQ, purL, purF), valine, leucine and isoleucine biosynthesis (leuA, leuB, leuC), and oligopeptide transport (opp3C, opp3D, opp3F, opp3A). In contrast, up regulation of genes encoding cation transport systems (mnhB_1, mnhC_1, mnhD_1, mnhE_1, mnhG_1) was found.

Here, we have clearly demonstrated thatagr is the major “on-off” switch for virulence in ST93 CA-MRSA, but we also found that other genetic changes are impacting virulence gene regulation in a clone-specific manner. We speculate that the inactivation of aryK may have been an evolutionary response by ST93 CA-MRSA to modulate or fine-tune the amount of Hla and other factors required for host persistence. There are six AraC/XylS family regulators inS. aureus(SA0097, SA0215, SA0622, SA1337, SA2092, SA2169;S. aureus strain N315 locus tags). Two of these, Rbf (SA0622) and Rsp (SA2169) have been studied and demonstrated in otherS. aureus
strains to regulate biofilm formation and modulate expression of surface-associated proteins [24,25,31]. In contrast, we found that *aryK* increases Hla expression and virulence, acting as a positive regulator of virulence by directly or indirectly upregulating exotoxin expression, without an apparent effect on *agr* expression in stationary phase.

**Conclusions**

In this study, we have obtained insights into the genetic basis for the increased virulence of ST93 by using a combination of comparative and functional genomics. We have demonstrated the key role of Hla and *agr* and shown how an additional novel regulatory gene, *aryK* by a loss-of-function point mutation, is modulating virulence in this clone. Quantification of exotoxin expression in a larger collection of ST93 strains demonstrated that the findings in strain JKD6159 are relevant to the majority of the ST93 population isolated from around Australia as exotoxin expression in JKD6159 is representative of most of the ST93 population. Our study highlights the power of comparative genomics to uncover new regulators of virulence but it also shows the complex nature of these changes even in closely related bacterial populations. Careful strain selection, detailed comparative genomics analyses, and functional genomic studies by creating multiple genetic changes in one strain will be required to gain a full insight into the genetic basis for the emergence and hypervirulence of ST93 CA-MRSA.

**Methods**

**Ethics statement**

This study was performed in accordance with the Australian Prevention of Cruelty to Animals Act 1986 and the Australian code of practice for the care and use of animals for scientific purposes. The protocol was approved by the Animal Ethics Committee of the University of Melbourne (Permit Number: 0911248.2).

**Bacterial strains and culture**

Bacterial strains used in this study are summarized in Table 1 (international clone collection) and Additional file 1 (ST93 strain collection), and include the ST93 reference strains JKD6159, USA300 strain FPR3757 [19], ST30 strain JKD6177, and the HA-MRSA ST239 clone JDK6009 [20], as well as 58 additional ST93 collected from around Australia and previously reported [17]. For all experiments except exotoxin expression bacteria were grown in brain heart infusion broth (BHI, Oxoid). Overnight cultures were diluted 1:100 into fresh media and then incubated at 37°C with shaking (180 rpm) until stationary phase was achieved. For LukF-PV detection, isolates were cultured for 8 hours (OD$_{600}$ ~ 1.8); for Hla detection, isolates were cultured for approximately 3 hours (OD$_{600}$ ~ 1.8); and for PSMα3 detection, isolates were cultured for 24 hours (OD$_{600}$ ~ 2.0). Culture supernatants were harvested by centrifugation and filter sterilized. These were performed in at least triplicate for each *S. aureus* strain.

**Detection of LukF-PV and Hla by western blotting**

Trichloroacetic acid was added to culture supernatants and incubated at 4°C overnight. Precipitates were then harvested by centrifugation, washed with acetonitrile, and solubilized in a SDS and 2-mercaptoethanol containing sample buffer. The proteins were separated on 12% SDS-PAGE. A peptide sequence specific to LukF-PV, HWIGNNYKDENRATHT was synthesized and HRP conjugated polyclonal chicken IgY was raised against this peptide (Genscript). This antibody was used to detect LukF-PV with enhanced chemiluminescence. Images generated from the western blots were quantitated using GS800 Calibrated Densitometer (BioRad) and Image J [32].

Hla was detected using a polyclonal rabbit anti-Hla (Sigma-Aldrich), in buffer containing 20 mM DEPC to inhibit non-specific protein A binding and HRP conjugated sheep anti-rabbit secondary antibody (Millipore) with enhanced chemiluminescence detection [33]. For comparison of JKD6159 versus international clone collection (Table 1), images generated from the Hla western blots were quantitated using GS800 Calibrated Densitometer (BioRad) and Image J [32].

**Detection of PSMα3 expression**

HPLC chromatography was performed on an Agilent Technology 1200 Series system with an analytical Agilent Eclipse XDB-C18 (4.6 mm x 150 mm) column. A water/acetonitrile gradient (0.1% trifluoroacetic acid) from 0 – 100% acetonitrile over 28 min at a flow rate of 1 mL/min.
was used. The total run time was 32 min, and peaks were quantified at a wavelength of 214 nm. The deformylated and formylated form of PSMa3 MEFVAKLFKFDLL GKFLGN was identified in the S. aureus TSB culture supernatants by comparison of their retention times to a commercially synthesized PSMa3 standard (GenScript) and by spiking the samples with the synthesized standards. The identity of the deformylated peptide present in the samples was confirmed by analysing collected fractions by ESI-MS. There was only one peptide present in this fraction; the deformylated form of PSMa3. In contrast, other peptides were observed in the fractions of USA300, JKD6272, TPS3104, TPS3105r, and JKD6159_AraCr containing the N-formylated form of PSMa3. In these cases, the percentage of N-formylated PSMa3 peptide was determined using the total ion count of the major peaks in the ESI-MS and the peak area of the HPLC chromatogram was adjusted accordingly. The concentrations of the deformylated and formylated forms of PSMa3 were determined by comparison of their peak areas to those of the synthesized standards. The standard curves were constructed in the concentration range of 6.2 – 100 μg/ mL and were linear over this range.

**DNA methods, molecular techniques and construction of mutants**

DNA was extracted using the GenElute kit according to the manufacturer’s instructions (Sigma-Aldrich). A lukSF-PV knockout, hla knockout and a repaired agrA of TPS3105 were generated according to the published method [34]. For the knockouts, flanking sequences were amplified and ligated prior to cloning with pKOR1. For allelic replacement to generate TPS3105r, a PCR product of agrA from JKD6159 was cloned with pKOR1. For allelic replacement JKD6159_AraCr, a PCR product of this AraC regulator from TPS3106 was cloned with pKOR1. The deletion of the whole psmA locus in JKD6159, chromosomal restoration of psmA in JKD6159ΔpsmA and the restoration of Hla expression in JKD6159Δhla were conducted using the plMAY protocol described by Monk et al. [35]. Knockout and restoration amplifiers were cloned into plMAY by SLIC [36]. The primers used are listed in Additional file 11. The knockout and restoration clones were confirmed by PCR and Sanger sequencing of the mutated locus. Mutations were further validated with functional assays of activity, which included sheep blood hemolysis, western blot, and/or HPLC.

**Mouse skin infection assay**

Mice were infected with S. aureus as previously described [14]. Briefly, six-week-old female BALB/c mice were infected by intradermal injection with 10⁸ CFU of S. aureus. Mice were assessed and weighed daily for five days. Mice were culled on the 5th day and lesion size measured and CFU recovered from infected tissues by homogenization and colony enumeration on BHI. For each S. aureus strain, at least 10 mice were assessed.

**Genome sequencing**

Genome sequences for three ST93 strains (TPS3104, TPS3105, TPS3106) were obtained from an Illumina GAIIx analyzer using 100 bp paired-end chemistry with a mean fold coverage of 331×. Genome sequencing of the two laboratory-induced mutants JKD6159Δhla (TPS3265) and JKD6159_AraCr (TPS3268) was performed using Ion Torrent sequencing technology.

**Comparative genomics**

A read mapping approach was used to compare the sequences from all isolates used in this study, as previously described [14,37]. Briefly, the reads from all genomes were aligned to the JKD6159 reference using SHRiMP 2.0 [38]. SNPs were identified using Nesoni v0.60 [www.bioinformatics.net.au]. Using the whole genome sequence of JKD6159 as a reference, a global SNP analysis was performed, and allelic variability at any nucleotide position was tallied to generate a global SNP analysis for every genome compared to JKD6159.

**Quantitative RT-PCR for RNAIII expression**

To investigate activity of the agr locus (RNAIII) qRT-PCR was performed for RNAIII as previously described [37]. Briefly, RNA was prepared as previously described with two on-column DNase I digestion steps and cDNA synthesis using SuperScript II reverse transcriptase (Invitrogen). Relative expression was determined as previously described and was normalised against gyrB. Results were obtained from 3 biological replicates each performed in triplicate.

**RNA sequencing**

*Staphylococcus aureus* strains JKD6159 and JKD6159_AraCr were grown to early stationary culture as described above. For RNA protection, 0.5 volumes of RNAlater® RNA stabilization reagent (Qiagen) was added immediately to the liquid culture and allowed to incubate with the bacterial suspension for 15 minutes at room temperature. Cells were pelleted at 5,000 x g for 5 minutes followed by RNA extraction using RNeasy mini kit (Qiagen) and two rounds of DNase I digestion (Qiagen) according to the manufacturer’s instruction. RNA concentration was quantified using Qubit® 2.0 Fluorometer and RNA quality assessed using Agilent 2100 Bioanalyzer. Ten μg of total RNA from the stationary growth phase with RNA integrity number (RIN) greater than 7 was used in RNA-seq. Ribosomal depletion, cDNA library preparation and pair ended sequencing using HiSeq2000 sequencing platform was performed by Beijing Genome Institute.
(Hong Kong, China). RNAseq was performed on two biological samples for each strain.

RNAseq reads were mapped onto the JKD6159 reference genome [23], using SHRIMP 2.2.2 [39]. Alignment to CDS features from each biological replicate of each strain provided counts that were a measure of mRNA levels. Counts were normalized using the trimmed-mean normalization function in edgeR, part of the BioConductor package [40]. A heat map was created based on log$_2$ transformed counts to identify consistent changes in expression profiles between strains. To be included in the heat map, genes were required to have at least 1000 counts, totaled over all samples, where and the standard deviation of the log$_2$ expression levels had to exceed two.

Statistical analysis
Percentage mouse weight change at day 5, viable counts of _S. aureus_ in mouse tissues and skin lesion area of each isolate, Hla, LukF-PV and PSMα3 expression versus JKD6159 were analyzed using an unpaired t test. A similar analysis was used to analyze virulence outcome measures and exotoxin expression between TPS3105 and TPS3105r. (There was no difference in results when Bonferroni analysis was performed). All analyses were performed using Prism 5 for Macintosh v5.0b (GraphPad Software Inc.).

Availability of supporting data
The data sets supporting the results of this article are in the NCBI Sequence Read Archive under study accession SRP004474.2 and the NCBI BioProject Archive under study accession PRJNA217697.

Additional files

Additional file 1: Staphylococcus aureus ST93 strains used in this study.

Additional file 2: Expression of PSMα3 by ST93 strains and USA300. (A) Expression of defomylated PSMα3. (B) Expression of N-formylated PSMα3. Data shown are mean concentration (µg/ml) and SEM.

Additional file 3: Expression of Hla by ST93 strains and USA300. Hla expression measured by quantitative Western blot. Data shown are mean intensity of bands in arbitrary units and SEM.

Additional file 4: Hla Western Blot of JKD6159, JKD6159Δhla and JKD6159Δhla r (A) Western Blot demonstrating that JKD6159Δhla does not express Hla by Western Blot and that complementation of this mutant (JKD6159Δhla r) results in restoration of Hla expression. (B) Arrangement of PCR primers used PCR screen of JKD6159Δhla and JKD6159Δhla r. (C) PCR screen of 25 randomly selected _S. aureus_ colonies obtained from two mice (mouse 4 and mouse 7) post skin infection with JKD6159Δhla r. The PCR primers used flank the region deleted in hla for the mutant and show incomplete penetration of the bacterial population with the repaired version of hla (17/25) with an intact allele for mouse 4 and 21/25 for mouse 7, thereby explaining the inability of the repaired mutant to fully restore the virulence phenotype in this infection model.

Additional file 5: Detection of formylated PSMα3 in JKD6159, JKD6159Δpsma and JKD6159Δpsma r by HPLC of culture filtrates. JKD6159Δpsma did not produce formylated PSMα3. Complementation of this strain resulted in restoration of formylated PSMα3 expression. In all strains δ-toxin expression was maintained.

Additional file 6: LukF-PV Western Blot of JKD6159 and JKD6159ΔlukSF-PV. Western Blot demonstrating that JKD6159ΔlukSF-PV does not express LukF-PV.

Additional file 7: Table of de novo assembly characteristics for _S. aureus_ strains TPS3104, TPS3105 and TPS3106.

Additional file 8: Table of single nucleotide differences between JKD6159 and TPS3105.

Additional file 9: Table of single nucleotide differences between JKD6159 and TPS3106.

Additional file 10: Table of primers used in this study.

Competing interest
No author has any competing interests to declare.

Authors’ contributions
Conceived the project, TPS, BPH, KYLC, JKD; performed the experiments, KYLC, IRM, YHL, JLP, GWC, JS, KLT; analysed the data, KYLC, YHL, TPS, BPH, TS, KLT; wrote the manuscript, KYLC, BPH, TPS. All authors read and approved the final manuscript.

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