Staphyloferrin B (SB) is an iron-chelating siderophore produced by *Staphylococcus aureus* in invasive infections. Proteins for SB biosynthesis and export are encoded by the *sbnABCDEF-GHI* gene cluster, in which SbnI, a member of the ParB/Srx superfamily, acts as a heme-dependent transcriptional regulator of the *sbn* locus. However, no structural or functional information about SbnI is available. Here, a crystal structure of SbnI revealed striking structural similarity to an ADP-dependent free serine kinase, SerK, from the archaea *Thermococcus kodakaren-* sis. We found that features of the active sites are conserved, and biochemical assays and 31P NMR and HPLC analyses indicated that SbnI is also a free serine kinase but uses ATP rather than ADP as phosphate donor to generate the SB precursor O-phospho-l-serine (OPS). SbnI consists of two domains, and elevated B-factors in domain II were consistent with the open–close reaction mechanism previously reported for SerK. Mutagenesis of Glu20 and Asp28 in SbnI disclosed that they are required for kinase activity. The only known OPS source in bacteria is through the phosphoserine aminotransferase activity of SerC within the serine biosynthesis pathway, and we demonstrate that an *S. aureus* serC mutant is a serine auxotroph, consistent with a function in l-serine biosynthesis. However, the serC mutant strain could produce SB when provided l-serine, suggesting that SbnI produces OPS for SB biosynthesis *in vivo*. These findings indicate that besides transcriptionally regulating the *sbn* locus, SbnI also has an enzymatic role in the SB biosynthetic pathway.

*Staphylococcus aureus* is a prominent human pathogen that also asymptomatically colonizes a proportion of the human population (1, 2). Though colonization is typically not harmful to the host, *S. aureus* is frequently associated with minor skin and soft-tissue infections. In more serious cases, it is capable of breaching host innate immune responses to gain access to deep tissues, causing more severe and invasive infections, including endocarditis, osteomyelitis, and necrotizing pneumonia (2, 3).

Successful iron uptake from the human host is integral to infection and pathogenesis of *S. aureus* and most other microbial pathogens (4, 5). Iron is reactive and tightly regulated in the human body to restrict toxicity and to limit bioavailability to invading microbial pathogens as a type of innate, nutritional immunity (6). As such, iron in the mammalian host is found either within heme and bound to hemoproteins, intracellularly bound to proteins as free ions or iron-sulfur clusters, stored in ferritin, or extracellularly complexed with glycoproteins like transferrin (7, 8). *S. aureus* has evolved several mechanisms to counter iron restriction and exploit a variety of host iron sources. Iron uptake strategies employed include heme uptake via the iron-responsive surface determinant system and through synthesis and secretion of two iron-chelating siderophores, staphylferrin A (SA)3 and staphylferrin B (SB) (9–11). Siderophores are small molecules with high iron affinity capable of outcompeting extracellular host iron-binding proteins for ferric iron. Iron-bound siderophores are selectively imported by dedicated surface receptors and cognate ABC transporters, HtsABC and SirABC, for SA and SB, respectively (12).

SA and SB have both been implicated in pathogenesis of *S. aureus*, but SB has been recognized for its importance in severe disease phenotypes and promotion of staphylococcal virulence in abscess and endocarditis models of infection (13, 14). Additionally, the SB biosynthetic locus and *sirABC* are among the most strongly up-regulated genes in bacteria isolated from the iron-restricted host (15–17). Importantly, SB biosynthesis can occur independent of TCA cycle activity, which is down-regulated in *S. aureus* as part of the iron-sparking response (18, 19). In contrast, SA biosynthesis is dependent on the TCA cycle for citrate precursors (18).

Staphylferrin biosynthesis is achieved through non-ribosomal peptide synthetase-independent siderophore (NIS) bio-

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3 The abbreviations used are: SA, staphyloferrin A; SB, staphyloferrin B; NIS, non-ribosomal peptide synthetase-independent siderophore; α-KG, α-ke-to glutarate; l-Dap, l-2,3-diaminopropionate; OPS, O-phospho-L-serine; PLP, pyridoxal 5'-phosphate; PK/LDH, pyruvate kinase/lactate dehydroge-nase; RMSD, root mean square deviation; DLS, dynamic light scattering; TCEP, tris(2-carboxyethyl)phosphine; CDMG, carbon defined medium with glucose; TMS, Tris minimal succinate; AMP-PNP, 5'-adenyl-β,γ-imido- diphosphate; TCA, tricarboxylic acid; PDB, Protein Data Bank.

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**SbnI serine kinase activity and role in SB biosynthesis**

**Figure 1. Metabolic pathways for the production of L-Dap and α-KG from glucose or L-serine in *S. aureus*.** Highlighted in light green is the contribution of SbnI in this pathway, which feeds into the previously characterized SB biosynthetic pathway (light blue). **3P**, 3-phospho.

SbnI, the ninth gene product, is a heme-responsive transcriptional regulator for SB production (27). SbnI is required for full expression of *sbnD-H* and thus controls SB-mediated iron acquisition. SbnI binds a 271-bp DNA fragment in the *sbnC* coding region. However, SbnI can bind heme, which abrogates interaction with DNA, thereby limiting SB production. A model is proposed by which SbnI is required for transcription of the full SB biosynthetic operon and senses heme to reduce SB synthesis in favor of heme acquisition (27). This model also provides a mechanism for the findings that *S. aureus* demonstrates a heme-iron preference in vitro (28).

Based on primary sequence analysis, homology was not detected between SbnI and any characterized transcription factors or heme-binding proteins (27), but it is annotated as containing an N-terminal ParB-like domain. ParB is an essential component of the chromosome segregation system in bacteria (29). However, the conserved N-terminal ParB-like domain in SbnI is not responsible for interaction with DNA in ParB (30), leaving the role of the ParB-like domain in SbnI unknown.

The aim of this study was to gain insight into how SbnI functions using X-ray crystallography. The SbnI structure revealed striking structural homology to a recently characterized free serine kinase, SerK, from the archaea *Thermococcus kodakar densis* (31). SerK can phosphorylate free L-serine using ADP to generate OPS for cysteine biosynthesis (32). Herein, we demonstrate that SbnI is also a free serine kinase that uses ATP to phosphorylate L-serine to yield OPS and ADP (Fig. 1). The structure of SbnI, supported by site-directed mutagenesis, suggests that it follows a similar open–close reaction mechanism as proposed for SerK. Additionally, SbnI-generated OPS can be used by SbnA in vitro and serves as the in vivo source of OPS for SB production. To our knowledge, this is the first example of a bacterial free serine kinase and the first described free serine kinase that is ATP-dependent. This enzymatic function earns SbnI an enzymatic role in the SB biosynthetic pathway in addition to its heme-dependent transcriptional regulatory function.

**Results**

**Structure determination of SbnI**

Full-length (254-amino acid) SbnI was not amendable for X-ray crystallography due to its propensity to precipitate. To improve solubility and stability in solution, several expression constructs were made containing varying N- and C-terminal truncations based on predicted secondary structure and disorder identified using PSIPRED and DISOPRED, respectively (33, 34). One construct containing a 14-amino acid C-terminal truncation, SbnI (1–240), had improved stability in solution and produced well-diffracting crystals. The structure of selenomethionine-labeled SbnI (1–240) was determined to 2.5 Å resolution using single wavelength anomalous dispersion in space.
group $P_3$, with one molecule in the asymmetric unit (Fig. 2a).

Data collection and refinement statistics are summarized in Table 1. All 240 residues were modeled with 98% of residues in the most favored regions of the Ramachandran plot. SbnI(1–240) is composed of two domains, domain I and II. Domain I consists of residues Met1–Gln83 and the most favored regions of the Ramachandran plot. Domain II consists of residues Tyr84–Ile205–Ala240, and domain II is composed of residues Asn204. Domain I includes a conserved core ParB/Srx fold, corresponding to the annotated ParB-like domain based on primary sequence. This fold has been described in a functionally diverse ParB/Srx superfamily of proteins. Members are found in varied biological contexts and thus far are described to bind a nucleotide for kinase, ATPase, or DNase activity (32, 36). To our knowledge, no ParB/Srx family member has been found to be directly involved in siderophore biosynthesis. The ParB/Srx core domain is composed of a four-strand mixed $\beta$-sheet and two $\alpha$-helices, $\alpha_2$ and $\alpha_3$ (Fig. 2a), and contains an absolutely conserved GXXR motif, $^{59}$GVHR$^{62}$ in SbnI.

Domain II is composed of a mixed $\alpha/\beta$ fold with a central four-stranded antiparallel $\beta$-sheet surrounded by five $\alpha$-helices. A pair of antiparallel $\beta$-strands abut the main sheet and serve as a linker to domain I. $B$-Factor analysis reveals that domain II has a relatively high average $B$-factor of 109 Å$^2$, compared with 75 Å$^2$ in domain I, suggesting that domain II has more disorder in the crystal and the domains are connected by a flexible linker.

To gain functional insight into SbnI, a search of the SbnI(1–240) structure against structures in the PDB was performed with the Dali server. Five proteins all belonging to the functionally diverse superfamily of ParB/Srx proteins were identified (Z score < 3.9). The most striking observation was the high structural similarity SbnI shared with the top search result, SerK (PDB code 5X0B). Superimposition of SbnI(1–240) with SerK using PDBeFOLD has a root mean square deviation (RMSD) of 2.0 Å for 194 Ca despite sharing only 19% amino acid sequence identity across the aligned residues (Fig. 2b). The other structures also share modest sequence identity (12–23%) and include sulfiredoxin (Srx) from Homo sapiens (PDB code 2RII, RMSD of 2.7 Å across 84 residues), chromosome partitioning protein (ParB) from Sulfolobus solfataricus (PDB code 5K5D, RMSD of 2.5 Å across 71 residues), oncogenic suppressor (Osa) from Shigella flexneri (PDB code 4OVB, RMSD of 3.7 Å across 83 residues), and chromosome segregation protein (Spo0I) from Thermus thermophilus (PDB code 1VZ0, RMSD of 5.4 Å across 76 residues) (31, 36–39).

SerK is a free serine kinase from *T. kodakarensis* that uses ADP to phosphorylate L-serine to generate OPS for cysteine biosynthesis. Of the proteins annotated in the ParB/Srx family, SerK is the only identified kinase, although Osa and Srx both possess ATPase activity (36, 40). Overall, the structures of SerK

### Table 1

**Data collection and refinement statistics for SbnI(1–240)**

| Data collection* | Value |
|------------------|-------|
| Resolution range (Å) | 42–2.50 (2.59–2.50) |
| Space group | $P_3_1$ |
| Unit cell dimensions | |
| $a$, $b$, $c$ (Å) | 55.1, 55.1, 92.7 |
| Unique reflections | 10,883 (1,564) |
| Completeness (%) | 99.9 (100) |
| Redundancy | 2.9 (2.9) |
| Average $I/\sigma(I)$ | 15.8 (2.2) |
| $R$merge | 0.051 (1.007) |
| Wilson $B$-factor (Å$^2$) | 56.4 |
| Anisotropy | 0.501 |

| Refinement | Value |
|-------------|-------|
| $R_{work}$ ($R_{free}$) | 0.219 (0.259) |
| No. of water molecules | 12 |
| RMSD bond length (Å) | 0.003 |
| Average $B$-values (Å$^2$) | 95.0 |
| Ramachandran plot (%) | |
| Most favored regions | 97.5 |
| Disallowed regions | 0.4 |

| PDB code | 5X0E |

* Data collection values in parentheses represent the data for the highest-resolution shell.
and SbnI are very similar. The SerK domain II has a high average $B$-factor and superimposition of SbnI(1–240) with the SerK structure in a “closed” conformation (PDB code 5X0E) suggests how SbnI may possess similar domain flexibility in solution (Fig. 2c). Additionally, structural superimposition and multiple sequence alignments revealed that several residues important for substrate and product binding identified in the SerK crystal structure (PDB code 5X0E) are conserved in SbnI. Moreover, the active-site architecture is highly conserved between the two proteins (Fig. 2d and Fig. S1). Of the active-site residues, SerK Glu$^{30}$ was identified as a catalytically essential residue, and Asp$^{69}$ is required for magnesium ion binding; site-directed mutagenesis of either of these residues abolished SerK kinase activity (31). The homologous residues in SbnI are Glu$^{20}$ and Asp$^{58}$.

The genomic context of $sbnI$ was analyzed to compare with the genomic neighborhoods of homologs. All staphylococcal $sbnI$ homologs are part of the nine-gene SB biosynthetic cluster. More distant homologs co-occur with $sbnA$ and $sbnB$ homologs, either alone or in combination with different putative siderophore biosynthetic enzymes (Fig. 3). More distant SbnI homologs are shorter, and alignments suggest that they have an abbreviated domain II (Fig. S1). Interestingly, a putative $sbnI$ ortholog was identified upstream of the $sbnA$–$H$ gene locus in Ralstonia solanacearum, which was previously thought to lack a SbnI homolog but still produces SB (41). A multiple-sequence alignment of SerK and SbnI homologs used in the genomic neighborhood analysis reveals that certain key residues important for catalysis, substrate, and product binding identified in SerK are fully conserved (Fig. S1). Notably, these include the SerK catalytic residue Glu$^{30}$; magnesium ion-binding residue Asp$^{69}$; residues implicated in interacting with the $\beta$-phosphate of ADP or phosphate group of OPS, His$^{72}$ and Arg$^{73}$; and residues that interact with the serine moiety of OPS, Trp$^{102}$ and Thr$^{223}$ (Fig. S1). More variability is seen with the SerK residues interacting with the adenosine group, raising the possibility that SbnI and other homologs may use a different phosphate donor or binding mode. Overall, the genomic neighborhood and sequence analyses suggest that free serine kinases are found in diverse species belonging to Firmicute and Proteobacteria phyla.

Figure 3. Illustration of gene neighborhoods containing SbnI homologs from diverse species from Firmicute and Proteobacteria phyla. Each predicted gene is represented by an arrow showing the direction of transcription. Gray links connect protein homologous with e value $\leq 1 \times 10^{-40}$, and orthologous genes are indicated in the same color. This figure highlights that SbnI homologs appear in the same genomic context as $sbnA$ and $sbnB$ homologs in these bacterial genomes. The bottom scale shows the length of depicted genomic regions in nucleotide base pairs. Region coordinates used for each species are as follows (GenBank$^{34}$/nucleotide region): S. aureus USA300 FPR3757 (CP000255.1/134324–145881), Staphylococcus pseudintermedius E104 (LAWU0100001.1/151604–163186), Brevibacillus brevis NBRC 110488 (NZ_BDFB01000004.1/355053–396579), Paenibacillus larvae SAG 10367 (NZ_CP020557.1/4429364–4439754), Bacillus badius DSM 5610 (NZ_LVT001000018.1/5060–15111), Marininema mesophilum DSM 45610 (NZ_FNNQ0100001.1/200002–206967), Methylobacterium nodulans ORS 2060 (NC_011894.1/7032971–7044380), Ralstonia solanacearum CQPS-1 (NZ_CP016915.1/169574–186973), and Shewanella denitrificans OS217 (NC_007954.1/663307–674451).
Consurf analysis of SbnI(1–240) was used to map conserved regions to the molecular surface (Fig. 4). Highly conserved residues, including those that form the kinase active site, delineate a groove between domain I and II (Fig. 4, a and b). Structural alignment with the structure of the SerK ternary product complex (PDB code 5X0E) revealed that among SbnI homologs, the putative active site is highly conserved, whereas the remainder of the protein surface is variable (Fig. 4c).

**SbnI is a dimer in solution**

The oligomeric state of SbnI was analyzed using dynamic light scattering (DLS). Because SbnI contains seven Cys residues, the analysis was conducted in the presence of GSH as a reductant. The calculated molecular mass based on amino acid sequence of full-length SbnI is 30 kDa, and the molecular mass measured by DLS was 61 ± 6 kDa with an average of 24 ± 7% of polydispersity, implying that it predominantly forms a dimer in solution (Fig. S2). The molecular mass of SbnI(1–240) by DLS was 28 ± 3 kDa with 34 ± 1% polydispersity (Fig. S2). SbnI(1–240) has a calculated mass of 28 kDa, implying that it is primarily a monomer in solution. These data indicate that the C-terminal 14 amino acids excluded from the SbnI(1–240) construct are important for dimerization of the full-length protein.

**SbnI is a serine kinase that uses L-serine and ATP to generate OPS**

OPS is a substrate for SbnA in SB biosynthesis (25, 26), lending support to our hypothesis that SbnI produces OPS for use by SbnA. To thus test if SbnI is capable of producing OPS, the spectral changes that occur when SbnA binds OPS were used to assay SbnI activity. SbnA has a characteristic absorption maximum at 412 nm attributable to an internal Schiff base formed between its pyridoxal 5’-phosphate (PLP) cofactor and an active-site lysine. Adding OPS to SbnA causes a rapid change in UV-visible spectra with the appearance of absorption peaks at 324 and 467 nm (Fig. 5a), characteristic of the formation of an external aminoacrylate intermediate (25). This spectral change is specific to OPS and does not occur with O-acetyl-L-serine or L-serine (25). SbnA incubated with SbnI, L-serine, and ATP resulted in no change in the UV-visible spectrum of SbnA. However, SbnA incubated with SbnI, L-serine, and ATP led to a shift in the UV-visible spectra indicative of OPS production and reaction with SbnA-PLP to form the external aminoacrylate (Fig. 5, b and c). No spectral change was observed when SbnI was omitted, indicating that only the SbnI enzymatic product could react with SbnA-PLP. Therefore, we conclude that the reaction product is most likely OPS, and SbnI activity is ATP-dependent.

Phosphate acceptors, alternative to L-serine, were tested using a pyruvate kinase/lactate dehydrogenase (PK/LDH) assay for detection of ATP conversion to ADP. L-Threonine, α-KG, and L-Dap were not phosphate acceptors (data not shown). Additionally, SbnI did not phosphorylate the serine residue in a His-Ser dipeptide (data not shown).

SbnI-mediated conversion of ATP to ADP and generation of OPS were monitored using HPLC and 31P NMR. Incubation of SbnI with ATP and excess L-serine led to turnover of ATP to ADP, as detected by HPLC (Fig. 6a), indicating that SbnI has ATPase activity. Unlike SerK, no turnover of ADP to AMP was detected with ADP as a phosphate donor by HPLC (Fig. 6a). ADP was not generated when L-serine was excluded, indicating that this activity requires the presence of L-serine. 31P NMR was also used to monitor SbnI-mediated ATPase activity and generation of OPS from L-serine. The observed disappearance of the ATP γ-phosphate 31P signal with the concomitant appearance of a 31P signal of OPS demonstrates transfer of the ATP γ-phosphate to L-serine to yield ADP and OPS (Fig. 7). The chemical shift of SbnI-generated OPS was consistent with the 31P NMR spectrum measured for an OPS standard (Fig. S3). Attempts to obtain a co-crystal structure of SbnI with identified substrates (or ATP analog, AMP-PNP) or products, both in the presence and absence of heme, have not been met with success.

**SbnI active-site variants**

The role of Glu20 and Asp58 in SbnI kinase function were tested by site-directed mutagenesis. Two mutants, each containing a single alanine substitution, SbnI E20A and SbnI D58A, were generated. Using the PK/LDH assay, we determined that the mutants were incapable of turning over ATP to ADP in the presence of L-serine. Additionally, no ADP could be detected by HPLC in reactions containing the SbnI mutants incubated with ATP and L-serine (Fig. 6b); together, these data allow us to conclude that these mutants are catalytically inactive. These data also correlate with SbnA-PLP UV-visible absorption spectra that demonstrated that these SbnI variants also do not produce OPS (Fig. 5, d and e). The importance of these residues in catalysis is consistent with the reaction mechanism presented for SerK in which substrate binding promotes conformational closure positioning the catalytic Glu30 (Glu20 in SbnI) close to the hydroxyl group of bound L-serine. Glu30 is a catalytic base that deprotonates the hydroxyl group of L-serine. The deprotonated hydroxyl can then attack the phosphorus atom of the ADP β-phosphate to yield OPS and AMP (31). Our results suggest SbnI uses a similar two-ligand binding sequential mechanism but instead uses L-serine and ATP to yield ADP and OPS.

**Sbnl serine kinase activity and role in SB biosynthesis**

Figure 4. Conservation of surface residues of SbnI(1–240) generated using ConSurf. a and b, ribbon and surface representations of SbnI(1–240) have AMP and OPS modeled based on a structural alignment with the SerK ternary product complex (PDB code 5X0E). c, surface representation of SbnI(1–240) after 180° rotation about the x axis. Conserved amino acids are colored maroon, residues of average conservation are white, and variable amino acids are turquoise.

Figure 5, a and b, ribbon and surface representations of SbnI(1–240) have AMP and OPS modeled based on a structural alignment with the SerK ternary product complex (PDB code 5X0E). c, surface representation of SbnI(1–240) after 180° rotation about the x axis. Conserved amino acids are colored maroon, residues of average conservation are white, and variable amino acids are turquoise.
Kinetic analysis of SbnI kinase activity

To obtain kinetic parameters for the SbnI kinase activity, SbnI enzymatic turnover was monitored using an established coupled assay for ADP using PK/LDH. The steady-state kinetic parameters of SbnI reaction with ATP and L-serine were determined and are summarized in Table 2. The saturating concentration of L-serine was beyond conditions permissive to the assays, and thus $K_m$ could not be accurately determined. A second assay to measure SbnI enzymatic turnover employed SbnA-dependent turnover of OPS coupled to a phosphate release detection assay. SbnA is the most likely in vivo acceptor of OPS generated by SbnI as SbnA requires OPS as a substrate for synthesis of SB precursors, L-Dap and $\alpha\$-KG, in concert with SbnB. Using excess concentrations of SbnA, the coupled assay demonstrated that SbnA could use SbnI-generated OPS and supplied L-glutamate to generate its products, N-[(1-amino-1-carboxyl-2-ethyl)]glutamic acid and inorganic phosphate. The kinetic parameters of SbnI reaction with ATP and L-serine using the SbnA coupled assay were determined and are summarized in Table 2. The kinetic parameters measured using both methods agree with rates and $K_m$ values in the same order of magnitude. SbnI mutants E20A and D58A were catalytically insufficient to accurately measure enzyme rates. Compared with SerK, SbnI has a $K_m$ for ATP 1 order of magnitude lower than SerK has for ADP and a higher $K_m$ for L-serine by 2 orders of magnitude.

The enzymatic activity of SbnI(1–240) was also measured because the 14-amino acid C-terminal truncation does not exclude any regions in the SerK structures identified as required for substrate and product binding or catalysis. SbnI(1–240) displayed decreased kinase activity compared with full-length SbnI. These data correlate with the HPLC and spectral data with SbnA for SbnI(1–240) having intermediate activity relative to full-length SbnI (Figs. 5f and 6b). Plots of initial velocities used for determination of kinetic constants are included in Fig. S4. Given that SbnI(1–240) is a monomer in solution (see above), we conclude that dimerization/oligomerization of SbnI is not essential for its kinase activity.
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Physiological function of ATP-dependent serine kinase activity in S. aureus

The serC gene encodes the enzyme responsible for OPS synthesis from 3-phosphohydroxypropionate in the S. aureus serine biosynthesis pathway. To our knowledge, before this work, SerC activity was the only identified metabolic source of OPS in S. aureus and was the assumed source of OPS for SB biosynthesis (18, 26). To test whether other metabolic sources of OPS exist in S. aureus, wildtype USA300 and a serC transposon insertion mutant USA300 strain (serC) were grown in chemically defined media containing glucose with and without L-serine (18, 26). Whereas SbnI free L-serine kinase activity is functionally and enzymatically distinct from that of Ser/K, an enzyme in a biosynthetic pathway for cysteine in the thermophilic archaea T. kodakarensis (31). ATP is used as the phosphate donor in the reaction catalyzed by SbnI, whereas SerK uses ADP. Interestingly, hyperthermophiles generally use ADP in place of ATP in key glycolytic enzymes as an adaptation to life at high temperatures, presumably because ADP is more stable than ATP (43). Whereas SbnI is capable of phosphorylation of free L-serine, it was not capable of mediating phosphorylation to serine residues within a His-Ser dipeptide. Bacterial Ser/Thr kinases such as S. aureus Stk1 (also named PknB) (44) and cognate phosphatases function as molecular switches that have key roles in bacterial cell signaling as in eukaryotic systems (45). However, SbnI free L-serine kinase activity is functionally and enzymatically distinct from that of Ser/Thr kinases.

SbnI(1–240) retains partial catalytic efficiency compared with full-length SbnI. As SbnI(1–240) is lacking 14 residues at

Discussion

SbnI is a free L-serine kinase that makes OPS, which serves as a substrate for SbnA and is a precursor for SB biosynthesis. Insight into the kinase function of SbnI was gained based on homology identified by structural similarity with SerK, an enzyme in a biosynthetic pathway for cysteine in the thermophilic archaea T. kodakarensis (31). ATP is used as the phosphate donor in the reaction catalyzed by SbnI, whereas SerK uses ADP. Interestingly, hyperthermophiles generally use ADP in place of ATP in key glycolytic enzymes as an adaptation to life at high temperatures, presumably because ADP is more stable than ATP (43). Whereas SbnI is capable of phosphorylation of free L-serine, it was not capable of mediating phosphorylation to serine residues within a His-Ser dipeptide. Bacterial Ser/Thr kinases such as S. aureus Stk1 (also named PknB) (44) and cognate phosphatases function as molecular switches that have key roles in bacterial cell signaling as in eukaryotic systems (45). However, SbnI free L-serine kinase activity is functionally and enzymatically distinct from that of Ser/Thr kinases.

SbnI(1–240) retained partial catalytic efficiency compared with full-length SbnI. As SbnI(1–240) is lacking 14 residues at...
SbnI serine kinase activity and role in SB biosynthesis

Figure 8. a, growth kinetics of S. aureus USA300 wildtype (black lines) and serC transposon insertion mutant (red lines) strains in CDMG (solid lines) and CDMG without l-serine (CDMG – l-Ser) (dashed lines). b, agar plate disc diffusion bioassays were performed using culture supernatants prepared from S. aureus USA300 strains (wildtype and sbnI and serC transposon insertion mutants), as indicated on the x axis, that were grown for 16 h in Chelex treated Tris minimal succinate (c-TMS) medium. The black dots (labeled SA (ΔsirA)) are a measure for the presence of SA in culture supernatants, and the gray dots (labeled SB (ΔhtsABC)) are a measure of SB in culture supernatants based on the growth radius around the disc. The disc radius (3 mm) is subtracted from the reported growth radius. Lines represent the S.D. ***, p < 0.0002; ****, p < 0.0001.

the C terminus and is monomeric, either the C terminus or dimerization is required for full kinase function. The residues shown to be required for phosphotransfer in SerK (Glu30 and Asp69) are conserved in SbnI and present in the truncated protein. Moreover, three of four residues interacting with the serine substrate in SerK are conserved (Glu30, Trp102, and Thr223), and the fourth position is a conservative substitution of His225 with Phe203 in SbnI (Fig. 2d). In contrast, interactions of SerK with ADP/AMP are poorly conserved. Of eight residues making key contacts, only three residues (Ser43, His72, and Arg73) are conserved in SbnI, and two of these interact with the phosphate groups. Attempts to obtain crystals with substrates or products bound to SbnI or SbnI(1–240) have not been met with success, and the binding mode of ATP to SbnI remains elusive.

Structural analysis of SerK suggests that the conformational closure upon binding both substrates positions the catalytic glutamate (Glu30) to deprotonate l-serine to attack the terminal phosphate of ADP. With no published structure of substrate or product-free SerK, the apo-SbnI(1–240) structure supports this proposed mechanism in that the unbound form is in an open conformation to expose the binding pocket. Measurement of SbnI enzyme kinetic parameters revealed that, compared with SerK, it has a relatively low selectivity for l-serine. The comparatively low $k_{cat}/K_m$ could relate to the physiological role of SbnI in S. aureus. SerK supplies cysteine synthase with OPS to produce cysteine and may represent an ancient heterothrophic mechanism of amino acid metabolism (32). Interestingly, this cysteine synthase is a distant SbnA homolog and a true OPS sulfhydrilase. Also, SerK is postulated to provide an advantage by enabling carbon from serine to be directed to glycolysis and gluconeogenesis by conversion to OPS (32). In vitro evidence suggests that S. aureus uses amino acids to support gluconeogenesis (46). However, serine is used to generate ATP and acetate rather than to facilitate gluconeogenesis as in T. kodakarense (32, 46). In contrast, SbnI kinase activity fulfills a distinct physiological role, providing substrate necessary for SB production. The comparatively low $k_{cat}/K_m$ may allow SbnI to respond to a greater range of substrate concentrations, such that at high l-serine concentrations, SbnI increases the rate of OPS production for SB production and possibly other metabolic processes. Although SbnI has lower catalytic efficiency than SerK (Table 2), the related ParB/Srx family member, Srx, is similar to SbnI as it is not a highly efficient enzyme. The Srx ATP-dependent reduction of peroxiredoxin sulfenic acid activities described in mammals and plants so far is slow, with turnover rates of 0.2–0.5 min$^{-1}$ for ATP, which are an order of magnitude lower than SbnI (47–49). However, the catalytic efficiencies of Srx are 0.8–8.4 mm$^{-1}$min$^{-1}$ and in the range measured for SbnI (47, 49).

SbnI is a sufficient biological source of OPS for SB biosynthesis and contributes to the functional modularity of the sbn locus. A recognized characteristic of bacterial networks is a high degree of modularity and sparse connectivity between individual functional modules, where a functional module refers to a group of biological components that are spatially isolated or chemically specific and work together for a discrete biological function (50). The functional redundancy of enzymes encoded in the sbn locus for generation of precursor substrates, SbnG and SbnI, decreases dependence on central metabolism and contribute to the modularity of the sbn locus. SbnG is functionally redundant with the TCA cycle citrate synthase, CitZ (23). Moreover, SbnG activity allows for SB biosynthesis to occur independent of TCA cycle activity (18). This functional independence is important because S. aureus elicits an iron-sparing response during infection resulting in down-regulation of the TCA cycle. We propose that the functional redundancy of SbnI with SerC for OPS production allows SB biosynthesis to occur independent of glycolysis as SerC substrate is funneled from 3-phosphoglycerate. Additionally, the serine biosynthetic pathway is regulated by negative feedback where SerA, metabolically upstream of SerC, is allosterically inhibited by l-serine and could limit the amount of SerC-derived OPS available to support SB synthesis when serine is abundant. Thus, an alternative, SB-dedicated, OPS synthetic route via SbnI is advantageous. Together with SbnG, SbnI allows SB biosynthesis to occur autonomously from glycolysis and TCA cycle activity by generating precursor substrates dedicated to SB biosynthesis.

This observed metabolic redundancy may improve robustness and help buffer environmental perturbations S. aureus encounters during infection. Glucose is the preferred carbon source by S. aureus and available at concentrations to support growth in human blood (51). However, within staphylococcal
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Experimental procedures

Cloning and site-directed mutagenesis

Constructs with an N-terminal His6 tag and thrombin cleavage site were generated in pET28a vectors for recombinant expression of S. aureus full-length SbnI, residues 1–254, with the first codon mutated from the native TTG to a common start codon, ATG, and the S. aureus C-terminal truncated construct SbnI(1–240) (residues 1–240). The S. aureus SbnI nucleotide sequence can be accessed in the GenBank™ database under accession code NC_009641.1 (positions 90178–90942) (gene locus NWMN_RS00380), and the amino acid sequence can be accessed through NCBI Protein Database under NCBI accession WP_001015549.1. Briefly, a megaprimer-based whole-plasmid synthesis PCR cloning protocol was used to clone constructs amplified from chromosomal DNA from S. aureus strain Newman (54). S. aureus SbnI variants E20A and D58A were produced using a single-primer mutagenesis method (55). Primers used in this study are summarized in Table S1. All clones were introduced into E. coli BL21 (ADE3) and confirmed by DNA sequencing. Bacterial strains and plasmids used in this study are summarized in Table 3.

Protein expression and purification

Recombinant full-length SbnI, SbnI(1–240), SbnI E20A, and SbnI D58A constructs were overexpressed in E. coli BL21 (ADE3) cells. Cultures were grown in 2x YT medium supplemented with 25 µg/ml kanamycin at 30 °C to an $A_{600}$ of 0.7–0.9. Cultures were then induced with 0.5 mM isopropyl β-D-thiogalactopyranoside and grown for an additional 18 h at 20 °C. Cells were pelleted by centrifugation at 4,400 × g for 7 min at 4 °C and resuspended in buffer containing 50 mM HEPES (pH 7.4), 300 mM NaCl, 5% (v/v) glycerol, 2 mM tris(2-carboxyethyl) phosphine (TCEP), and 10 mM imidazole on ice. Approximately 5 mg of DNase was added to cell suspension before lysis at 4 °C using an EmulsiFlex-C5 homogenizer (Avestin). Insoluble material was removed by centrifugation at 39,000 × g for 1 h, and recombinant protein was purified from soluble lysate using a HisTrap nickel affinity column (GE Healthcare) by elution with an imidazole gradient. Protein was dialyzed against 50 mM HEPES (pH 7.4), 100 mM NaCl, 5% (v/v) glycerol, and 2 mM TCEP and then cleaved with thrombin at a 1:500 ratio by weight of His$_6$ protein to remove the His$_6$ tag over 18 h at 4 °C. Subsequently, recombinant protein was dialyzed into 50 mM HEPES (pH 7.4), 5% (v/v) glycerol, and 2 mM TCEP and further purified by anion-exchange chromatography using a Source 15Q column (GE Healthcare). Purified protein was obtained by elution with a NaCl gradient and further dialyzed into 50 mM HEPES (pH 7.4), 300 mM NaCl, 5% (v/v) glycerol, 2 mM GSH. The sample was concentrated to ~20 mg/ml, flash-frozen, and stored at −80 °C. Selenomethionine-incorporated SbnI(1–240) was produced by methods described previously (56) and purified as described above for native SbnI(1–240).

His$_6$-tagged SbnA was expressed in E. coli BL21 (ADE3) cells from the plasmid pET28a, purified by His tag affinity chromatography, and digested with the thrombin to remove the His$_6$ tag. The protein was further purified by anion-exchange chromatography using the previously published method for
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Table 3
Bacterial strains and plasmids used in this study

| Strains             | Description                                                                 | Source or reference |
|---------------------|-----------------------------------------------------------------------------|---------------------|
| E. coli             | F−ompT gal dcm lon hsdsM (rads− m+λ) (DE3 [lacI lacUV5-T7 gene 1 ind1 sam7 nin5]) | Novagen             |
| BL21 (ADE3)         | USA300 LAC cured of antibiotic resistance plasmid                           | Ref. 68             |
| S. aureus           | USA300 E2 serC::FNS2                                                         | Ref. 69             |
| JE2 serC::FNS2      | USA300 E2 serC::FNS2                                                         | Ref. 69             |
| JE2 sbnl::FNS2      | USA300 sbnl::FNS2                                                           | This study          |
| serC                | USA300 sbnl::FNS2                                                           | This study          |
| sbnl                | RN6390ΔserA::KmR; SB transport-deficient mutant                            | Ref. 35             |
| sirA                | RN6390ΔsirA::KmR; SB transport-deficient mutant                            | Ref. 72             |
| Plasmids            |                                                                             |                     |
| pET28a-sbnl         | IPTG-inducible expression vector containing sbnl; KmR                      | This study          |
| pET28a-sbnl−240     | IPTG-inducible expression vector containing sbnl−240; KmR                 | This study          |
| pET28a-sbnl-E20A     | IPTG-inducible expression vector containing sbnlE20A; KmR                 | This study          |
| pET28a-sbnl-D58A     | IPTG-inducible expression vector containing sbnlD58A; KmR                 | This study          |

Improved SbnA solubility (26). SbnA was dialyzed into 50 mM Tris, pH 8, 100 mM NaCl, and 2 mM TCEP, concentrated to ~20 mg/ml, and stored at ~80 °C.

Crystallization, data collection, and structure determination of SbnI(1–240)

Selenomethionine-labeled SbnI(1–240) crystals were grown by sitting-drop vapor diffusion at 4 °C in 2-μl drops with a 1:1 mixture of ~20 mg/ml SbnI(1–240) in 50 mM HEPES (pH 7.4), 100 mM NaCl, 5% (v/v) glycerol, and 2 mM TCEP with reservoir solution containing 0.18 M HEPES (pH 7.5) and 20% (v/v) PEG 8000. Crystals were briefly soaked in reservoir buffer supplemented with 30% (v/v) glycerol for cryoprotection and flash-frozen in liquid nitrogen. A single-wavelength anomalous diffraction data set was collected at the Canadian Light Source on Beamline 08B1-1 (57). The data were processed and scaled using XDS (58, 59). Crystals were of space group P31, with one molecule in the asymmetric unit. Five selenomethionine sites were identified for phasing to build a preliminary model using AutoSol (initial figure of merit of 0.35) and Autobuild (187 of 240 residues built) programs in Phenix (60). Manual building was done using Coot (61), and refinement was performed with phenix.refine using translation liberation screw parameters with three groups (62). The refined structure contains Met1–Ala240, one glycerol, and 12 water molecules. Data collection and refinement statistics are summarized in Table 1. Structure figures were generated in PyMOL (PyMOL Molecular Graphics System, version 1.8, Schrödinger, LLC, New York). Domain analysis was done using the Dali server for comparison of the protein structure against structures in the PDB (63).

Genomic neighborhood and conservation analysis

Protein homology between genomic regions carrying SbnI orthologs was plotted using a custom Biopython script, bio.links.py, based on output from BLASTP 2.2.28 (e value ≤ 1.00e−40) (https://github.com/minevskiy/bioinformatics). Species used for comparison were found by BLAST search of SbnA or SbnI and STRING analysis of SbnI. Orthologous genes are indicated in the same color.

Sequence conservation was mapped onto the SbnI(1–240) structure using ConSurf (64). The multiple sequence alignment used for the analysis was generated using default ConSurf parameters and the SbnI amino acid sequence as the search sequence.

Dynamic light scattering

Samples of SbnI and SbnI(1–240) were analyzed by DLS using a DynaPro Plate Reader (Wyatt Technologies). Protein was diluted to 0.5 mg/ml with 50 mM HEPES (pH 7.4), 300 mM NaCl, 5% (v/v) glycerol, and 2 mM GSH, and results were generated based on averaging five 5-s acquisitions. Data were collected at room temperature. Values reported are an average of data collected.

UV-visible spectrophotometry analysis of SbnI OPS production using SbnA

UV-visible spectra were collected using a Varian Cary 50 UV-visible spectrophotometer. The SbnA-PLP spectrum was recorded at a concentration of 15 μM in 50 mM HEPES, pH 7.4, 100 mM NaCl, 5% (v/v) glycerol. The spectrum of SbnA amino-acrylate aldime complex was recorded immediately after the addition of 30 μM OPS (25). The spectral shift observed when SbnA binds OPS was used to evaluate OPS production by SbnI. The spectrum of 15 μM SbnA in 50 mM HEPES, pH 7.4, 100 mM NaCl, 5% (v/v) glycerol, 20 mM MgCl2, 25 mM L-serine, and 5 mM ATP was recorded before and after the addition of 15 μM SbnI, SbnI E20A, SbnI D58A, or SbnI(1–240). Phosphate donor specificity was also examined using 5 mM ADP in place of ATP.

HPLC

Kinase activity of SbnI was detected using HPLC to examine production of ADP from ATP. The reaction mixture was composed of 50 mM HEPES (pH 7.4), 100 mM NaCl, 2.5% glycerol, 50 mM L-serine, 0.25 mM ATP or ADP, 10 mM MgCl2, and 5 μM SbnI, SbnI E20A or SbnI D58A. The reaction was carried out for 1 h at room temperature (22 °C). The protein was removed by centrifugation using a 3K Nanosep column. The filtrate was
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0.2-μm filtered and analyzed by HPLC using a Waters 2695 Separations HPLC module (Milford, MA) equipped with a Waters 2996 photodiode array detector and a Luna 3-μm PFP(2) 50 X 4.6-mm LC column (Phenomenex) using a linear gradient of 0–15% methanol in 0.1 M ammonium acetate, pH 4.5, over 10 min at 1 ml min⁻¹. Analytes were detected by the absorbance at 258 nm.

### ATP-dependent serine kinase activity

ATP-dependent serine kinase activity of SbnI was measured using a PK/LDH coupled assay. The assay is based on a reaction in which the regeneration of hydrolyzed ATP is coupled to the oxidation of NADH (66). The rate of NADH absorbance decrease at 340 nm (A₃₄₀, nm = 6,220 M⁻¹ cm⁻¹) is proportional to the rate of ATP conversion to ADP by SbnI kinase activity. Coupled reactions contained 50 mM HEPES, pH 7.4, 100 mM NaCl, 2.5% (v/v) glycerol, 10 mM MgCl₂, 2 mm L-serine, 2 mM phosphoenolpyruvate, 1/3 of the final reaction mixture volume of PK/LDH enzyme (from rabbit muscle, Sigma-Aldrich, catalog no. P-0294), 5 mM ATP, and 100 mM L-serine. The mixture was incubated for 5 min to remove any contaminating ADP. Continuous measurement at 340 nm was recorded for 2 min before the addition of 0.5 μM SbnI or SbnI(1–240), SbnI E20A, or SbnI D58A enzyme to start the reaction. The assay was run for 10 min. To determine kinetic parameters, the initial velocities of SbnI and SbnI(1–240) kinase reactions in the presence of varying concentrations of ATP with 100 mM L-serine and in the presence of varying concentrations of L-serine with 10 mM ATP were recorded. All data were collected on a Varian Cary 50 UV-visible spectrophotometer at room temperature (22 °C) and a total of three replicates were collected for each reaction condition. The concentration of inorganic phosphate release from OPS was determined using the extinction coefficient A₅₀₆₀ nm = 11,000 M⁻¹ cm⁻¹ (67). Data were fit by nonlinear regression using a Michaelis–Menten model in GraphPad Prism version 6.

### Measurement of serine kinase activity

The reaction mixture contained 50 mM HEPES (pH 7.4), 100 mM NaCl, 2.5% (v/v) glycerol, 10 mM MgCl₂, 48 mM L-serine, 5 mM ATP, and 5% D₂O. NMR spectra were collected at 25 °C using a broadband frequency probe with Z-magnetic field gradient in a Bruker Avance III 500-MHz spectrometer. One-dimensional ³¹P NMR spectra were recorded at different time points before and after the addition of 4.8 μM SbnI until reaction completion. The spectra were referenced to 2,2,6,6-tetramethylpiperidine, which was set to 0 ppm. The spectra were processed and using TopSpin™ (Bruker). The chemical shifts of ATP (65) and OPS were assigned using reference spectra (Fig. S3).

### S. aureus bacterial strains and growth conditions

Experiments were performed with a derivative of S. aureus USA300 LAC cured of the 27- kb plasmid encoding macrolide resistance (68). The plasmid-cured USA300 LAC is referred to as USA300. Transposon insertion mutants JE2 serC::ΔΦNΔΣ;EmR (SAUSA300_1669) and JE2 sbni::ΔΦNΔΣ;EmR (SAUSA300_0126) were obtained from the Nebraska Transposon Mutant Library containing the resistance cassette ermB, which confers resistance to erythromycin (69). Transposons were transduced to USA300 background strain using phage λ from S. aureus USA300 LAC cured of the 27-kb plasmid encoding macrolide resistance (68). The plasmid-cured USA300 LAC is referred to as USA300 throughout. Transposon insertion mutants JE2 serC::ΔΦNΔΣ;EmR (SAUSA300_1669) and JE2 sbni::ΔΦNΔΣ;EmR (SAUSA300_0126) were obtained from the Nebraska Transposon Mutant Library containing the resistance cassette ermB, which confers resistance to erythromycin (69). Transposons were transduced to USA300 background strain using phage λ from S. aureus USA300 LAC cured of the 27-kb plasmid encoding macrolide resistance (68). The plasmid-cured USA300 LAC is referred to as USA300 throughout (Table 3). Bacterial growth curves to test serine auxotrophy were performed in chemically defined medium with 0.4% (w/v) glucose (CDMG) as described previously (18), with and without L-serine. Briefly, colonies of wildtype USA300 or serC transposon insertion mutant were inoculated from tryptic soy agar into 2 ml of CDMG overnight at 37 °C. Cells were normalized to an A₆₀₀ of 0.1 and washed twice with CDMG lacking L-serine (CDMG-Ser), and 5 μl of the resuspension was used to inoculate 200-μl aliquots of CDMG or CDMG-Ser in 96-well plates. Cultures were grown in a TECAN plate reader for 24 h at 37 °C with 10 s of shaking every 10 min, and the A₆₀₀ was assessed every 30 min. Data are representative of three independent experiments, and error bars signify S.E.

### Disc diffusion assays to assess siderophore production

Concentrated spent culture supernatants were prepared from 10-ml cultures of S. aureus USA300, serC transposon insertion mutant, and sbni transposon insertion mutant grown for 16 h in Chexlex-100–treated Tris minimal succinate (TMS), as described previously (71), in a flask/volume ratio of 10:1 at 37 °C with shaking at 200 rpm without antibiotic selection. Growth was assessed via A₆₀₀ and culture densities were normalized. Bacterial cells were pelleted by centrifugation, and culture supernatants were filter-sterilized and lyophilized overnight. Dried material was resuspended in 0.5 ml of sterile double-distilled H₂O. To assess growth promotion of concentrated culture supernatants, S. aureus strain RN6390 sirA
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mutant (growth of this mutant is dependent on SA in supernatant) or itsABC mutant (growth of this mutant is dependent on SB in supernatant) derivatives, as described previously (72), were seeded into TMS agar containing 10 μM ethylenediamine-N,N’-bis(2-hydroxyphenylacetic acid) to 2 × 10^5 cells/ml. 10 μl of concentrated supernatant was applied to sterile paper discs placed on TMS agar containing the seeded reporter strains, and growth radii about the discs were measured after 24 h of incubation at 37 °C. The reported growth radius has the disc radius (3 mm) subtracted. Statistical analyses were conducted using two-way analysis of variance.

Author contributions—M.M.V., D.E.H., and M.E.P.M. conceptualization; M.M.V., C.P.-B., and K.L.B. data curation; M.M.V., C.P.-B., K.L.B., and M.E.P.M. formal analysis; M.M.V., C.P.-B., and K.L.B. visualization; M.M.V., D.E.H., and M.E.P.M. methodology; M.M.V. and M.E.P.M. writing–original draft; M.M.V., D.E.H., and M.E.P.M. writing–review and editing; D.E.H. and M.E.P.M. resources; M.E.P.M. supervision; M.E.P.M. funding acquisition; M.E.P.M. investigation.

Acknowledgments—We thank Kateryna Ivanyakmenko for assistance with genomic neighborhood analysis and figure generation. We also thank Dr. Anson Chan for assistance with data collection and Mariko Ikehata and Angele Arrieta for technical assistance. We thank the staff at the CLS for assistance with crystallographic data collection. Support for infrastructure for structural biology was provided by the Canadian Foundation for Innovation (to M. E. P. M.). Research described in this paper was performed using beamlines 08B1-1 and 08ID-1 at the Canadian Light Source, which is supported by the NSERC, the National Research Council Canada, the Canadian Institutes of Health Research, the Province of Saskatchewan, Western Economic Diversification Canada, and the University of Saskatchewan.

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