Characterization of EssB, a protein required for secretion of ESAT-6 like proteins in Staphylococcus aureus

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Abstract

Background: Staphylococcus aureus secretes EsxA and EsxB, two small polypeptides of the WXG100 family of proteins. Genetic analyses have shown that production and secretion of EsxA and EsxB require an intact ESAT-6 Secretion System (ESS), a cluster of genes that is conserved in many Firmicutes and encompasses esxA and esxB. Here, we characterize EssB, one of the proteins encoded by the ESS cluster. EssB is highly conserved in Gram-positive bacteria and belongs to the Cluster of Orthologous Groups of protein COG4499 with no known function.

Results: By generating an internal deletion in essB, we demonstrate that EssB is required for secretion of EsxA. We use a polyclonal antibody to identify EssB and show that the protein fractionates with the plasma membrane of S. aureus. Yet, when produced in Escherichia coli, EssB remains mostly soluble and the purified protein assembles into a highly organized oligomer that can be visualized by electron microscopy. Production of truncated EssB variants in wild-type S. aureus confers a dominant negative phenotype on EsxA secretion.

Conclusions: The data presented here support the notion that EssB may oligomerize and interact with other membrane components to form the WXG100-specific translocon in S. aureus.

Keywords: ESAT-6 secretion, ESS, WXG100, EssB, Type 7 secretion, Staphylococcus aureus

Background

In Gram-positive bacteria, proteins released in the extracellular environment are synthesized as precursor polypeptides with a cleavable N-terminal leader peptide as the sole topogenic signal. Precursors are moved across the plasma membrane by a translocon and signal peptides act on newly translocated precursors to release the mature polypeptide from the membrane [1]. The events leading to protein translocation across the plasma membrane have been genetically dissected using the model organism Escherichia coli. Most precursor proteins travel in an unfolded state through the SecYEG translocon [2-5], pushed by the cytoplasmic ATPase SecA [6]. Precursor proteins bearing a leader peptide with the twin-arginine motif are moved across the plasma membrane by the Tat translocon [7,8]. Recently, it has been observed that some bacteria, in particular Firmicutes and Actinobacteria, can secrete proteins lacking a canonical leader peptide [9]. Many of these proteins share some distinguishing and conserved features that include small size (approximately 100-amino acid residues), a WXG amino acid motif in the middle of the protein [10] and a conserved three-dimensional structure (helix–turn–helix hairpin) [11,12]. Together, these proteins form the WXG100 family of proteins [10]. ESAT-6 and CFP-10 of Mycobacterium tuberculosis are the founding members of the WXG100 family of proteins and are identified with the acronym EsxA and EsxB for ESAT-6 extracellular protein A and B [10]. Bioinformatic and genetic approaches have revealed that the esxA and esxB genes cluster with both conserved and non-conserved genes of unknown function that are required for the stability and secretion of WXG100/Esx proteins into the extracellular milieu [13-16]. These clusters are conserved among several Firmicutes (Figure 1) but not with Mycobacteriaceae who only share EssC-like ATPases.
The name ESX has been used to refer to such gene clusters in Mycobacteriaceae and *M. tuberculosis* for example encodes five ESX clusters (ESX-1 through ESX-5) [17]. In more general term, ESX mediated secretion has been refereed as Type 7 secretion but it was noted that this general designation should not be used for Firmicutes owing to the lack of overall sequence conservation [18]. Clusters bearing *esx* genes have therefore been referred as ESAT-6 Secretion Systems (ESS) in *Staphylococcus aureus* and *Bacillus anthracis* where they have been experimentally examined [16,19-21] and sometimes as WXG100 Secretion Systems (WSS) [22]. It has been proposed that at least three factors, ESAT-6 secretion system genes A, B and C (EssA, EssB and EssC), are important for secretion of WXG100 proteins in *S. aureus* based on the phenotype of transposon insertions in the three corresponding genes [16]. Here, we present genetic and biochemical data that support this hypothesis for EssB. By generating a minimal deletion of *essB* in strain USA300, we observe that EssA remains in the cytoplasm and is no longer secreted into the extracellular milieu. Further, we demonstrate that EssB localizes to the plasma membrane of *S. aureus* and that truncated variants of EssB confer a dominant-negative phenotype on chromosomally encoded EssB (loss of EssA secretion). These results are consistent with the notion that EssB oligomerizes and/or interacts with a larger complex of proteins to mediate translocation of EssA across the plasma membrane of *S. aureus*.

**Results**

**EssB is required for the secretion of EssA by *S. aureus* USA300**

The ESS pathway has previously been examined in *S. aureus* strain Newman, where a transposon insertion in gene NWMN_0222 resulted in a severe loss of EsxA and EsxB production. A definitive function for the *ess* gene product in *S. aureus* Newman could not be revealed, owing to the instability of EsxA and EsxB in this strain. Nevertheless, it was hypothesized that NWMN_0222 may contribute to the secretion of EsxA and EsxB across the membrane. The gene was named EssB for ESAT-6 like secretion system gene B. Further examinations revealed low expression of the ESS cluster in *S. aureus* Newman as compared to the more virulent staphylococcal isolates *S. aureus* USA200, USA300 and USA400 [19,20]. We therefore sought to study the secretion of EssA in strain USA300 and generated an *essB* mutant via allelic replacement. This mutant harbors an internal deletion by fusing the first fifteen and last fifteen codons of the *essB* open reading frame, which otherwise encodes a 444 amino acid polypeptide. In parallel, we produced recombinant EssB in *E. coli* (see below) and used the purified protein to generate a polyclonal rabbit serum.
Cultures of wild-type \textit{S. aureus} USA300 and the isogenic \textit{essB} mutant were grown to mid-log phase and treated with lysostaphin to generate total protein extracts (T, as shown on Figure 2A). Proteins were precipitated with trichloroacetic acid and separated on SDS/PAGE followed by transfer to PVDF membrane for immunoblotting. Blots shown on Figure 2A identify an EssB-immune reactive species in \textit{S. aureus} USA300 that is absent in the extract of the \textit{essB} mutant. As a control, ribosomal protein (L6), \(\alpha\)-hemolysin (Hla) and sortase A (SrtA) were identified in all extracts. The EssB immune species migrated at about 52 kDa on SDS/PAGE. To evaluate the phenotype of the \textit{essB} mutant, staphylococcal cultures were centrifuged to separate bacterial cells (C) from the medium (M), and proteins in both fractions were examined by immunoblotting with EsxA-specific rabbit antibodies (Figure 2B). EsxA was found in bacterial cells and in the extracellular medium of \textit{S. aureus} USA300 cultures. In contrast, EsxA remained in the cytoplasm of \textit{essB} mutant staphylococci (Figure 2B). EsxA immune reactive signals were reduced to non-detectable levels in the extracellular milieu of an \textit{essB} mutant, supporting the notion that EssB is required for the secretion of EsxA. The deletion of the \textit{essB} gene did not affect the localization of the ribosomal protein L6 in the cytoplasm or the secretion of Hla into the extracellular medium (Figure 2B). EsxA secretion was restored to wild-type levels when \textit{essB} was expressed from a plasmid (p\textit{essB}), suggesting that deletion of the \textit{essB} gene does not affect the expression of downstream genes also involved in the ESS pathway [16,19,20].

\section*{Subcellular localization of EssB}

We wondered whether EssB is itself secreted or localizes to a particular subcellular compartment (cytosol/membrane). A culture of \textit{S. aureus} USA300 was centrifuged to separate cells from the extracellular milieu. As expected Hla, but not EssB, was found in the extracellular medium (Figure 2C; lane M). Further fractionation was achieved by subjecting lysed cellular extracts to sedimentation at 100,000 \(\times\) g. As a control for subcellular fractionation, samples were examined by immunoblot for the ribosomal protein L6 (S, soluble) and membrane protein SrtA (I, insoluble). EssB was identified in the membrane sediment along with SrtA (Figure 2C), suggesting that EssB may either be inserted into the lipid bilayer or associated with one or more proteins in the membrane. This finding is in good agreement with a recent report suggesting that YukC the \textit{B. subtilis} homologue of EssB (Figure 1) belongs to the membrane proteome of \textit{B. subtilis} [23].

The TMHMM algorithm (www.cbs.dtu.dk/services/TMHMM-2.0) was used to perform sequence-based prediction of EssB, which identified a string of hydrophobic residues amino acids 229–251 (W229VAIGMTTLSVL-LIAFLAFLYFS251) at the center of the EssB polypeptide. Hereafter we refer to the segment of hydrophobic amino

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure2.png}
\caption{Identification and characterization of EssB. (A) \textit{S. aureus} USA300 (WT) or isogenic mutant \textit{essB} were examined for production (T: total culture extracts) and subcellular localization of EssB (C: cell extracts followed by 100,000 \(\times\) g sedimentation and separation of soluble, S and insoluble I proteins; M: medium). Proteins in each fraction were precipitated with trichloroacetic acid, separated by SDS-PAGE and detected by immunoblotting with specific antibodies [\(\alpha\)-EssB, as well as \(\alpha\)-L6, \(\alpha\)-Hla, \(\alpha\)-SrtA, as cytoplasmic, secreted and membrane protein controls, respectively]. (B) Plasmid complementation analysis of bacterial cultures separated between cells (C) and medium (M). \textit{S. aureus} USA300 (WT) or \textit{essB} mutants harboring or not a complementing plasmid (p\textit{essB}) were examined for their ability to secrete EsxA in the culture medium. Samples were analyzed as in panel A.}
\end{figure}
acids within EssB as the Putative Trans Membrane Domain (PTMD).

Deleting essB affects the production of several ESS factors

Recently, we reported that the last gene of the ESS cluster, esaD, is required for the effective secretion of EsxA (Figure 1) [20]. We therefore wondered whether the EsxA secretion phenotype of the essB mutant could be explained by the possible loss of expression of other EsaD factors. To examine this possibility, extracts of bacterial cultures (medium and lysed cells) derived from wild-type or the essB mutant carrying either a plasmid control without insert (vector) or the complementing plasmid (pessB), were subjected to immunoblot analysis using antibodies against EsaD as well as the control protein SrtA (Figure 3A). Interestingly, EsaD appeared to accumulate in the essB mutant. Intrigued by this finding, we performed a similar analysis using antibodies against EsaB, a small cytoplasmic protein that modulates the ESS pathway by an unknown mechanism [19]. EsaB is conserved in the minimal ESS cluster of B. subtilis where it is designated YukD (Figure 1). We observed that deletion of essB also led to the accumulation of EsaB (Figure 3A). These observations were quantified by performing each experiment in triplicate and comparing the average abundance of proteins in wild-type and essB mutant strains. EsaD and EsaB were found to accumulate with 2.5-fold and 5-fold increase over wild type, respectively (Figure 3B). Expression of wild-type essB from the complementing plasmid rescued this phenotype, albeit that only partial complementation was achieved. Perhaps, the physiological ratio between EssB and EsaB could not be achieved upon overexpression of essB using a plasmid. Taken together, these observations suggest that EssB is a critical component of the ESS pathway required for secretion of EsxA and proper accumulation of EsaB and EsaD.

Recombinant EssB is soluble and prone to multimerization

EssB is a 444 amino acid protein with relative molar mass $M_r = 52023.94$ (Figure 4A). Its production could be achieved to high yield in E. coli BL21(DE3) harboring pET15b encoding essB. In order to purify the protein, cells were lysed in a French pressure cell and lysates were subjected to ultracentrifugation at 100,000 × g for 60 min. To our surprise most EssB remained in the supernatant (>75%). Assuming that amino acids 229–251 represent a hydrophobic buried segment, the primary sequence of EssB can be roughly divided in two soluble N-terminal and C-terminal domains (Figure 4A). We generated five recombinant variants encompassing the predicted soluble N- or C-terminal domain with or without the PTMD as well as a variant lacking PTMD (Figure 4A). The variants were named EssB$^N$, EssB$^C$, EssB$^{NM}$, EssB$^{MC}$, EssB$^{ΔM}$, respectively. Similar to full length EssB, over 75% of the overproduced proteins could be recovered from the supernatant of E. coli.
lysates subjected to ultracentrifugation (100,000 × g for 60 min) with the exception of EssBΔM that was poorly expressed. Full length EssB along with all variants were purified to homogeneity using affinity chromatography and the affinity tags were removed. (C) Size exclusion chromatography of full length EssB and truncated variants shown in panel B. Proteins (~100 μg) were loaded onto a Superdex™ 75 10/300 GL and fractions (0.5 ml) were collected and analyzed by SDS-PAGE. Proteins in the gel were visualized by Coomassie staining. Masses of protein standards used for calibration are shown above the gels (158, 75, 43, 17 kDa) and correspond to the exclusion volumes of Aldolase, Conalbumin, Ovalbumin and Myoglobin, respectively. (D-E) TEM of purified recombinant EssB (D) and EssBΔM (E). The proteins were allowed to bind to glow discharged grids and were negatively stained using 2% uranyl acetate. This analysis reveals a rod-like structure for EssB and more spherical, aggregated-like structure for EssBΔM. Scale bar = 20 nm.

Figure 4 Purification and characterization of recombinant EssB and truncated variants. (A) Diagrammatic representation of full length EssB and truncated variants produced in E. coli. Numbers indicate amino acid positions in the primary sequence and the grey box labeled PTMD depicts the hydrophobic sequence. (B) Coomassie gel of purified recombinant proteins as shown in panel A. Proteins were purified from E. coli by affinity chromatography and affinity tags were removed. (C) Size exclusion chromatography of full length EssB and truncated variants shown in panel B. Proteins (~100 μg) were loaded onto a Superdex™ 75 10/300 GL and fractions (0.5 ml) were collected and analyzed by SDS-PAGE. Proteins in the gel were visualized by Coomassie staining. Masses of protein standards used for calibration are shown above the gels (158, 75, 43, 17 kDa) and correspond to the exclusion volumes of Aldolase, Conalbumin, Ovalbumin and Myoglobin, respectively. (D-E) TEM of purified recombinant EssB (D) and EssBΔM (E). The proteins were allowed to bind to glow discharged grids and were negatively stained using 2% uranyl acetate. This analysis reveals a rod-like structure for EssB and more spherical, aggregated-like structure for EssBΔM. Scale bar = 20 nm.

lyates subjected to ultracentrifugation (100,000 × g for 60 min) with the exception of EssBΔM that was poorly expressed. Full length EssB along with all variants were purified to homogeneity using affinity chromatography and the affinity tags were removed by thrombin digestion. The purity of the polypeptides was evaluated on Coomassie-stained SDS/PAGE (Figure 4B). Next, these polypeptides were subjected to gel filtration onto Sephacryl S-200 column and aliquots of eluted fractions were evaluated once more on Coomassie-stained SDS/PAGE (Figure 4C). When subjected to gel filtration, EssB eluted as a homogenous peak with $M_r \approx 158,000$ (Figure 4C). The elution profile did not change when the protein concentration was increased or decreased by a factor of 10 and EssB protein did not scatter UV light suggesting that the polypeptide remained soluble (not shown). Variants that lacked PTMD, EssBΔM and EssBΔC, eluted with $M_r \approx 22-25,000$, close to their calculated masses (Figure 4C). In contrast, variants that retained PTMD, EssBΔNM and EssBΔMC, eluted with $M_r > 158,000$ following size exclusion chromatography (a somewhat higher mass than the full length protein). Removal of PTMD caused EssBΔM to elute with a $M_r$ of $\approx 47,000$ suggesting that quite like EssBΔN and EssBΔC, this variant did not multimerize (Figure 4C).

Visualization of purified EssB protein by transmission electron microscopy suggested that the sample is homogenous. Small dense structures could be seen throughout the field and at larger magnification they revealed a clear rod-shaped organization of the molecule (Figure 4D). A similar analysis was performed for affinity purified EssBΔM. Transmission electron micrography revealed that overall the protein preparation was homogeneous (not shown), however the rod-shaped structure of EssB is lost in this variant (Figure 4E). Together, these results suggest that the PTMD segment is required for the multimerization of EssB and that the rod-shaped structure may be an energetically favorable conformation in the cytoplasm of E. coli. Interestingly, the structure for a so-called “cytoplasmic component of EssB” has been deposited in the databank and made publicly available (http://www.ncbi.nlm.nih.gov/Structure/mmdb/...
Truncated EssB variants display a dominant negative phenotype in *S. aureus*

We wondered whether truncated EssB variants may trigger misassembly of the ESS secretion machinery and interfere with the secretion of EsxA in *S. aureus*. To test this, the EssB variants illustrated in Figure 4A were cloned into the expression plasmid pWWW412 and transformed into *S. aureus* USA300 wild-type and essB mutant strains. First, complementation of Ess function was assessed in the essB mutant, using plasmids carrying either no insert or wild-type essB controls or essB variants encoding EssB<sup>N</sup>, EssB<sup>C</sup>, EssB<sup>NM</sup>, EssB<sup>MC</sup>, EssB<sup>ΔM</sup>, respectively (Figure 5A). Cell extracts were fractionated to reveal synthesis and subcellular localization of full length or truncated EssB proteins following sedimentation of lysed cells at 100,000 × g (Figure 5A). As a control, sortase A (SrtA) was found in the sediment (I, insoluble fraction) of ultracentrifugation samples. EssB protein expressed from pessB in the essB mutant strain also sedimented during ultracentrifugation (Figure 5A; right two lanes) similar to endogenous EssB from *S. aureus* USA300 (Figure 2A). An additional immune reactive species was observed when EssB was overproduced from the plasmid (Figure 5A, white asterisk). Variants carrying the PTMD sequence, EssB<sup>NM</sup> and EssB<sup>MC</sup>, sedimented during ultracentrifugation, whereas EssB<sup>ΔM</sup>, the variant that lacks the PTMD sequence, did not. Two proteins assumed aberrant behavior. The EssB<sup>N</sup> protein was either poorly produced or very unstable in *S. aureus* essB mutant (Figure 5A; white arrow). EssB<sup>C</sup> partitioned into both the soluble and the insoluble fractions. Perhaps, this domain interacts weakly with components of the secretion machine embedded in the membrane. Of note, only the plasmid encoding full-length EssB restored EsxA secretion into the extracellular medium of essB mutant cultures (M); all other plasmids failed to complement essB for EsxA secretion (Figure 5B). As expected, the control ribosomal protein L6 was found in cell lysates (C) (Figure 5B).
When transformed into wild-type \textit{S. aureus} USA300, plasmid produced EssB and variants fractioned as before following \(100,000 \times g\) ultracentrifugation (Figure 5C). Briefly, EssB, EssB\textsuperscript{NM} and EssB\textsuperscript{MC} were found in the sediment, EssB\textsuperscript{AM} remained soluble and EssB\textsuperscript{AT} fractionated equally in the soluble and insoluble compartments (Figure 5C). Expression of EssB\textsuperscript{NM} led to some degradation of EssB (Figure 5C, black asterisk). As before, very little EssB\textsuperscript{NI} immunoreactive material could be detected in \textit{S. aureus} USA300 cells (Figure 5C, white arrow). Interestingly, its production caused a reduction of wild-type EssB (Figure 5C, blue arrow). EssB was also unstable in the merodiploid strain expressing EssB\textsuperscript{MC} (Figure 5C; purple arrow). Not surprisingly, destabilization of EssB by either EssB\textsuperscript{NI} or EssB\textsuperscript{MC} led to altered expression and secretion of EssA (Figure 5D). Sedimentable variants encompassing the PTMD, EssB\textsuperscript{NM} and EssB\textsuperscript{MC}, caused a dominant-negative phenotype on the activity of wild-type EssB and as a result expression or secretion of EssA were altered. On the contrary, EssB\textsuperscript{AM} lacking PTMD remained soluble and did not interfere with EssB function. Taken together, these data suggest that EssB variants that sediment with staphylococcal membranes interfere with the stability or function of endogenous EssB and as a consequence EssA production and secretion are also affected. Thus, EssB is part of the secretion machine and its multimerization and possible association with other Ess components enables the secretion of EssA.

**Discussion**

Secreted proteins are generally tagged with topogenic sequences for recognition by a specific secretion machine and transport across the plasma membrane. Over a third of all proteins synthesized by a bacterial cell carry leader peptides, the topogenic signal for recognition by a third of all proteins synthesized by a bacterial cell carry chime and transport across the plasma membrane. Over sequences for recognition by a specific secretion machinery. Secreted proteins are generally tagged with topogenic sequences that enable the secretion of EsxA. This enables the secretion of EsxA.

By raising a polyclonal antibody against EssB, we find that the protein sediments with \textit{S. aureus} membranes in a manner similar to SrtA, a well-characterized membrane embedded protein [29]. Residues 229–251 roughly define a hydrophobic sequence reminiscent of a transmembrane spanning segment (PTMD). Interestingly, recombinant EssB behaves as a soluble oligomer in \textit{E. coli} with a rod-shaped like structure and the PTMD sequence appears to be necessary and sufficient for this oligomerization process. Obviously, this conformation may simply represent an energetically favorable state for an otherwise membrane-spanning. Nonetheless, recombinant EssB\textsuperscript{NM} and EssB\textsuperscript{MC} are more prone to multimerization than intact EssB suggesting that the full-length sequence limits or regulates the oligomerization of the protein. Protein translocators of other secretion systems such as the Tat or holin pathways undergo regulated multimerization to facilitate pore function in the membrane [30,31]. In \textit{S. aureus}, the presence of the PTMD targets EssB\textsuperscript{NM} and EssB\textsuperscript{MC} to the membrane. This targeting appears to affect the function of endogenous EssB in wild-type staphylococci. On the contrary, EssB\textsuperscript{AM} (lacking PTMD) is soluble. It is unable to complement the \textit{essB} mutant and it displays no dominance over wild-type for EssA secretion. As such, none of the truncated EssB variant could complement wild-type EssB for secretion. Further studies are needed to determine whether the PTMD sequence serves as an autonomous membrane-spanning domain or whether it provides a mean to associate with another integral membrane protein encoded within the ESS cluster.

Deletion of \textit{essB} in strain USA300 leads to loss of EssA secretion and EssA remains in the cell. Because overproduction of EssB is not toxic in \textit{E. coli}, we do not believe that this protein alone is capable of forming a pore for the passage of secreted substrates. Interestingly, two other proteins EsaB and EsaD also accumulate in the \textit{essB} mutant. While the exact role of EsaB
is still unknown, it does not appear to be a secreted substrate [19], and thus the reason for this increase is unclear but it points to additional biochemical interactions within proteins of the ESS cluster. Recent evidence suggests that EsaD is a membrane protein also required for EsxA secretion [20]. Perhaps EssB interacts physically with EsaD to either complete or regulate formation of the translocon. Future studies are needed to address this possibility and determine whether EssB is an integral or peripheral element of the ESS translocon.

Conclusions
The ESS pathway is an alternate and conserved secretion system of several Gram-positive bacteria. Here, we show that EssB is found in the membrane of *S. aureus* and deletion of the corresponding gene abrogates secretion of EsxA. We show that a hydrophobic segment in the middle of the protein referred as PTMD is required for targeting to the plasma membrane. We observe that recombinant EssB harboring PTMD folds into an oligomeric rod-shaped structure that allows the protein to remain soluble in *E. coli*. Interestingly, truncated EssB variants harboring an intact PTMD display a dominant negative phenotype over wild type EssB for secretion of EsxA. The data indicate that EssB is an essential component of the ESS translocon and likely interacts with itself and other machine components. Together, this study provides the first genetic and biochemical characterization of the ESS translocon in *S. aureus*.

### Table 1 Oligonucleotides used in this study

| Name               | Nucleotide sequence                                                                 | Usage                                                                 |
|--------------------|-------------------------------------------------------------------------------------|----------------------------------------------------------------------|
| essB-attB1-F       | GGGGACAAAGTTTGATACAAAAAGCAAGGCTCTACCTTGAATGTTGATTTTAATCTATAG                    | Cloning of the essB deletion mutant in pKOR1 for allelic replacement |
| essB-15codons-R    | AAGATCTCAACATCTTTGACTTTTGATTAG                                                     | Same as above                                                        |
| essB-15codons-F    | AAAGATCTCAAAAGATAAAGAAACGCCAAAG                                                   | Same as above                                                        |
| essB-attB2-R       | GGGGACAACTTTGATAAAGAAAGCTGGTGCAACCCGGATATGTAATTTC                                | Same as above                                                        |
| essB-XhoI-F        | AAACCTGAGATTGTTAAAAATCATAACCTATCTTAATCTATAG                                     | Gene expression in *E. coli* or *S. aureus* using pET15b or pWWW412, respectively |
| essB-BamHI-R       | AAGAGATCCCTATTGTTTTTTCTTCTAGCTTTGCTAGGTTT                                      | Same as above                                                        |
| essB(1-223)-BamHI-R| AAGAGATCCCTACCTTTCTTTGCTGATATGATATGATA                                          | Same as above                                                        |
| essB(23-444)-NdeI-F| AAACATATGAAAAAGACGGCATTGAAGGCAAGGC                                                | Same as above                                                        |
| essB(1-252)-BamHI-R| AAGAGATCCCTATTGTTTTTTCTTCTAGCTTTGCTAGGTTT                                      | Same as above                                                        |
| essB(220-444)-NdeI-F| AAACATATGAAAAAGACGGCATTGAAGGCAAGGC                                                | Same as above                                                        |
| essB(A224-252)-EcoRI-R | AAGGAATCTAGAAGCATAAGTAGAGGGGATTGAAAG                                            | Same as above                                                        |
| essB(A224-252)-EcoRI-F   | AAGGAATCTAGAAGCATAAGTAGAGGGGATTGAAAG                                            | Same as above                                                        |
| essB-BamHI-F       | AAGAGATCCCTATTGTTTTTTCTTCTAGCTTTGCTAGGTTT                                      | Production of GST hybrids using pGEX-2TK                              |
| essB-EcoRI-R       | AAGGAATCTATTGTTTTCTTCTAGCTTTAGG                                                  | Same as above                                                        |
| essB(1-223)-EcoRI-R | AAGGAATCTATTGTTTTCTTCTAGCTTTAGG                                                  | Same as above                                                        |
| essB(253-444)-BamHI-F | AAGGAATCTAAGATAGAAATGAGGGCATTGAAAG                                             | Same as above                                                        |
deletion of the Putative Trans Membrane Domain (PTMD), two DNA fragments were amplified with two sets of primers prior to ligation in pWWW412. The pET15b (Novagen) and pGEX-2T (GE Healthcare) vectors were used for expression of recombinant essB and truncated variants in E. coli. The DNA sequences of the full-length gene and variants were amplified by PCR using primers listed in Table 1. Vector pET15b was used for production of recombinant EssB, EssB<sup>NM</sup>, EssB<sup>MC</sup>, EssB<sup>AM</sup>, and pGEX-2T for production of recombinant EssB<sup>N</sup> and EssB<sup>C</sup>. All clones were validated by nucleotide sequencing performed by the DNA Sequencing Facility of the Cancer Research Center at the University of Chicago. All plasmids and strains are listed in Table 2.

**Culture and bacterial fractionation, and western blot experiments**

Bacterial strains were grown overnight from isolated colonies in TS broth supplemented with 0.2% serum at 37°C with shaking. Cultures were diluted 1:100 in fresh broth and allowed to shake at 37°C until they reached an absorbance of 1 at 600 nm (A<sub>600nm</sub>) corresponding to exponentially growing bacteria. For whole culture lysates (samples labeled T, for total culture extracts as shown in Figures 2A and 3), cultures (6 ml) were incubated in the presence of lysostaphin (100 μg/ml) for 30 min at 37°C. To separate proteins in the culture medium (M) from those in the bacterial cell (C), cultures (6 ml) were centrifuged (10,000 × g for 10 min) and the supernatant was transferred to a new tube prior to lysostaphin treatment of intact cells. For subcellular localization of EssB (Figures 1A and 5 top panel), cultures were centrifuged to separate medium and cells. Staphylococci were washed, and peptidoglycan digested with lysostaphin. Staphylococcal extracts were subjected to ultracentrifugation at 100,000 × g for 40 min at 4°C. The supernatant, containing soluble proteins (S), was transferred to a new tube. The sediment containing insoluble membrane proteins (I), was suspended in 6 ml PBS buffer. Proteins in all samples were precipitated with 10% trichloroacetic acid on ice for 30 min. Precipitates were sedimented by centrifugation at 15,000 × g, washed, dried and solubilized in 100 μl of 0.5 M Tris–HCl (pH 8.0)/4% SDS and heated at 90°C for 10 min. Proteins were separated on SDS/PAGE and transferred to poly(vinylidene difluoride) membrane for immunoblot analysis with appropriate polyclonal antibodies. Immunoreactive signals were revealed by using a secondary antibody coupled to IRDye<sup>680</sup>. Quantification of western blots was conducted using a Li-Cor Biosciences Odyssey imager.

**Table 2 Strains and plasmids used in this study**

| Strains          | Description                                                                 | Reference          |
|------------------|-----------------------------------------------------------------------------|--------------------|
| pWWW412         | pOS1 derivative carrying the constitutive hprK promoter, Cm<sup>R</sup>     | [33]               |
| pKOR1           | temperature sensitive plasmid for allelic replacement, Cm<sup>R</sup>      | [32]               |
| pET15b          | Vector carrying an N-terminal hexa-histidine repeat followed by a thrombin cleavage site and T7 transcription start; Amp<sup>R</sup> | Novagen            |
| pGEX-2TK        | Vector carrying GST followed by thrombin cleavage and phosphorylation sites, and a taq promoter; Amp<sup>R</sup> | GE Healthcare      |
| pESSB<sup>B</sup> | pWWW412 expressing wild-type essB codons 1-444                            | This study         |
| pESSB<sup>B,M</sup> | pWWW412 expressing codons 1–223 of essB                                  | This study         |
| pESSB<sup>B</sup> | pWWW412 expressing codons 1–252 of essB                                  | This study         |
| pESSB<sup>M</sup> | pWWW412 expressing codons 253–444 of essB                                | This study         |
| pESSB<sup>B,M</sup> | pWWW412 expressing essB lacking codons 224-252                           | This study         |
| phi1-essB<sup>B</sup> | pET15b expressing histidine tagged essB codons 1-444                    | This study         |
| phi1-essB<sup>B,M</sup> | pET15b expressing histidine tagged essB codons 1–252                    | This study         |
| phi1-essB<sup>M</sup> | pET15b expressing histidine tagged essB codons 220–444                 | This study         |
| phi1-essB<sup>B,M</sup> | pET15b expressing histidine tagged essB lacking codons 224-252          | This study         |
| pGEX-2TK         | pGEX-2TK expressing GST fused to codons 1–223 of essB                    | This study         |
| pGST-essB<sup>B</sup> | pGEX-2TK expressing GST fused to codons 253–444 of essB                  | This study         |
Briefly, cells were grown to the same optical density. All strains reached similar density in the same time period suggesting that either deletion or cis-expression of genes did not affect growth of bacteria. Signal intensity of immune reactive signals for EssA, EssB, EssBΔC and EssD was compared to that obtained for WT, WT/vector, essB/pessB or WT/pessB sample extracts for Figures 2, 3, 5 A, B, C and D, respectively. Immune reactive signals (as shown in Figure 3) were averaged in three independent experiments and the data was analyzed in pairwise comparisons between WT/vector and variant strains with the unpaired two-tailed Student’s t-test and found to be statistically significant.

**Protein and polyclonal antibody purification**

Briefly, recombinant EssB, EssBΔN, EssBΔC, EssBΔM tagged with N-terminal hexa-histidine were purified using Ni-NTA Agarose (Qiagen) following manufacturer’s recommendations. Recombinant EssBΔN and EssBΔC fused to glutathione S-transferase were purified by using sepharose-immobilized glutathione (Glutathione Sepharose, GE Healthcare) following manufacturer’s recommendations. Bound proteins were incubated in 50 mM Tris–HCl buffer (pH 7.5) containing 300 mM NaCl (buffer A) with thrombin (10 U/mg, GE Healthcare) at 4°C for 12 h to cleave the hexa-histidine and glutathione S-transferase moieties, respectively. Released proteins were dialyzed in buffer B (50 mM Tris–HCl [pH 8.0] containing 150 mM NaCl and 1 mM DTT) and stored at 4°C for use within the next 48 h. A 100-μl volume of each recombinant protein (~100 μg) was loaded onto a Superdex™ 75 10/300 GL (GE Healthcare) in buffer B at 4°C. The chromatography was performed at a flow rate of 0.5 ml/min, and fractions of 0.5 ml were collected and analyzed by SDS-PAGE. The gel filtration was compared to that obtained for WT, WT/vector, ΔΔm, ΔΔc, ΔΔn and variant strains.

Transmission electron microscopy (TEM) and image processing

Purified recombinant proteins EssB and EssBΔM were prepared as described above, dialyzed in Buffer B (without DTT) and diluted to approximately 10 to 50 μg/ml. Proteins were bound to glow discharged, carbon coated (Edwards Auto 306 Evaporator) copper grids (400 mesh), washed, and subsequently negatively stained using 2% uranyl acetate (Electron Microscopy Services). Images were recorded using a Tecnai F30 (Philips/FEI) transmission electron microscope (Field emission gun, 300-kV accelerating voltage, with a magnification of 49,000 to 75,000×) and a high performance CCD camera with a 4K × 4K resolution. Images were captured using Gatan DigitalMicrograph software and processed using Adobe Photoshop (Adobe, San Jose, CA, USA). Images of single protein were selected manually.

**Abbreviations**

Ess: ESAT-6 secretion system; TSS: Type 7 secretion system; Sec: Secretion; LB: Luria Bertani; TS: Tryptic Soy; SDS-PAGE: Sodium dodecyl sulphate polyacrylamide gel electrophoresis; TEM: Transmission electron microscopy; TM: Transmembrane domain; T: Total culture; M: Medium; C: Cell; S: Soluble proteins; I: Insoluble membrane proteins.

**Competing interests**

The authors declare no competing interest.

**Authors’ contributions**

YHC conducted most of the experiments in the study and wrote a preliminary draft. MA generated some of the *S. aureus* reagents. APAH performed the transmission electron microscopy. DM defined the concept of the study and wrote the manuscript. All authors have read and approved the final manuscript.

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