Functional analysis of the EsaB component of the
Staphylococcus aureus Type VII secretion system

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Abstract

Type VII secretion systems (T7SS) are found in many bacteria and secrete proteins involved in virulence and bacterial competition. In Staphylococcus aureus the small ubiquitin-like EsaB protein has been previously implicated as having a regulatory role in the production of the EsxC substrate. Here we show that in the S. aureus RN6390 strain, EsaB does not genetically regulate production of any T7 substrates or components, but is indispensable for secretion activity. Consistent with EsaB being an essential component of the T7SS, loss of either EsaB or EssC are associated with upregulation of a common set of iron acquisition genes. However, a further subset of genes were dysregulated only in the absence of EsaB. Quantitative western blotting indicates that EsaB is present at very low levels in cells. Substitution of a highly conserved threonine for alanine or arginine resulted in a loss of EsaB activity and destabilisation of the protein. Taken together our findings show that EsaB is essential for T7SS activity in RN6390.

INTRODUCTION

Protein secretion systems are nanomachines employed by bacteria to transport protein substrates across their cell envelopes. Gram-negative bacteria produce a number of different secretion machineries that export proteins involved in a wide variety of processes including signalling, nutrient scavenging, host interaction and virulence [1]. The Type VII secretion system (T7SS) is found in some Gram-negative and many Gram-positive bacteria, and is particularly common among organisms of the actinobacteria and firmicutes phyla [2]. The T7SS was initially described in the pathogenic mycobacteria Mycobacterium tuberculosis and Mycobacterium bovis, where the ESX-1 T7SS was shown to be essential for virulence, due to the secretion of two major T-cell antigens EsxA (formerly known as ESAT-6) and EsxB (formerly known as CFP-10) [3–5]. EssA and EssB are founding members of the WXG100 protein family that appear to be exclusively linked to T7SSs, and all characterised T7 systems are associated with at least one family member. The presence of a membrane-bound ATPase of the SpoIIE/FtsK family (termed EccC in actinobacteria and EssC in firmicutes) is another hallmark of all T7SSs [6]. In Mycobacteria, three further membrane proteins EccB, EccD and EccE assemble with EccC to form a large 1.5 MDa core complex [7, 8]. This complex further associates with a membrane-bound mycosin serine protease, MycP, that is essential for T7 protein secretion and for stability of the membrane complex [9].

Staphylococcus aureus, an opportunistic pathogen of humans and animals, also elaborates a T7SS that is distantly related to the T7SSs found in mycobacteria [10]. Mutational analysis has indicated that it plays an important role in persistence in mouse models of infection, intra-species competition and potentially iron homeostasis [10–15]. In commonly-studied strains of S. aureus such as Newman, USA300 and RN6390, the secretion system is encoded by the 12 gene ess locus [10, 12, 16]. The first six genes at this locus encode essential components of the secretion machinery, including the WXG100 protein EssA and the SpoIIE/FtsK ATPase EssC (Fig. 1a, b). However, S. aureus and other firmicutes lack homologues of EccB, EccD, EccE and MycP and instead have an apparently unrelated set of membrane-bound secretion components (EssA, EssA and EssB in S. aureus) [12, 17–19]. The sixth component of the S. aureus T7SS is EsaB, which is predicted to be a small cytoplasmic protein of 80 amino acids that is structurally related to ubiquitin [20]. In S. aureus strains Newman and USA300, a
Fig. 1. EsaB is not a transcriptional regulator. (a) The ess locus in S. aureus RN6390. Genes encoding essential secretion components are shaded in grey, secreted proteins in blue and a cytoplasmic antitoxin in yellow. The regions analysed by RT-PCR are indicated. (b) Predicted subcellular locations of Ess-encoded components. cw–cell wall, cm–cytoplasmic membrane. (c) RT-PCR analysis of esxA (region 1) and esxC/B (region 2) from the RN6390 and isogenic esaB and esxA mutant strains. Shading is as for Fig. 1(a) with essential secretion components in grey, secreted proteins in blue and a cytoplasmic antitoxin in yellow. Equivalent amounts of mRNA from each strain were used to generate cDNA. RT: reverse transcriptase. (d) Total mRNA counts of ess genes from RNA-Seq analysis of RN6390 and the esaB mutant strain. RPKM – reads of transcript per kilobase per million of mapped reads.
transposon insertion in esaB does not abolish secretion of T7 substrates but is linked with an increase in RNA transcripts covering the gene encoding the substrate EsxC [11]. By contrast, in-frame deletion of esaB abolished EssA and EssC secretion in strain RN6390 but did not detectably affect production of these substrate proteins [12]. Similarly, inactivation of yukD, which encodes the Bacillus subtilis esaB homologue, also abolished T7 secretion [17, 18].

In this study, we have addressed the role of EsaB in S. aureus T7 secretion using strain RN6390. We show that EsaB does not regulate exxC transcripts or those of other ess-encoded genes. Instead our findings show that EsaB behaves as an essential component of the T7SS. Interestingly, however, RNA-Seq analysis identified a subset of genes from the AirSR regulon that showed altered regulation in the absence of EsaB. This suggests that loss of esaB has additional unexpected effects on S. aureus physiology.

METHODS

Bacterial strains and growth conditions

S. aureus strain RN6390 (NCTC8325 derivative, rbsU, tcaB, cured of φ11, φ12, φ13; [21]) and the isogenic ΔesaB, ΔessC and Δexx (ΔessA – esaG) strains [12] were employed in this study, along with strain Newman [22]. The esaB deletion strain is an in-frame deletion of the gene that maintains the first ten and final three codons of esaB (as there is a 9 codon overlap between the end of essA and the start of esaB). S. aureus strains were cultured in Tryptic Soy Broth (TSB) at 37°C with shaking unless otherwise stated. For calculation of cell numbers we estimated by dilution analysis that 37°C and the start of 10 million c.f.u. for strain. When required, chloramphenicol (Cml, final concentration 100 µg ml⁻¹) was added for plasmid selection. E. coli strain JM110 (Stratagene) was used for cloning purposes and BL21(DE3) [23] for EsaB overproduction and purification. E. coli was grown in Luria-Bertani (LB) medium at 37°C with agitation. When appropriate, ampicillin was used for plasmid selection (final concentration 125 µg ml⁻¹).

Genetic constructs

All plasmids used in this study are listed in Table 1. The esaB gene with its own RBS was PCR amplified from S. aureus RN6390 genomic DNA using primers EsaB-fw and EsaB-rev (Table S1, available with the online version of this article). The 0.3 kb HpaI/EcoRI restriction fragment was cloned into pRAB11 under control of the tetracycline inducible promoter, giving pRAB11-esaB. Clones were selected in E. coli and verified by DNA sequencing. Plasmid pRAB11-esaB-YFP was generated by cloning the 0.3 kb HpaI/EcoRI restriction fragment into pRAB11-YFP [15]. Clones were selected in E. coli and verified by DNA sequencing. Nucleotide variants of esaB were generated by the Quickchange site-directed mutagenesis protocol (Stratagene) using pRAB11-esaB or pRAB11-esaB-YFP as a template and primers listed in Table S1. Modified plasmids were digested using DpnI for at least 1 h at 37°C and transformed into E. coli. Single point mutations were verified by DNA sequencing.

RNA isolation and RT-PCR

For RNA-Seq analysis, three biological repeats of the S. aureus esaB strain was grown aerobically in TSB up to an OD₆₀₀ of 1 at which point mRNA was prepared (in three technical replicates). This experiment was carried out alongside the RN6390 and essC strains [15] and followed identical methodology.

For RT-PCR, the indicated S. aureus strains were grown aerobically in TSB and harvested at an OD₆₀₀ of 1. At this point, the mRNA was extracted using the SV total RNA Isolation Kit (Promega) with some minor modifications. Cell samples were stabilised in 5% phenol/95% ethanol on ice for at least 30 min and then centrifuged at 2770 g for 10 min. Cells were then resuspended in 100 µl of TE buffer containing 500 µg ml⁻¹ lysozyme and 50 µg ml⁻¹ lysozyme and incubated at 37°C for 30 min. Subsequently, the manufacturer’s instructions were followed. Isolated RNA was subjected to a second DNsae treatment using the DNA-free kit (Ambion). RNA was stored at –80°C until use. RT-PCR to probe transcription of genes in the indicated strains was carried out using 500 ng of mRNA as template with the indicated primers (Table S1). PCR products were visualized on 1% agarose gels.

Purification of 6His-EsaB and generation of polyclonal antisera

The EsaB coding sequence (UniProt code ESAB_STAAM) was PCR amplified from a synthetic gene (codon optimised for E. coli K12 (Genscript)) using the primers EsaB-pET1 and EsaB-pET2 (Table S1) and cloned into the Ndel/XhoI site of a modified pET15b vector (Novagen). The plasmid produces an N-terminal His₆-tagged protein with a TEV (tobacco etch virus) protease cleavage site. The protein was expressed and purified as described previously [24], except the tag-free EsaB was not collected in the flow-through of the negative purification but required a 30 mM imidazole elution. The final size exclusion chromatography step used a 24 ml HR 30/100 GL Superdex75 column (GE healthcare), equilibrated with 20 mM Tris pH 7.8, 100 mM NaCl and was calibrated with molecular mass standards (thryoglobulin, 670 kDa; γ-globulin, 158 kDa; serum albumin, 67 kDa; ovalbumin; 44 kDa; myoglobin, 17 kDa; and vitamin B12, 1 kDa). Two mg purified EsaB (retaining a Gly–Ala–Ser–Thr sequence at the N-terminus after the cleavage step) was utilised as antigen to immunise rabbits for polyclonal antibody production in a standard three injections protocol (Seqlab).

Secretion assays, subcellular fractionation and western blotting

The indicated strains were grown overnight in TSB, diluted 1/100 in fresh medium and grown up to mid-log phase, at which point whole cells and supernatant fractions were harvested as described previously [12]. Briefly, cells and supernatant were separated by a 10 min centrifugation step at
2770 g. Cells were washed twice with PBS, adjusted to and OD600 of 1 and digested using 50 µg ml⁻¹ of lysostaphin by incubation at 37 °C for 30 min. Supernatants were filtered using a 0.22 µm filter and TCA-precipitated in the presence of 50 µg ml⁻¹ deoxycholate, as described. For S. aureus subcellular fractionation, cells were grown to mid-log phase with shaking and treated as previously described [12]. Briefly, cells were harvested by centrifugation and resuspended in TSM buffer (50 mM Tris-HCl pH 7.6, 0.5 M sucrose, 10 mM MgCl₂). Lysostaphin was added to a final concentration of 50 µg ml⁻¹ and cells were incubated at 37 °C for 30 min to digest the cell wall. At this point, protoplasts were sedimented to recover the cell wall (supernatant fraction). Protoplasts were disrupted by sonication and the cell wall (supernatant fraction). Protoplasts were sedimented to recover the cell wall (supernatant fraction). Protoplasts were disrupted by sonication and the cell wall (supernatant fraction). Protoplasts were disrupted by sonication and the cell wall (supernatant fraction). Protoplasts were disrupted by sonication and the cell wall (supernatant fraction). Protoplasts were disrupted by sonication and the cell wall (supernatant fraction).

Polyclonal antisera were used at the following dilutions: α-EsxA 1:2500 [12], α-EsxB 1:1000 [15], α-EsxC 1:2000 [12], α-EsaB 1:500, α-TrxA 1:20000 [25] and α-SrtA (Abcam) 1:3000. Anti-GFP antibody was obtained from Roche and used according to manufacturer’s instructions.

RESULTS
EsxB does not regulate the level of esxC transcripts in strain RN6390

A previous study has shown that a transposon insertion in the esxB gene results in an increase in esxC transcripts in the Newman and USA300 strain backgrounds, and a concomitant increase in the EsxC polypeptide, implicating it as a regulator [11]. To investigate whether loss of esxB by in-frame deletion affects the level of esxC mRNA in strain RN6390, we isolated mRNA from the parental strain and the isogenic esxB mutant, prepared cDNA and undertook reverse transcriptase PCR with primers covering either esxA (the first gene at the ess locus, included as a negative control) or esxC (Fig. 1a). It can be seen (Fig. 1c) that the level of transcripts for each of these genes was qualitatively similar in the wild-type and esxB backgrounds.

To examine this quantitatively, we undertook RNA-Seq analysis on RNA prepared from three biological repeats of the RN6390 and esxB strains grown aerobically in TSB to an OD600 of 1. Note that these experiments were performed at the same time as the RN6390 versus essC RNA-Seq analysis described in [15] and used the same RN6390 dataset. Fig. 1(d) shows that the level of esxC transcripts were indistinguishable between the wild-type and esxB strains. Analysis of the transcript levels of the other genes at the ess locus indicates that in general they were also not significantly altered by the loss of esxB although there was a small increase in the level of essB. We conclude that there is no evidence that esxB regulates the level of esxC transcripts in RN6390.

We next examined the entire transcript profile of the esxB mutant to investigate the transcriptional/post-transcriptional response to the loss of this small protein. We found 101 genes de-regulated in the esxB mutant compared to the parental strain (using a cut off of logFC >2 or <-2 and

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Table 1. Plasmids used in this study

| Plasmid          | Relevant genotype or description                              | Source or reference |
|------------------|---------------------------------------------------------------|---------------------|
| pRAB11-esaB-T8A  | pRAB11 producing T8A-substituted EsaB                         | This study          |
| pRAB11-esaB-T8E  | pRAB11 producing T8E-substituted EsaB                         | This study          |
| pRAB11-esaB-T8R  | pRAB11 producing T8R-substituted EsaB                         | This study          |
| pRAB11-esaB-T8H  | pRAB11 producing T8H-substituted EsaB                         | This study          |
| pRAB11-esaB-T8K  | pRAB11 producing T8K-substituted EsaB                         | This study          |
| pRAB11-esaB-T8S  | pRAB11 producing T8S-substituted EsaB                         | This study          |
| pRAB11-esaB-D10A | pRAB11 producing D10A-substituted EsaB                         | This study          |
| pRAB11-esaB-D20A | pRAB11 producing D20A-substituted EsaB                         | This study          |
| pRAB11-esaB-L21A | pRAB11 producing L21A-substituted EsaB                         | This study          |
| pRAB11-esaB-K30A | pRAB11 producing K30A-substituted EsaB                         | This study          |
| pRAB11-esaB-K52A | pRAB11 producing K52A-substituted EsaB                         | This study          |
| pRAB11-esaB-K56A | pRAB11 producing K56A-substituted EsaB                         | This study          |
| pRAB11-esaB-L66A | pRAB11 producing L66A-substituted EsaB                         | This study          |
| pRAB11-esaB-G74A | pRAB11 producing G74A-substituted EsaB                         | This study          |
| pRAB11-esaB-D75A | pRAB11 producing D75A-substituted EsaB                         | This study          |
| pRAB11-esaB-YFP  | pRAB11 producing EsaB-YFP                                      | This study          |
| pRAB11-esaB-T8A-YFP | pRAB11 producing T8A-substituted EsaB-YFP                     | This study          |
| pRAB11-esaB-T8K-YFP | pRAB11 producing T8K-substituted EsaB-YFP                    | This study          |
| pRAB11-esaB-D10A-YFP | pRAB11 producing D10A-substituted EsaB-YFP                  | This study          |
| pRAB11-esaB-K30A-YFP | pRAB11 producing K30A-substituted EsaB-YFP                  | This study          |
| pRAB11-esaB-K52A-YFP | pRAB11 producing K52A-substituted EsaB-YFP                 | This study          |
| pRAB11-esaB-K56A-YFP | pRAB11 producing K56A-substituted EsaB-YFP                 | This study          |
| pRAB11-esaB-D20A-YFP | pRAB11 producing D20A-substituted EsaB-YFP                  | This study          |
| pRAB11-esaB-L21A-YFP | pRAB11 producing L21A-substituted EsaB-YFP                 | This study          |
| pRAB11-esaB-K30A-YFP | pRAB11 producing K30A-substituted EsaB-YFP                 | This study          |
| pRAB11-esaB-K52A-YFP | pRAB11 producing K52A-substituted EsaB-YFP                 | This study          |
| pRAB11-esaB-K56A-YFP | pRAB11 producing K56A-substituted EsaB-YFP                 | This study          |
| pRAB11-esaB-L66A-YFP | pRAB11 producing L66A-substituted EsaB-YFP                  | This study          |
| pRAB11-esaB-G74A-YFP | pRAB11 producing G74A-substituted EsaB-YFP                 | This study          |
| pRAB11-esaB-D75A-YFP | pRAB11 producing D75A-substituted EsaB-YFP                 | This study          |
| pRAB11-esaB-D10A-YFP | pRAB11 producing D10A-substituted EsaB-YFP                  | This study          |
| pRAB11-esaB-D20A-YFP | pRAB11 producing D20A-substituted EsaB-YFP                  | This study          |
| pRAB11-esaB-L21A-YFP | pRAB11 producing L21A-substituted EsaB-YFP                 | This study          |
| pRAB11-esaB-K30A-YFP | pRAB11 producing K30A-substituted EsaB-YFP                 | This study          |
| pRAB11-esaB-K52A-YFP | pRAB11 producing K52A-substituted EsaB-YFP                 | This study          |
| pRAB11-esaB-K56A-YFP | pRAB11 producing K56A-substituted EsaB-YFP                 | This study          |
qvalue <0.05, as applied previously [15]), Fig. 2(a). Of these, 43 were upregulated by the loss of esaB whereas 58 were downregulated when esaB was absent — these genes are listed in Table 2. Interestingly, almost all of the genes that were differentially regulated in the essC mutant [15] were also similarly regulated in the esaB strain (Fig. 3b), although there was a substantive subset of genes that were differentially expressed in the esaB mutant but not the essC strain (Table 2). It can be seen that almost all of the iron acquisition genes, including those for heme acquisition, staphyloferrin synthesis and uptake and ferrichrome import were commonly upregulated by loss of either esaB or essC (Table 2). Furthermore, six of the eight downregulated genes from the essC strain were also down regulated in the esaB strain (note that one of the two genes unaffected in the esaB dataset is essC itself, which appears downregulated in the essC dataset because it has been deleted). The finding that almost the entire subset of genes differentially regulated in the absence of essC is also similarly altered by loss of esaB strongly suggests that EsaB is, like EssC, a component that is essential for activity of the secretion machinery in strain RN6390.

As mentioned above, a subset of transcripts were differentially expressed in the esaB but not the essC strain. These include downregulated genes required for anaerobic nitrate respiration (narGHJ/narK), some secreted proteases (sspa/B/C, aur), capsular polysaccharide synthesis (capG/F/hysA), lactose metabolism (lacB/C/D) and antimicrobial peptide synthesis (epiA/C/D/P). Many of these genes are under control of the essential two component regulatory system AirSR (formerly YhcSR) [26–29]. This observation indicates that EsaB has additional effects on S. aureus physiology. This could be indirect and arising from its role in T7 secretion, for example through altered membrane permeability when EsaB is absent. Alternatively, EsaB may have additional roles in the cell in addition to its requirement for T7 protein secretion.

EsaB is present at low amounts in cells of S. aureus RN6390

To explore the biological role of EsaB in T7 secretion, we overproduced recombinant EsaB with a cleavable His-tag in E. coli. The purified protein eluted from gel filtration as a monomer, in agreement with structural analysis of the B. subtilis EsaB homologue, YukD, which also appears to be monomeric [20]. Polyclonal antisera were raised against purified EsaB and the antibody was affinity purified against the EsaB antigen, before being used to detect the protein in whole cells of S. aureus. Fig. 4(a) shows that although the purified antiserum could clearly recognise purified EsaB, it did not detect a band of the expected size of EsaB in whole cells. We have shown previously that expression of the T7SS genes in RN6390 is upregulated approximately 2–3-fold in the presence of exogenous hemin, and fourfold by hemin in a ΔessC background [15]. However, supplementation of either of these strains with hemin did not result in detectable EsaB in the cellular fraction (Fig. 4b) and it could also not be detected in cells of strain Newman (Fig. 4b). Probing a dilution series of purified EsaB indicated that the antibody was able to cross-react with as little as 25 ng of protein (Fig. 4a), which is equivalent to 1.6 × 10^11 EsaB molecules. Since the antibody was unable to detect EsaB in whole cells of RN6390 from 9.6 × 10^10 colony forming units that were loaded onto the SDS gel, we conclude that there are less than 170 molecules of EsaB per cell.

Since we were unable to detect native EsaB in S. aureus cell extracts, we constructed a series of tagged variants for which commercial antisera were available. To this end we introduced His6, Myc, hemagglutinin (HA) and Strept epitopes onto the N-terminus of EsaB, and His6, Myc, HA, mCherry or FLAG epitopes onto the C-terminus, but in each case were unable to detect the tagged protein (not shown). We also introduced His6 and His6 epitopes into two predicted loop regions internal to the EsaB sequence but again were unable to detect tagged EsaB (not shown). The only tag we introduced that allowed detection of EsaB was a C-terminal yellow fluorescent protein (YFP) tag. Fig. 4(c) shows that basal production of either native (untagged) EsaB or EsaB-YFP from plasmid vector pRAB11 was sufficient to restore secretion of the T7SS extracellular protein EssA and of substrates EssB and EssC to the culture supernatant. Blotting the same cell samples for the presence of the YFP fusion protein (Fig. 4d) showed that it migrated at close to the predicted mass (37 kDa) of the EsaB fusion. There was no evidence for degradation of the fusion protein even after prolonged exposure of the immunoblot (Fig. 4d). We conclude that the YFP-tagged variant of EsaB probably retains functionality.

EsaB-YFP partially localises to the cell membrane

EsaB is predicted to be a soluble cytoplasmic protein [10], and is known to share structural homology with ubiquitin [20]. Interestingly, a domain sharing the same fold is also associated with the actinobacterial T7SS, being found at the cytoplasmic N-terminus of EccD [30], indicating that such proteins may be essential features of all T7SSs. To determine the subcellular location of EsaB-YFP, we blotted secreted and whole cell samples of the esaB mutant strain producing plasmid-encoded EsaB-YFP with the YFP antiserum. Fig. 3(a) shows that EsaB-YFP was associated exclusively with the cellular fraction.

We next fractionated these cells to obtain cytoplasm, cell wall and membrane fractions. Immunoblotting with antisera to control proteins known to localize to the cell membrane (SrtA) and cytoplasm (TrxA) indicated that the fractionation had been largely successful, although some SrtA was found in the cell wall fraction (Fig. 3b). Blotting these same fractions for the presence of EsaB-YFP showed that the protein localised to both the cytoplasm and membrane fractions. Some degradation of the fusion protein was also noted in these experiments which may result from the activation of proteases during fractionation. Indeed, in the membrane fraction it appears that EsaB-YFP migrated as a doublet band of around 37 kDa, indicating some probable truncation of the protein. When unfused YFP was produced
in the wild-type strain it did not localise to the membrane (Fig. 3c), indicating that membrane binding was unlikely to be mediated through the YFP portion of the fusion.

Next we tested whether EsaB-YFP localised to the membrane through interactions with membrane components of the T7SS. To this end we repeated the fractionation in a

Fig. 2. RNA-Seq analysis of differentially regulated genes in the esaB mutant strain. (a) Volcano plot representation of the differentially expressed genes in RN6390 strain compared to the isogenic esaB mutant. The orange and grey spots represent, respectively, genes that are up- or down-regulated in the esaB mutant relative to the parental strain. (b). Overlap between up- and down-regulated genes in the esaB and essC datasets.
Table 2. Genes differentially regulated (>log 2 fold) in the RN6390 esaB deletion mutant, sorted by ascending fold change (FC)

Genes highlighted in grey are also differentially regulated in the essC deletion strain. The column on the right shows the fold change (FC) of the same gene in the essC dataset where NS indicates no statistically significant change in expression level relative to the same gene in the wild-type dataset.

| Locus ID      | Gene name       | FC in esaB mutant | Proposed function                  | FC in essC mutant |
|--------------|-----------------|-------------------|------------------------------------|------------------|
| SAOUHSC_00986 | espC             | −23.7             | Cysteine protease                  |                 |
| SAOUHSC_00988 | espA             | −22.3             | Glutamyl endopeptidase             |                 |
| SAOUHSC_00987 | espB             | −20.8             | Cysteine protease                  |                 |
| SAOUHSC_01573 | −                | −19.0             | Unknown, hypothetical protein       |                 |
| SAOUHSC_01941 | espD             | −18.8             | Serine protease SplB               | −4.3            |
| SAOUHSC_02971 | aub              | −17.1             | Zinc metalloproteinase aureolysin  |                 |
| SAOUHSC_01942 | splA             | −16.4             | Highly specific serine protease    | −5.4            |
| SAOUHSC_02680 | narH             | −15.7             | Nitrate reductase subunit beta     |                 |
| SAOUHSC_01944 | −                | −14.3             | Unknown, hypothetical protein       | −4.5            |
| SAOUHSC_02681 | narG             | −14.3             | Nitrate reductase subunit alpha    |                 |
| SAOUHSC_01121 | hla              | −13.5             | a-hemolysin                        | −4.1            |
| SAOUHSC_02241 | lukF             | −13.0             | Unknown, hypothetical protein       | −3.3            |
| SAOUHSC_02163 | hib              | −12.3             | β-hemolysin                        |                 |
| SAOUHSC_01938 | splD             | −12.2             | Serine protease SplD               | −4.3            |
| SAOUHSC_02679 | narJ             | −12.2             | Nitrate reductase subunit delta    |                 |
| SAOUHSC_02671 | narK             | −11.6             | Putative nitrate transporter        |                 |
| SAOUHSC_02455 | lacA             | −11.0             | Galactose-6-phosphate isomerase    |                 |
| SAOUHSC_01530 | −                | −10.9             | Hypothetical phage protein         |                 |
| SAOUHSC_01542 | −                | −10.9             | Unknown, SNF2 family protein       |                 |
| SAOUHSC_01535 | −                | −10.9             | Phage capsid protein               |                 |
| SAOUHSC_02240 | hib              | −10.5             | Truncated β-hemolysin              |                 |
| SAOUHSC_02423 | lukG             | −10.4             | Leukocidin like toxin              | −4.5            |
| SAOUHSC_02685 | nmrR             | −10.3             | Unknown, hypothetical protein       |                 |
| SAOUHSC_01939 | espC             | −10.3             | Serine protease SplC               | −3.2            |
| SAOUHSC_01937 | −                | −10.3             | Unknown, hypothetical protein       | −2.8            |
| SAOUHSC_02970 | argR             | −8.8              | Arginine repressor family protein  |                 |
| SAOUHSC_00113 | adhE             | −8.6              | Bifunctional acetaldehyde-CoA/alcohol dehydrogenase | |
| SAOUHSC_00051 | plc              | −8.1              | 1-phosphatidylinositol phosphodiesterase | −2.5 |
| SAOUHSC_00898 | argH             | −6.7              | Argininosuccinate lyase             |                 |
| SAOUHSC_02684 | nulD             | −6.6              | Assimilatory nitrite reductase [NAD(P)H] large subunit | |
| SAOUHSC_02187 | hlgC             | −6.5              | γ-hemolysin component C precursor  | −1.8            |
| SAOUHSC_02682 | nufF             | −6.4              | Uroporphyrin-III C-methyltransferase | |
| SAOUHSC_02426 | −                | −6.4              | Unknown, hypothetical protein       |                 |
| SAOUHSC_00401 | −                | −6.3              | Putative exported protein           | −1.6            |
| SAOUHSC_01950 | epbD             | −6.3              | Flavoprotein                       |                 |
| SAOUHSC_01936 | splE             | −6.3              | Serine protease SplE               | −3.3            |
| SAOUHSC_02454 | lacB             | −6.3              | Galactose-6-phosphate isomerase    | −3.4            |
| SAOUHSC_00899 | argG             | −6.2              | Argininosuccinate synthase          |                 |
| SAOUHSC_02108 | ftn              | −6.1              | Ferritin                           |                 |
| SAOUHSC_00368 | −                | −6.1              | Unknown, hypothetical protein       |                 |
| SAOUHSC_00411 | −                | −5.9              | Unknown, hypothetical protein       | −2.2            |
| SAOUHSC_01951 | epbC             | −5.8              | Epidermin biosynthesis protein EpbC |                 |
| SAOUHSC_02683 | nseE             | −5.6              | Assimilatory nitrite reductase [NAD(P)H] small subunit | |
| SAOUHSC_01935 | splF             | −5.3              | Serine protease SplF               | −2.7            |
| SAOUHSC_02452 | lacD             | −5.2              | Tagatose 1,6-diphosphate aldolase  | −2.6            |
| SAOUHSC_01953 | epA              | −5.2              | Gallidermin superfamily EpA protein | |
| SAOUHSC_02541 | nrgK             | −4.9              | Anaerobic ribonucleotide reductase activating protein | |
| SAOUHSC_00717 | sacP             | −4.7              | Putative lipoprotein               | −1.4            |
| SAOUHSC_01990 | gltQ             | −4.6              | Glutamine transport ATP-binding protein | |
| SAOUHSC_02557 | ynt              | −4.5              | Putative urea transporter           |                 |
Table 2. cont.

| Locus ID       | Gene name | FC in esaB mutant | Proposed function                                                      | FC in essC mutant |
|---------------|-----------|-------------------|-----------------------------------------------------------------------|------------------|
| SAOUHSC_01949 | epiP      | −4.4              | Intracellular serine protease                                         | NS               |
| SAOUHSC_00120 | capG      | −4.4              | UDP-N-acetylglucosamine 2-epimerase                                   | NS               |
| SAOUHSC_01952 | bsaB      | −4.4              | Lantibiotic epidermin biosynthesis protein EpiB                       | NS               |
| SAOUHSC_03015 | hisZ      | −4.4              | ATP phosphoribosyltransferase regulatory subunit                       | NS               |
| SAOUHSC_00119 | capF      | −4.4              | Capsular polysaccharide biosynthesis protein CapF                      | NS               |
| SAOUHSC_02463 | hysA      | −4.3              | Hyaluronate lyase                                                      | NS               |
| SAOUHSC_02453 | lacC      | −4.1              | Tagatose-6-phosphate kinase                                           | −2.2             |
| SAOUHSC_00608 | adh1      | −4.1              | Alcohol dehydrogenase                                                 | NS               |

Upregulated genes

| SAOUHSC_02767 | opp-1A    | 4.0               | Peptide ABC transporter substrate-binding protein                      | 2.6              |
| SAOUHSC_02655 |            | −                 | Unknown, hypothetical protein                                          | 6.3              |
| SAOUHSC_01292 |            | −                 | Putative DNA-binding protein                                           | NS               |
| SAOUHSC_00130 | isdI      | 4.4               | Heme-degrading monoxygenase IsdI                                       | 5.7              |
| SAOUHSC_00176 |            | −                 | Extracellular solute-binding protein                                   | NS               |
| SAOUHSC_02435 | sfaA      | 4.5               | Putative transporter                                                  | NS               |
| SAOUHSC_02799 | sarT      | 4.6               | Staphylococcal accessory regulator                                     | NS               |
| SAOUHSC_02432 |            | −                 | Unknown, hypothetical protein                                          | 6.2              |
| SAOUHSC_02425 |            | −                 | Unknown, hypothetical protein                                          | 6.5              |
| SAOUHSC_00652 | fhuA      | 5.1               | Ferrichrome ABC transporter ATP-binding protein FhuA                    | 7.0              |
| SAOUHSC_00071 | sirC      | 5.3               | Involved in staphyloferrin B transport into the cytoplasm              | 4.6              |
| SAOUHSC_00131 |            | −                 | Putative membrane protein                                              | 6.1              |
| SAOUHSC_02821 |            | −                 | Putative membrane protein                                              | NS               |
| SAOUHSC_00070 | sarH1     | −                 | Unknown, hypothetical protein                                          | 3.9              |
| SAOUHSC_02428 | htsB      | 6.3               | Heme transport system permease HtsB                                     | 5.4              |
| SAOUHSC_000974 |          | −                 | Unknown, hypothetical protein                                          | NS               |
| SAOUHSC_01081 | isdA      | 6.5               | Iron-regulated heme-iron binding protein                               | 5.4              |
| SAOUHSC_00072 | strB      | 6.5               | Involved in staphyloferrin B transport into the cytoplasm              | 7.4              |
| SAOUHSC_02554 | fhuD2     | 6.5               | Ferric hydroxamate receptor 1 FhuD2                                     | 6.8              |
| SAOUHSC_00074 |            | −                 | Unknown, hypothetical protein                                          | 3.9              |
| SAOUHSC_00073 |            | −                 | Unknown, hypothetical protein                                          | NS               |
| SAOUHSC_01086 | isdF      | 8.5               | ABC permease IsdF                                                      | 6.1              |
| SAOUHSC_01085 | isdE      | 8.6               | Heme-receptor lipoprotein IsdE                                        | 5.6              |
| SAOUHSC_01089 | isdG      | 8.7               | Heme-degrading monoxygenase IsdG                                       | 4.7              |
| SAOUHSC_01087 |            | −                 | Iron compound ABC transporter permease                                 | 6.3              |
| SAOUHSC_01082 | isdC      | 8.9               | Heme transporter IsdC                                                  | 5.5              |
| SAOUHSC_00748 | strC      | 9.6               | Ferrichrome ABC transporter ATP-binding protein SstC                    | 9.1              |
| SAOUHSC_00545 | strD      | 10.0              | Fibrinogen-binding protein SdrD                                         | NS               |
| SAOUHSC_02246 | fhuD1     | 10.0              | Iron compound ABC transporter FhuD1                                    | 8.0              |
| SAOUHSC_000972 |          | −                 | Unknown, hypothetical protein                                          | NS               |
| SAOUHSC_01088 | strB      | 10.2              | Sortase StrB                                                           | 6.2              |
| SAOUHSC_00747 | sstB      | 10.4              | Ferrichrome ABC transporter permease SstB                               | 9.0              |
| SAOUHSC_00070 | sarH1     | 11.2              | Unknown, hypothetical regulatory-like protein                           | NS               |
| SAOUHSC_02430 | hisA      | 11.2              | Heme transport system lipoprotein HisA                                  | 10.5             |
| SAOUHSC_00074 | strA      | 11.9              | Ferrichrome ABC transporter permease SarA                               | 10.9             |
| SAOUHSC_01084 | isdD      | 13.3              | ATP-hydrolysing and heme-binding protein IsdD                            | 6.2              |
| SAOUHSC_01087 | strA      | 13.6              | Receptor component of staphyloferrin B                                 | 16.3             |
| SAOUHSC_01514 |            | −                 | Unknown, hypothetical protein                                          | NS               |
| SAOUHSC_02232 |            | −                 | Hypothetical phage protein                                             | NS               |
| SAOUHSC_02084 |            | −                 | Phage repressor protein                                                | NS               |
| SAOUHSC_02218 |            | −                 | Unknown, hypothetical protein                                          | NS               |
| SAOUHSC_00069 | spa       | 51.5              | Protein A                                                              | NS               |
Fig. 3. EsaB-YFP localises to the cytoplasm and membrane. (a) EsaB-YFP is not secreted in *S. aureus* strain RN6390. RN6390 harbouring empty pRAB11 and the isogenic ΔesaB strain harbouring empty pRAB11 or pRAB11 encoding EsaB-YFP were cultured in TSB medium until mid-log phase and separated into cellular and supernatant fractions (sn). For each gel, 10 µl of OD_{600} adjusted cells and 15 µl of TCA-precipitated culture supernatant were loaded. Blots were probed with anti-EsxA, anti-TrxA (cytoplasmic control) and anti-GFP antisera. Cell and supernatant samples have been blotted on the same gel but intervening lanes have been spliced out. Subcellular localisation of (b) EsaB-YFP in RN6390 and an isogenic Δesx (ΔesxA-esaG) strain or (c) YFP in RN6390. Cells were grown aerobically in TSB to mid-log phase and fractionated as indicated in the Methods. Equivalent amounts of each fraction was probed with anti-TrxA (cytoplasmic control), anti-SrtA (membrane control), anti-EsxA and anti-GFP antisera.
strain carrying a chromosomal deletion in all twelve genes at the css locus (Fig. 1a). However this did not alter the localisation of EsaB-YFP, which was still detected in both cytoplasm and membrane fractions (Fig. 3b). It is possible that EsaB-YFP localises to the membrane through interaction with additional membrane proteins. Alternatively, it may be that the membrane localisation arises as an artefact of the C-terminal YFP tag, since this tag is known to influence protein behaviour (e.g. [31]).

**Mutagenesis of conserved residues in EsaB**

An alignment of EsaB homologues encoded across firmicutes (Fig. 5a) identifies a number of highly conserved amino acids. Many of these are hydrophilic and fall on one face of the predicted structure of EsaB including T8 (S. aureus numbering) which is highly conserved as either threonine or serine, and the invariant K56. The presence of an invariant lysine is intriguing since there are a number of highly conserved lysine residues on the structurally-related protein ubiquitin, which are used to assemble polyubiquitin chains [32]. To probe potential roles of these conserved residues we mutated each of T8, D10, L21, K30, K52, K56, L66, G74 and D75 to alanine on plasmid-encoded EsaB and assessed whether the variant EsaB proteins were able to restore T7 secretion activity to the esaB deletion strain.

Fig. 5(c) shows that alanine substitutions of each of these conserved residues was tolerated by EsaB with the exception of T8A, which completely abolished EsaB activity. To test whether other side chain substitutions were permissive at T8, we subsequently constructed EsaB T8S, T8E, T8H, T8K and T8R substitutions. As seen in Fig. 5(d), in addition to T8A the T8R substitution also abolished EsaB activity, but the other substitutions resulted in active protein. Finally we attempted to assess whether the T8A and T8R inactivating substitutions altered the subcellular location of EsaB-YFP. However, we introduced these into EsaB-YFP we found that they destabilised the protein as it was almost undetectable in whole cells (Fig. 5e), precluding further analysis. We are therefore unable to determine whether substitution of T8 directly alters EsaB function or has an indirect effect by disrupting folding.

**DISCUSSION**

In this work we have investigated the role of EsaB in Type VII secretion. EsaB proteins are conserved in firmicutes that produce the T7SS and are encoded at the same loci. Previous work had implicated EsaB in the regulation of cssC transcripts [11], although this cannot be a conserved role for EsaB proteins as they are found in all S. aureus strains, including the subset that do not encode cssC [16]. Here we show that EsaB does not regulate cssC in strain RN6390, nor any of the other genes encoded at the css locus. Instead, deletion of esaB is associated with upregulation of genes involved in iron acquisition, mirroring the upregulation of

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Fig. 4. EsaB is present in cells at low amounts. (a) Titration of α-EsaB antibodies. The indicated amounts of purified EsaB, alongside 30 µl of OD₆₀₀₅ adjusted cells were loaded on a SDS-PAGE as indicated and blotted using α-EsaB antibodies. Two exposures of the blot are shown. (b) RN6390 and the isogenic ΔesxC strain (top) or strain Newman (bottom panel) were grown aerobically in TSB medium with or without hemin, as indicated, until an OD₆₀₀ of 2 was reached, at which point cells were harvested as described in Methods. In each case, for detection of EsaB, 25 µl of OD₆₀₀ adjusted cells were loaded and 25 ng of purified EsaB protein was loaded as a positive control. 5 µl of OD₆₀₀ adjusted cells were probed against anti-TrxA antisera as a cytoplasmic control. (c) RN6390 harbouring empty pRAB11 (labelled RN6390), and the isogenic esaB deletion strain harbouring pRAB11 (labelled ΔesaB), or pRAB11 encoding native EsaB or EsaB-YFP was cultured aerobically in TSB medium until an OD₆₀₀ of 2 was reached. Samples were fractionated to give cells and supernatant (sn), and supernatant proteins were precipitated using TCA. For each gel, 10 µl of OD₆₀₀ of 2 was reached. Whole cell samples (20 µl of OD₆₀₀ adjusted cells) were loaded and blots were probed with anti GFP antibodies. Two exposures of the blot are shown.
Fig. 5. Site-directed mutagenesis of conserved residues of EsaB. (a) Sequence alignment of EsaB homologues from: Sau-Staphylococcus aureus; Slu-Staphylococcus lugdunensis; Lmo - Listeria monocytogenes; Lgr - Listeria grayi; Bce - Bacillus cereus; Bam - Bacillus amylo-liquefaciens; Bsu - Bacillus subtilis; Bli - Bacillus licheniformis; Bhc - Bhargavaea cecembensis; Ssi - Solbacillus silvestris; Sgr - Streptococcus orals; Sga - Streptococcus galalolyticus. * indicate conserved residues and † indicates residues forming a potential hydrophobic patch that were mutated in this work. (b) Model of S. aureus EsaB with positions of conserved residues targeted for mutagenesis highlighted. The N- and C-termini are also indicated. (c) and (d) RN6390 harbouring empty pRAB11, and the isogenic esaB deletion strain harbouring pRAB11, or pRAB11 encoding native, the indicated variants of EsaB were cultured aerobically in TSB medium until an OD₆₀₀ of 2 was reached. Samples were fractionated to give cells and supernatant (sn), and supernatant proteins were precipitated using TCA. For each gel, 10 µl of OD₆₀₀ adjusted cells and 15 µl of culture supernatant were loaded. Blots were probed with anti-EsaA, and anti-TrxA (cytoplasmic control) antisera. (e) The ΔesaB strain harbouring pRAB11 encoding EsaB-YFP (WT-YFP) or the T8A or T8R amino acid-
substituted variants were cultured in TSB medium until mid-log phase and separated into cellular and supernatant fractions (sn). For each sample, 10 μl of OD600 adjusted cells and 15 μl of culture supernatant were loaded and blots were probed with anti-EsxA, anti-TrxA or anti-GFP antisera. The cell samples and supernatant samples have been blotted on the same gels but intervening lanes have been spliced out.

iron-acquisition genes seen when the core T7 component, EssC, is absent [15]. This supports the notion that EsaB is an essential component of the secretion machinery in RN6390 that is necessary for activity, and in agreement with this, deletion of esaB prevented export of the T7-dependent extracellular proteins EsxA, EsxB and EssC. This conclusion is also in agreement with related studies in B. subtilis, where the EsaB homologue YukD was shown to be essential for secretion of the WXG100 protein YukE [17, 18].

The precise role of EsaB in T7 secretion is unclear. Structural analysis of B. subtilis YukD shows that it shares a very similar fold to ubiquitin but that it lacks the ability to be conjugated with other proteins [20]. Interestingly, a structurally-related domain is associated with the actinobacterial T7SS, being found at the cytoplasmic N-terminus of the polypeptide EccD membrane component [30], suggesting that EsaB-like domains may be essential features of all T7SSs. Given its small size and the observation that highly conserved residues fall primarily on one face of the protein, we reasoned that EsaB may interact with one or more components of the S. aureus T7SS, potentially regulating activity. Post-translational regulation of the S. aureus T7SS has been suggested because in some growth conditions the secretion machinery is present but there is no or very little substrate secretion [12, 19]. Other protein secretion systems are also post-translationally regulated, for example the flagellar Type III secretion system is regulated through interaction of the FilF component with the second messenger cyclic di-GMP [33], and Type VI secretion systems are regulated by phosphorylation [34]. In this context, EsaB proteins contain a highly conserved threonine (or serine) residue close to their N-termini which we considered as a potential site for phosphorylation. Intriguingly, substitution of EsaB T8 for alanine abolished the function of EsaB, although introduction of either the phospho-mimetic glutamate at this position or a positively charged lysine did not affect EsaB activity.

RNA-Seq analysis of the esaB mutant strain showed that in addition to a common set of genes showing similar regulation in the esaB and essC strains, a further subset of genes were uniquely deregulated in the esaB mutant. Many of the genes in this EsaB-specific subset are part of the AirSR regulon [26–29]. The AirSR two component system responds to oxidation signals via a redox-active [2Fe-2S] cluster in the sensor kinase AirS to regulate diverse sets of genes involved in anaerobic respiration, lactose metabolism and capsule biosynthesis. It is possible that these genes are dysregulated indirectly, for example in the absence of EsaB the T7 machinery may be in a state that causes ion leakage or membrane stress. Alternatively it is possible that EsaB interacts directly with a component of the Air system. In future it will be interesting to further decipher the roles of EsaB in T7 protein secretion and S. aureus physiology.

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Conflicts of interest
The authors declare that there are no conflicts of interest.

References
1. Costa TR, Felisberto-Rodrigues C, Meir A, Prevost MS, Redzej A et al. Secretion systems in Gram-negative bacteria: structural and mechanistic insights. Nat Rev Microbiol 2015;13:343–359.
2. Unnikrishnan M, Constantindou C, Palmer T, Pallen MJ. The Enigmatic Esx proteins: looking beyond mycobacteria. Trends Microbiol 2017;25:192–204.
3. Hsu T, Hingley-Wilson SM, Chen B, Chen M, Dai AZ et al. The primary mechanism of attenuation of bacillus Calmette-Guerin is a loss of secreted lytic function required for invasion of lung interstitial tissue. Proc Natl Acad Sci USA 2003;100:12420–12425.
4. Pym AS, Brodin P, Majlesi L, Brosch R, Demangel C et al. Recombinant BCG exporting ESAT-6 confers enhanced protection against tuberculosis. Nat Med 2003;9:533–539.
5. Stanley SA, Raghavan S, Hwang WW, Cox JS. Acute infection and macrophage subversion by Mycobacterium tuberculosis require a specialized secretion system. Proc Natl Acad Sci USA 2003;100:13001–13006.
6. Pallen MJ. The ESAT-6/WXG100 superfamily – and a new Gram-positive secretion system? Trends Microbiol 2002;10:209–212.
7. Houben EN, Bestebroer J, Ummels R, Wilson L, Piersma SR et al. Composition of the type VII secretion system membrane complex. Mol Microbiol 2012;86:472–484.
8. Beckham KS, Ciccarelli L, Bunduc CM, Mertens HD, Ummels R et al. Structure of the mycobacterial ESX-5 type VII secretion system membrane complex by single-particle analysis. Nat Microbiol 2017;2:17047.
9. van Winden VJ, Ummels R, Piersma SR, Jiménez CR, Korotkov KV et al. Myosins are required for the stabilization of the ESX-1 and ESX-5 type VII secretion membrane complexes. MBio 2016;7:e01471–16.
10. Burts ML, Williams WA, Debord K, Missiakas DM. EsxA and EsxB are secreted by an ESAT-6-like system that is required for the pathogenesis of Staphylococcus aureus infections. Proc Natl Acad Sci USA 2005;102:1169–1174.
11. Burts ML, Dedent AC, Missiakas DM. EsAC substrate for the ESAT-6 secretion pathway and its role in persistent infections of Staphylococcus aureus. Mol Microbiol 2008;69:736–746.
12. Kneuper H, Cao ZP, Towmey KB, Zoltner M, Jäger F et al. Heterogeneity in ess transcriptional organization and variable
contribution of the Ess/Type VII protein secretion system to virulence across closely related Staphylococcus aureus strains. Mol Microbiol 2014;93:928–943.

13. Wang Y, Hu M, Liu Q, Qin J, Dai Y et al. Role of the ESAT-6 secretion system in virulence of the emerging community-associated Staphylococcus aureus lineage ST398. Sci Rep 2016;6:25163.

14. Cao Z, Casabona MG, Kneuper H, Chalmers JD, Palmer T. The type VII secretion system of Staphylococcus aureus secretes a nuclease toxin that targets competitor bacteria. Nat Microbiol 2016;2:16183.

15. Casabona MG, Kneuper H, Alferes de Lima D, Harkins CP, Zoltner M, Fyfe PK, Palmer T, Hunter WN. The Ess/Type VII secretion system of Staphylococcus aureus. BioRxiv 2017.

16. Warne B, Harkins CP, Harris SR, Vatsiou A, Stanley-Wall N et al. The Ess/Type VII secretion system of Staphylococcus aureus shows unexpected genetic diversity. BMC Genomics 2016;17:222.

17. Baptista C, Barreto HC, Sáo-José C. High levels of DegU-P activate an Esat-6-like secretion system in Bacillus subtilis. PLoS One 2013;8:e67840.

18. Huppert LA, Ramsdell TL, Chase MR, Sarracino DA, Fortune SM et al. The ESX system in Bacillus subtilis mediates protein secretion. PLoS One 2014;9:e96267.

19. Jäger F, Zoltner M, Kneuper H, Hunter WN, Palmer T. Membrane interactions and self-association of components of the Ess/Type VII secretion system of Staphylococcus aureus. FEBS Lett 2016;590:349–357.

20. van den Ent F, Löwe J. Crystal structure of the ubiquitin-like protein YukD from Bacillus subtilis. FEBS Lett 2005;579:3837–3841.

21. Novick RP, Ross HF, Projan SJ, Kornblum J, Kreiswirth B. The Esa/Type VII secretion system of Bacillus subtilis mediates protein secretion. Proc Natl Acad Sci USA 1994;91:3967–3975.

22. Baba T, Bae T, Schneewind O, Takeuchi F, Hirama H. Genome sequence of Staphylococcus aureus strain Newman and comparative analysis of staphylococcal genomes: polymorphism and evolution of the major virulence factors. J Bacteriol 2001;183:300–310.

23. Studier FW, Moffatt BA. Use of bacteriophage T7 RNA polymerase to direct selective high-level expression of cloned genes. J Mol Biol 1986;189:113–130.

24. Zoltner M, Fyfe PK, Palmer T, Hunter WN. Characterization of Staphylococcus aureus EssB, an integral membrane component of the Type VII secretion system: atomic resolution crystal structure of the cytoplasmic segment. Biochem J 2013;449:469–477.

25. Miller M, Donat S, Rakette S, Stehle T, Kouwen TR et al. Staphylococcal PknB as the first prokaryotic representative of the proline-directed kinases. PLoS One 2010;5:e9057.

26. Yan M, Yu C, Yang J, Ji Y. The essential two-component system YhcSR is involved in regulation of the nitrate respiratory pathway of Staphylococcus aureus. J Bacteriol 2011;193:1799–1805.

27. Yan M, Hall JW, Yang J, Ji Y. The essential yhcSR two-component signal transduction system directly regulates the iac and opu-CABCD operons of Staphylococcus aureus. PLoS One 2012;7:e50408.

28. Sun F, Ji Q, Jones MB, Deng X, Liang H et al. AirSR, a [2Fe-2S] cluster-containing two-component system, mediates global oxygen sensing and redox signaling in Staphylococcus aureus. J Am Chem Soc 2012;134:305–314.

29. Hall JW, Yang J, Guo H, Ji Y. The AirSR two-component system contributes to Staphylococcus aureus survival in human blood and transcriptionally regulates sspABC operon. Front Microbiol 2015;6:682.

30. Wagner JM, Chan S, Evans TJ, Kahng S, Kim J et al. Structures of EccB1 and EccD1 from the core complex of the mycobacterial ESX-1 type VII secretion system. BMC Struct Biol 2016;16:5.

31. Swulius MT, Jensen GJ. The helical MreB cytoskeleton in Escherichia coli MC1000/pLE7 is an artifact of the N-Terminal yellow fluorescent protein tag. J Bacteriol 2012;194:6382–6386.

32. Li W, Ye Y. Polyubiquitin chains: functions, structures, and mechanisms. Cell Mol Life Sci 2008;65:2397–2406.

33. Trampari E, Stevenson CE, Little RH, Wilhelm T, Lawson DM et al. Bacterial rotary export ATPases are allosterically regulated by the nucleotide second messenger cyclic-di-GMP. J Biol Chem 2015;290:24470–24483.

34. Mougous JD, Gifford CA, Ramsdell TL, Mekalanos JJ. Threonine phosphorylation post-translationally regulates protein secretion in Pseudomonas aeruginosa. Nat Cell Biol 2007;9:797–803.

35. Corrigan RM, Foster TJ. An improved tetracycline-inducible expression vector for Staphylococcus aureus. Plasmid 2009;61:126–129.

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