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Penicillin-Binding Protein 1 (PBP1) of *Staphylococcus aureus* Has Multiple Essential Functions in Cell Division

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ABSTRACT Bacterial cell division is a complex process requiring the coordination of multiple components to allow the appropriate spatial and temporal control of septum formation and cell scission. Peptidoglycan (PG) is the major structural component of the septum, and our recent studies in the human pathogen *Staphylococcus aureus* have revealed a complex, multistage PG architecture that develops during septation. Penicillin-binding proteins (PBPs) are essential for the final steps of PG biosynthesis; their transpeptidase activity links the peptide side chains of nascent glycan strands. PBP1 is required for cell division in *S. aureus*, and here, we demonstrate that it has multiple essential functions associated with its enzymatic activity and as a regulator of division. Loss of PBP1, or just its C-terminal PASTA domains, results in cessation of division at the point of septal plate formation. The PASTA domains can bind PG and thereby potentially coordinate the cell division process. The transpeptidase activity of PBP1 is also essential, but its loss leads to a strikingly different phenotype of thickened and aberrant septa, which is phenocopied by the morphological effects of adding the PBP1-specific β -lactam, meropenem. Together, these results lead to a model for septal PG synthesis where PBP1 enzyme activity is required for the characteristic architecture of the septum and PBP1 protein molecules enable the formation of the septal plate.

IMPORTANCE Bacterial cell wall peptidoglycan is essential, and its synthesis is the target of clinically important antibiotics such as β -lactams. β -lactams target penicillin-binding proteins (PBPs) that assemble new peptidoglycan from its building blocks. The human pathogen *Staphylococcus aureus* only has two essential PBPs that can carry out all the functions necessary for growth and division. In the absence of the confounding antibiotic resistance-associated PBP PBP2A, PBP1 is required for cell division, and here, we have found that it has several essential functions, both as an enzyme and as a coordinator by binding to cell division proteins and to its peptidoglycan product, via its PASTA domains. This has led to a new model for cell division with PBP1 responsible for the synthesis of the characteristic architectural features of the septum.

KEYWORDS *Staphylococcus aureus*, cell division, penicillin-binding proteins, peptidoglycan

Peptidoglycan (PG) is the major structural component of the bacterial cell wall and is essential for maintaining cell shape, integrity, and survival (1–3). The final stages of assembly of this large polymeric molecule are mediated by penicillin-binding proteins (PBPs), key PG synthases that, through their transglycosylase (TG) and

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transpeptidase (TP) activities, polymerize glycan chains and cross-link them into a mesh-like hydrogel (4, 5). Since the cell wall is essential for maintaining bacterial life, PBPs and PG synthesis are a target of some of the most important antibiotics, β -lactams (penicillins) and glycopeptides (vancomycin) (6, 7). The major human pathogen *Staphylococcus aureus* has a minimalist PBP system, as it encodes only four PBPs, PBP1 to PBP4 (8). Only PBP1 (class B PBP with only TP activity, bPBP) and PBP2 (class A bifunctional PBP with both TG and TP activities, aPBP) are essential and sufficient for septal and peripheral PG synthesis in *S. aureus* (8, 9). PBP2 is the major PG synthase of *S. aureus*, and the septum formation activity of PBP2 is mediated by its substrate, lipid II (10). Although PBP2 is essential, loss of its TP activity can be compensated for by a horizontally acquired class B PBP2A in methicillin-resistant *S. aureus* (MRSA) (11). PBP2A, however, cannot replace PBP1, whose loss is detrimental to the viability of *S. aureus* (12). PBP1 and PBP3 (bPBP) form cognate pairs with the monofunctional TGs, FtsW and RodA, belonging to the SEDS (shape, elongation, division, and sporulation) family (13) to facilitate septum formation (PBP1-FtsW) and to maintain the prolate cell shape (PBP3-RodA) of *S. aureus*, respectively (14). Activation of the transglycosylase activity of FtsW requires complex formation with PBP1 (15). PBP4 is a class C PBP with D,D-carboxypeptidase activity (cPBP) and has a TP activity that contributes to the high-level cross-linking of PG and MRSA resistance to β -lactams (16, 17).

The cell wall of Gram-positive bacteria is decorated with wall teichoic acid (WTA) glycopolymers (18). WTA regulates cell shape, ion homeostasis, autolytic enzymes, growth, and division (19). In *S. aureus*, WTA plays a crucial role in virulence, MRSA resistance to β -lactam antibiotics, PBP4 localization at the septum, and PG cross-linking (20–23).

Although *S. aureus* PBPs have been studied over many years, the specific roles of PBP1 in cell division, PG synthesis, and architecture are not well understood. Previous studies have shown that while PBP1 is essential, its TP activity is not, implying another role (12, 14). However, this work was performed in an MRSA background that contains PBP2A, encoded by *mecA*, which is non-native to *S. aureus* (24). While PBP2A cannot replace PBP1, how these proteins interact is unknown. We have recently shown that the presence of *mecA* has a profound effect on cellular physiology (25). Thus, it is important to understand individual and combined roles of *S. aureus* PBPs in both the presence and absence of the exogenous PBP2A, as the vast majority of *S. aureus* infections are caused by methicillin-sensitive strains.

RESULTS

***S. aureus* PBP1 PASTA domains are essential for growth and PBP1 functionality.**

PBP1 has a short cytoplasmic fragment, a membrane-spanning sequence, an exocytosolic N-terminal pedestal domain, and a C-terminal region consisting of the TP domain and two PASTA domains (for penicillin-binding protein and serine/threonine kinase-associated domain) (26, 27). We created a set of conditional mutants of *pbp1* to investigate the role of PBP1 in cell division and PG synthesis. An ectopic copy of *pbp1* under the control of the *P_{spac}* promoter (*P_{spac}-pbp1*) was placed at the lipase locus (*geh::P_{spac}-pbp1*) of *S. aureus* SH1000, and a series of changes were made in this genetic background at the native *pbp1* locus: (i) an in-frame deletion of *pbp1* ($\Delta pbp1$), (ii) a deletion of the region encoding the two PASTA domains (*pbp1* $_{\Delta PASTA}$), and (iii) the substitution of the catalytic Ser314 to Ala in the TP domain (*pbp1*^{*}) (Fig. 1A and B). We examined the essentiality of PBP1, the PASTA domains, and the active TP domain with these mutants. Depletion of PBP1 via IPTG (isopropyl- β -D-thiogalactopyranoside) removal (Fig. 1C and Fig. S1A and B) resulted in cell death, confirming the essentiality of PBP1 (Fig. 1C and D and Fig. S1C and D). Deletion of the PASTA domains also led to growth inhibition and more than 99% cell death within 4 h (Fig. 1D and Fig. S1C and D). Importantly, this phenotype was not associated with PBP1 $_{\Delta PASTA}$ instability (Fig. 1C and Fig. S1A) or loss of its ability to bind its substrate analogue BocillinFL (Fig. S1B). In contrast, deletion of the PASTA domains of *Streptococcus pneumoniae* PBP2x, a PBP1 orthologue, resulted in a complete loss of

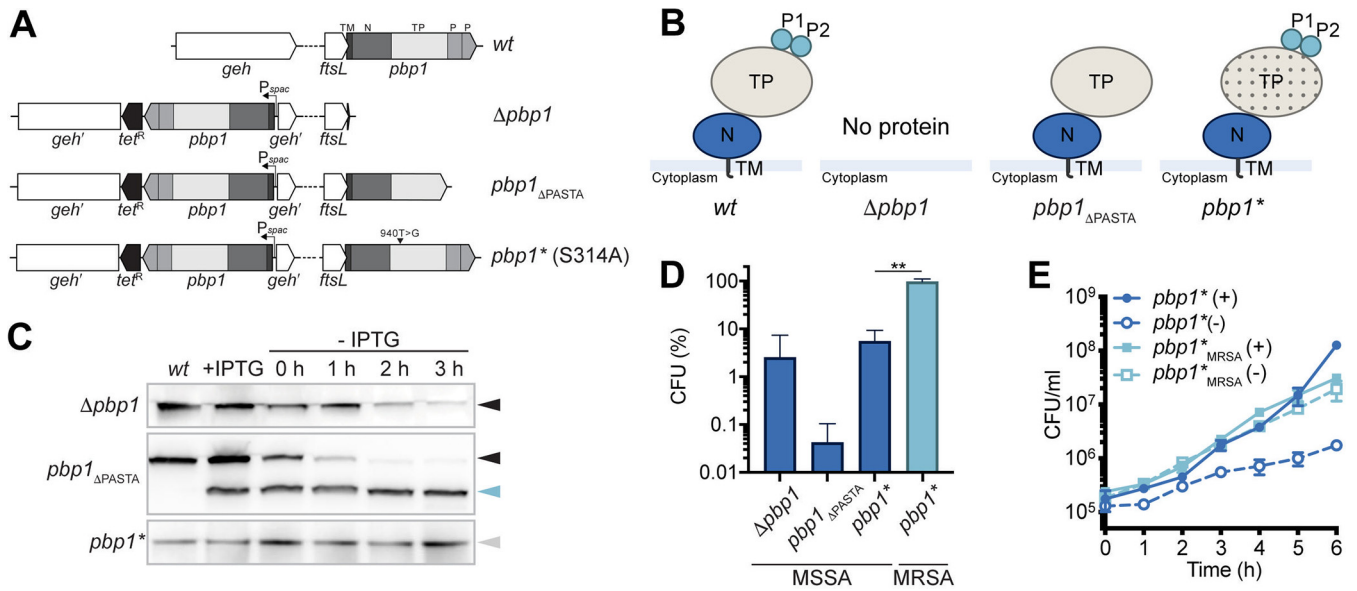


FIG 1 PBP1, its PASTA domains, and transpeptidase activity are essential in MSSA. (A) Schematic representation of genetic constructs used in this study. In *S. aureus* WT (*wt*) the 5' end of *pbp1* overlaps with the 3' end of *ftsL*. The *pbp1* gene encodes a protein containing the short cytoplasmic tail, transmembrane helix (TM), N-terminal pedestal domain (N), transpeptidase (TP) domain, and two PASTA domains (P1 and P2). In the mutants, an ectopic copy of *pbp1* is placed under the control of the *Pspac* promoter at the lipase (*geh*) locus, whereas the gene in the native *pbp1* locus is either deleted ($\Delta pbp1$), has P1 and P2 domains removed (*pbp1* _{Δ PASTA}), or has a point mutation which results in inactivation of the TP domain (*pbp1*^{*}). (B) Schematic representation of the domain architecture of PBP1 in *S. aureus* WT (*wt*) and PBP1 forms produced by $\Delta pbp1$, *pbp1* _{Δ PASTA}, and *pbp1*^{*} mutants in the absence of inducer. The TP domain inactivation is shown by dotted shading. (C) Immunoblot showing PBP1 levels in SH1000 *lacI* (*wt*) and in $\Delta pbp1$, *pbp1* _{Δ PASTA}, and *pbp1*^{*} grown with IPTG (+IPTG) and for 0, 1, 2, and 3 h without inducer (-IPTG) analyzed using anti-PBP1 antibody. Expected sizes: PBP1 and PBP1^{*} = 83 kDa (black and gray arrowheads, respectively) and PBP1 _{Δ PASTA} = 67 kDa (light blue arrowhead). (D) Plating efficiency of $\Delta pbp1$, *pbp1* _{Δ PASTA}, and *pbp1*^{*} (MSSA) cells and MRSA *pbp1*^{*} (*pbp1*^{*}_{MRSA}) cells upon inducer removal compared to the control groups grown in the presence of inducer. The *P* value was determined by Mann-Whitney *U* tests. *P* = 0.0043 (**, *P* < 0.01). Data represent the mean \pm standard deviation (SD). (E) Growth curves of *pbp1*^{*} (MSSA) and MRSA *pbp1*^{*} (*pbp1*^{*}_{MRSA}) in the presence (+) or absence (-) of IPTG. Data represent the mean \pm SD. Error bars that are smaller than the symbols are not shown. Data are representative of three (C and E) and at least four (D) independent experiments.

BocillinFL binding (28). These results indicate that the PASTA domains are essential for *S. aureus* growth and PBP1 functionality but not its stability.

During construction of the *pbp1*^{*} mutant we obtained, by serendipity, a *pbp1*_{STOP} mutant in which a single-nucleotide polymorphism (SNP) in the codon for Glu292 resulted in its replacement with a premature stop codon and the truncation of the entire TP and PASTAs region of PBP1 (Fig. S1E and F). However, immunoblot analysis using anti-PBP1 sera could not confirm the presence of the PBP1_{STOP} protein in the *pbp1*_{STOP} mutant (Fig. S1G), suggesting that stability of the N-terminal domain of PBP1 is dependent on its C terminus but not the PASTA domains. Although inactivation of PBP1 TP activity (PBP1^{*}) did not affect protein stability (Fig. 1C), it did remove the ability of PBP1 to bind BocillinFL (Fig. S1B). The loss of PBP1 TP activity resulted in severely compromised growth on solid medium (Fig. 1D and Fig. S1C) and reduced cellular viability in liquid culture (Fig. 1E and Fig. S1D). Thus, the TP activity of PBP1 is required for growth in the SH1000 background. Inactivation of the PBP1 TP activity was previously reported not to affect growth in the COL strain background (14). The differences in the necessity for the PBP1 TP activity could result from COL being MRSA, whereas SH1000 is a methicillin-sensitive *S. aureus* (MSSA) strain.

PBP1 TP activity is crucial in MSSA but not in MRSA. We have recently developed a set of defined strains where high-level β -lactam resistance of MRSA is mediated by *mecA* encoding PBP2A and a mutation in either *rpoB* or *rpoC* (25). This combination of genetic alterations (*mecA*⁺ *rpoB*) is present in COL (25). To test if the apparent disparity in PBP1's role is associated with MRSA, we developed a high-level resistant mutant of *pbp1*^{*} in the well-characterized *S. aureus* SH1000 by adding the *mecA rpoB*^{H929Q} to the MSSA *pbp1*^{*} mutant, resulting in SH1000_{MRSA} *pbp1*^{*} (Fig. S2A). Inactivating PBP1 TP did not affect the ability of SH1000_{MRSA} *pbp1*^{*} to grow in the absence of IPTG, whereas *pbp1* depletion led to growth inhibition in the isogenic $\Delta pbp1$ MSSA and MRSA strains

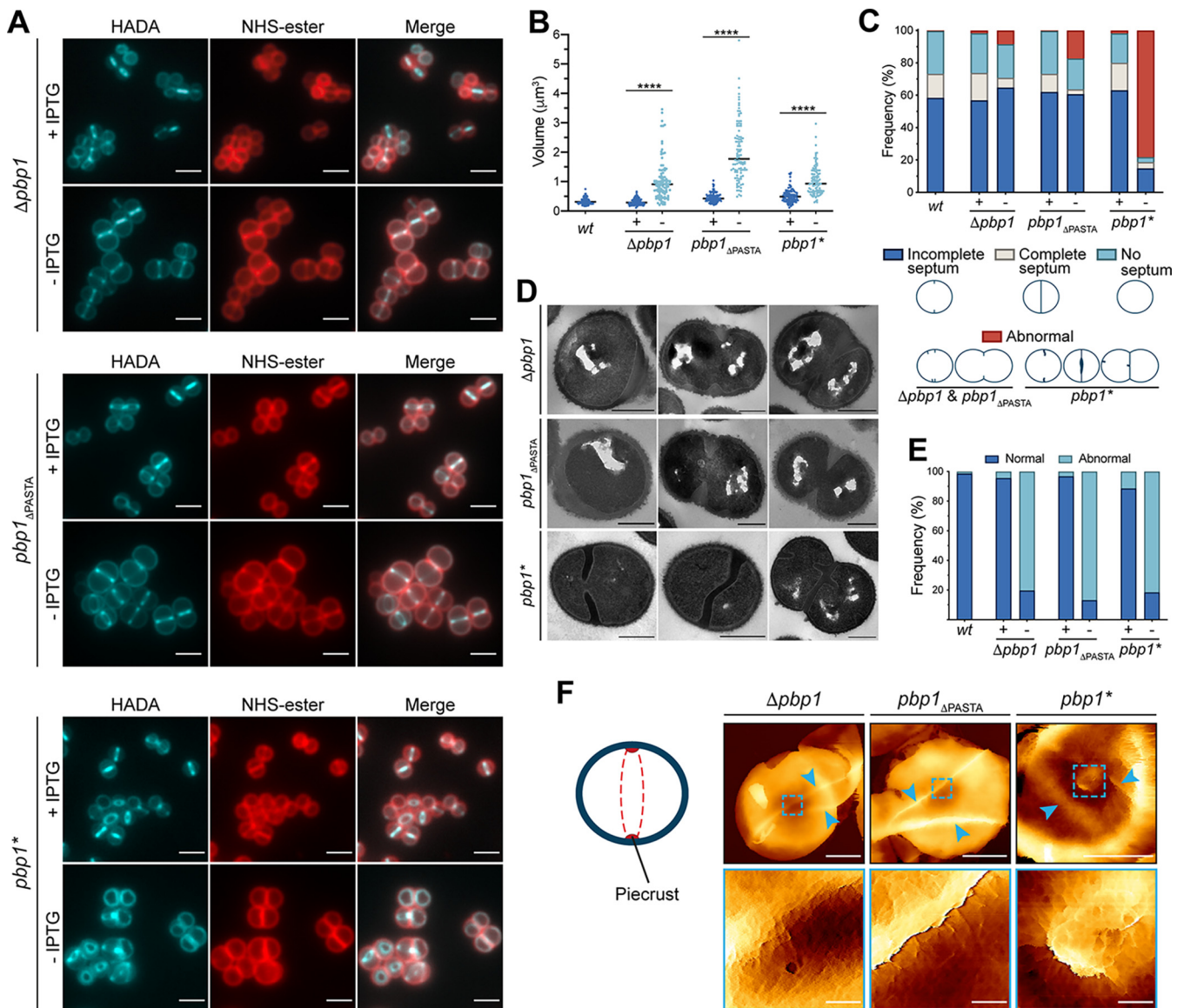


FIG 2 Role of PBP1 in cell division and PG synthesis in *S. aureus*. (A) $\Delta pbb1$, $pbb1_{\Delta PASTA}$ and $pbb1^*$ grown with or without IPTG for 2 h, incubated with HADA for 5 min to show nascent PG, and counterlabeled with NHS-ester Alexa Fluor 555 to image the cell wall. Images are average intensity projections of z stacks. Scale bars = 2 μm . (B) Cell volumes of SH1000 *lacI* (wt) and $\Delta pbb1$, $pbb1_{\Delta PASTA}$ and $pbb1^*$ grown with (+) or without (-) IPTG for 2 h as measured by fluorescence microscopy after NHS-ester Alexa Fluor 555 labeling. Each dot represents a single cell. The median of each distribution is indicated by a black line. The *P* value was determined by Mann-Whitney *U* tests (****, $P < 0.0001$). From left to right, $P = 3.033\text{e-}033$, $4.670\text{e-}049$, and $2.206\text{e-}022$. The number of cells analyzed for each mutant and condition was $n \geq 100$. (C) Quantification of cellular phenotypes for SH1000 *lacI* (wt) and $\Delta pbb1$, $pbb1_{\Delta PASTA}$ and $pbb1^*$ based on HADA incorporation (panel A) after incubation with (+) or without (-) IPTG for 2 h. From left to right, $n = 370$, 427 , 332 , 314 , 364 , 512 , and 331 . (D) TEM of $\Delta pbb1$, $pbb1_{\Delta PASTA}$ and $pbb1^*$ grown for 2 h in the absence of inducer. Scale bars = 500 nm. (E) Quantification of cellular phenotypes based on TEM data of SH1000 *lacI* (wt) and $\Delta pbb1$, $pbb1_{\Delta PASTA}$ and $pbb1^*$ grown for 2 h in the presence (+) or absence (-) of IPTG. Examples of cells classified as normal (blue) are shown in Fig. S4B. Cells with abnormal phenotypes (light blue) are shown in panel D and Fig. S4C. From left to right, $n = 304$, 391 , 329 , 314 , 377 , 263 , and 302 . (F) AFM topographic images of internal surface of purified sacculi from $\Delta pbb1$, $pbb1_{\Delta PASTA}$ and $pbb1^*$ grown in the absence of inducer for 2 h. (Left) Diagram of the inside of the cell before septal plate formation and (right) AFM images of sacculi (top images, scale bars = 500 nm, data scales [z]: 450, 300, and 100 nm from left to right, respectively) and higher magnification images (bottom images, scale bars = 50 nm, data scales [z]: 70, 100, and 50 nm from left to right, respectively) scanned within the boxed areas from the top images. The arrowheads indicate abnormal piecrusts. The boxed areas show details of piecrust features in $\Delta pbb1$ and $pbb1_{\Delta PASTA}$ and material agglomeration in $pbb1^*$. Data are representative of two (D to F) and (A to C) three independent experiments.

(Fig. 1D and E and Fig. S2B to D). Thus, the fundamental role of PBP1 in growth and division can only be studied in an MSSA background, as otherwise, the role of PBP1 can be confounded by the presence of the MRSA resistance apparatus.

PBP1 PASTA domains are required for septum progression. PG synthesis still occurred in $\Delta pbb1$, $pbb1_{\Delta PASTA}$ and $pbb1^*$ in the absence of IPTG, despite cell growth

inhibition, as measured by the incorporation of the fluorescent D-amino acid derivative HADA (Fig. 2A). This was not a consequence of the nonsynthesis, exchange reaction carried out by PBP4, as it occurred in *pbp4* as well as with the dipeptide ADA-DA (9, 29) (Fig. S3). All variants increased in cell volume upon depletion of *pbp1*, whereas *pbp1*_{ΔPASTA} was enlarged by almost twice as much as $\Delta pbp1$ and *pbp1*^{*} (Fig. 2A and B and Fig. S4A). Despite differences in cell size, both $\Delta pbp1$ and *pbp1*_{ΔPASTA} demonstrated a decrease in the proportion of cells with complete septa compared to the parent (Fig. 2A and C). Transmission electron microscopy (TEM) showed that more than 80% of the population had morphological defects, including cell wall thickening, PG blebs, and misshapen and/or multiple incomplete septa. (Fig. 2D and E and Fig. S4B and C). Such septa had abnormally thick bases and sharply pointed leading edges, suggesting that there is a problem with septal progression after initiation. Atomic force microscopy (AFM) previously revealed that the first step in cell division is the formation of a thick band of PG called the “piecrust” (30). Within this, the septal plate is formed, which has two PG architectures: disordered mesh facing the cell membrane and concentric rings in the septum core (5). Here, lack of PBP1 or the PBP1 PASTA domains led to formation of more than one, and often misplaced, piecrust. These mutations also caused an increase in unfinished septal annuli and alterations in the PG ring architecture (Fig. 2F and Fig. S5A to C, arrowheads), a feature that is revealed immediately after cell scission (5). Thus, depletion of PBP1 did not stop septum initiation, but the loss of the PASTA domains was enough to cause formation of irregular piecrusts, arrest septal plate formation, and lead to an altered septal PG architecture.

PBP1 TP activity is required for the characteristic septal PG architecture. The *pbp1*^{*} mutant gave a novel phenotype quite distinct from loss of entire PBP1 or PASTA domains. Inactivation of PBP1 TP activity did not prevent initiation and closing of the septa but, instead, resulted in accumulation of cells with aberrant septa and separation defects in about 80% of the population (Fig. 2A, C, and E). The septa in such cells had a rounded leading edge, were curved and abnormally thick (Fig. 2D and E and Fig. S4B and C), and had agglomerations of mesh-like material close to the septal center in addition to irregular piecrusts as observed by AFM (Fig. 2F and Fig. S5A and B). The intracellular agglomerations are PG, as they stain heavily with HADA and ADA-DA (Fig. 2A and Fig. S3C and F) and could be observed in purified sacculi (Fig. 2F and Fig. S5B). No ring architecture, only mesh-structured PG, could be observed on the surface of the *pbp1*^{*} mutant. Importantly, with fluorescence microscopy, the *pbp1*^{*} *pbp3* *pbp4* mutant, in which PBP2 is the only active TP, presented a similar phenotype upon IPTG removal as *pbp1*^{*}, exemplified by misshapen septa and agglomerations of PG material marked by HADA (Fig. S4D). Therefore, aberrant septal synthesis and progression occur in the *pbp1*^{*} mutant. Associated PG synthesis results from PBP2 transpeptidase activity and potentially the transglycosylase activity of FtsW, acting to produce un-cross-linked glycan strands.

The *pbp1*^{*} phenotype occurred specifically because of the loss of the TP activity of this essential enzyme. This phenotype is mirrored by the mode of action of β -lactam antibiotics, which bind to and inhibit the TP activity of PBPs (7). We have recently described the morphological effects of methicillin and oxacillin on *S. aureus*, which result in cell swelling and cessation of septal and peripheral cell wall synthesis (31). Our results suggest that PBP1 TP activity has a role in septal plate formation, and without this, the septum is misshapen. The conditional lethal strains made here allow for functional analysis of the genes concerned. However, phenotypes tend to accumulate on depletion of the wild-type protein over time, confusing the precise roles for individual components. To independently corroborate the role of the TP activity of PBP1, we utilized an approach to directly, and selectively, inhibit its activity. Meropenem (MEM) has a higher affinity for PBP1 than PBP2 (32, 33), and therefore, we hypothesized that its effect on *S. aureus* would match that of *pbp1*^{*}. In a MEM-titration, treatment with 1 \times MIC of MEM was sufficient to lead to cell death and a significant increase in SH1000 wild-type (WT) cell volume after 1 h (Fig. 3A and B and Fig. S6A). More than 70% of

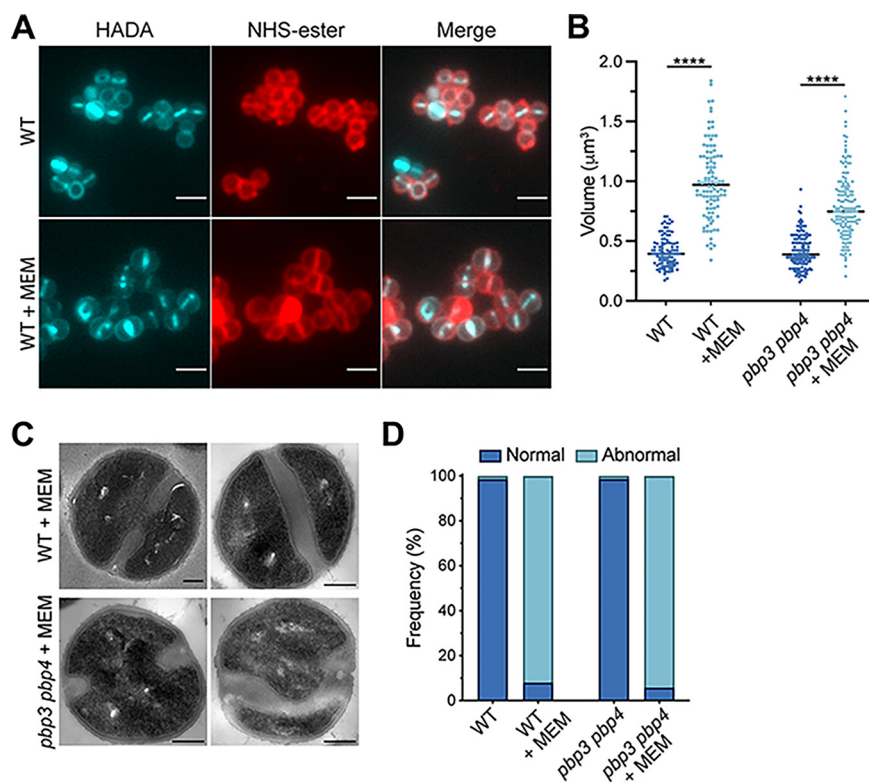


FIG 3 Effect of meropenem (MEM), an antibiotic with high affinity for PBP1, on *S. aureus*. (A) Fluorescence images of SH1000 WT treated with $1\times$ MIC of MEM for 1 h, labeled with HADA for 5 min to show nascent PG, and counterlabeled with NHS-ester Alexa Fluor 555 (cell wall). Images are average intensity projections of z stacks. Scale bars = $2\ \mu\text{m}$. (B) Cell volumes of SH1000 WT and *pbp3 pbp4* treated with $1\times$ MIC of MEM for 1 h as measured by fluorescence microscopy after NHS-ester Alexa Fluor 555 labeling (panel A). Each dot represents a single cell. The median of each distribution is indicated by a black line. The *P* value was determined by Mann-Whitney *U* tests (****, $P < 0.0001$). From left to right, $P = 1.276e-042$ and $1.303e-034$. The number of cells analyzed for each condition was $n \geq 100$. (C) TEM of SH1000 WT and *pbp3 pbp4* treated with $1\times$ MIC of MEM for 1 h. Scale bars = $200\ \text{nm}$. (D) Quantification of phenotypes of SH1000 WT and *pbp3 pbp4* treated with MEM ($1\times$ MIC) for 1 h based on TEM data (panel C and Fig. S6E and F). Examples of cells classified as normal (blue) are shown in Fig. S6E and F. Cells with abnormal phenotypes (light blue) are shown in panel C and Fig. S6E and F. From left to right, $n = 343, 287, 403,$ and 365 . Data are representative of two independent experiments.

MEM-treated cells had growth defects that manifested as aberrantly shaped septa and accumulation of PG as shown by HADA labeling (Fig. 3A, C, and D and Fig. S6C and E), similar to observations made with the *pbp1** mutants (Fig. 2A and C to E and Fig. S3C and F). The MEM phenotype of malformed septa was not linked to PBP3 or PBP4, as it was also observed in the corresponding double mutant (Fig. 3C and D and Fig. S6B, D, and F), which corroborated the role of PBP2 in misshapen septal genesis. The MEM phenotype differed from methicillin treatment, which inhibits both PBP1 and PBP2, as this results in a cessation of PG synthesis and apparent plasmolysis (31).

PASTA domains mediate PBP1 interaction with division-associated components.

The morphologies of the $\Delta pbp1$ and *pbp1* _{Δ PASTA} mutants resemble *S. aureus* depleted of DivIB in which EzrA and FtsZ form multiple rings and the synthesis of the cross wall is blocked, despite the normal recruitment of early cell division proteins and piecrust formation (34). EzrA is a marker of early division protein recruitment but also remains until septal completion (9, 35). In the $\Delta pbp1$ *ezrA-gfp* strain, EzrA, which here acts as an early cell division marker, was localized at midcell in the majority of cells and formed additional arcs or rings in 33% of the population (Fig. 4A and D). Multiple EzrA rings were observed in 43% of the *pbp1* _{Δ PASTA} *ezrA-gfp* mutant cells (Fig. 4B and D), supporting the requirement for PBP1 PASTA domains for correct selection of the division site and/or

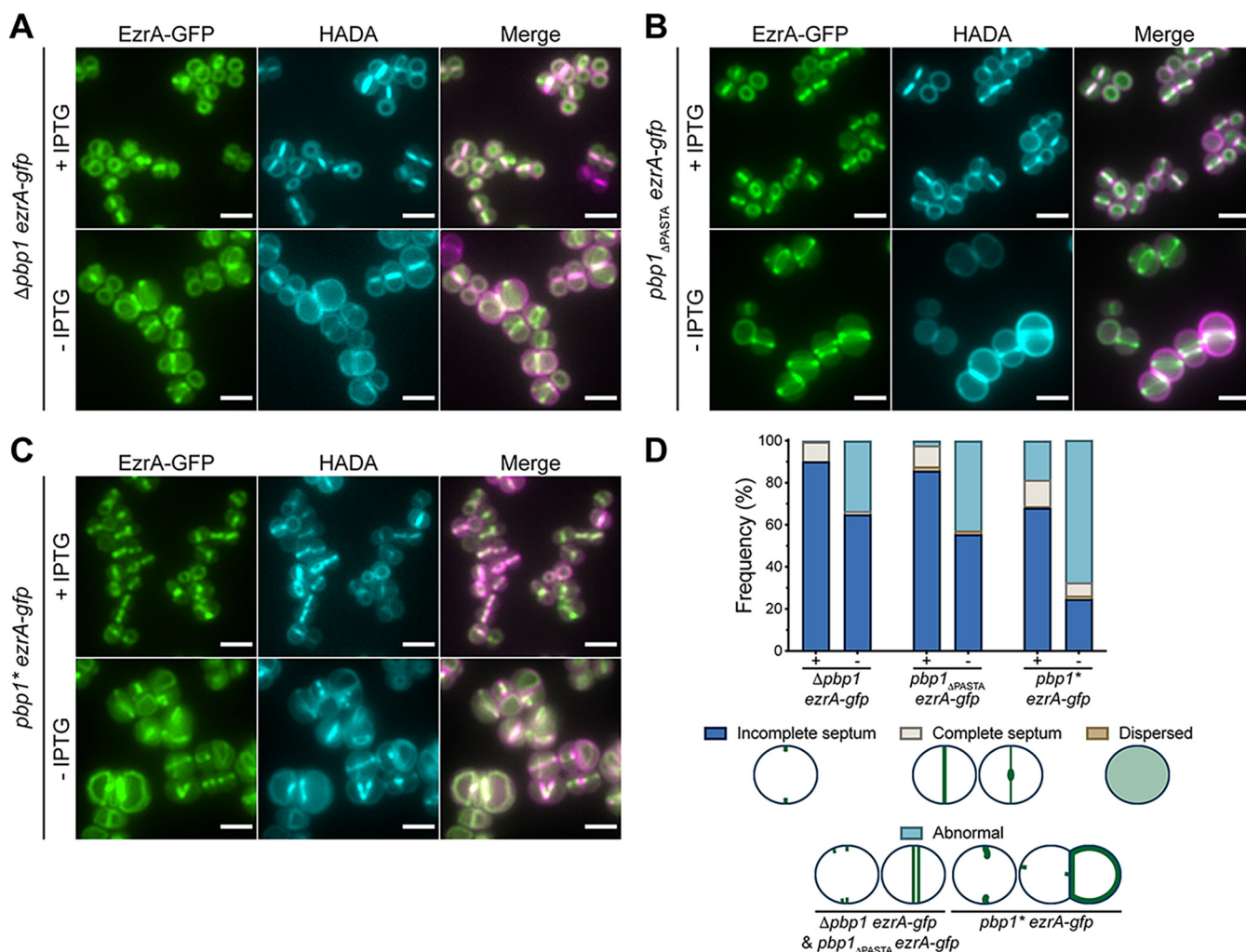


FIG 4 Role of PBP1, PASTA, and TP domains in EzrA localization in *S. aureus*. (A to C) Localization of EzrA-GFP in $\Delta pbp1$ *ezrA-gfp*, $pbp1_{\Delta PASTA}$ *ezrA-gfp*, and $pbp1^*$ *ezrA-gfp* grown in the presence or absence of IPTG for 2 h and labeled with HADA for 5 min to stain PG. Images are average intensity projections of z stacks. Scale bars = 5 μ m. (D) Quantification of EzrA-GFP localizations in $\Delta pbp1$ *ezrA-gfp*, $pbp1_{\Delta PASTA}$ *ezrA-gfp*, and $pbp1^*$ *ezrA-gfp* grown with or without IPTG. “Abnormal” includes those cells with multiple and/or misplaced EzrA rings. From left to right, $n = 395, 499, 481, 438, 360,$ and 382 . Data are representative of two independent experiments.

septal progression. Alternatively, the multiple division rings could result from a lack of the septal progression whereby the unproductive division machinery results in futile additional alternative initiation attempts, suggesting that PASTA domains are involved in the progression from piecruist to septal plate formation. While the number of cells with complete septa (EzrA-GFP visible as a line or focus) decreased by at least 6-fold in $\Delta pbp1$ *ezrA-gfp* and $pbp1_{\Delta PASTA}$ *ezrA-gfp*, it only halved in $pbp1^*$ *ezrA-gfp* (12.5% to 6.3% of $pbp1^*$ *ezrA-gfp* grown with or without IPTG, respectively; Fig. 4C and D), confirming that septum progression, although reduced, still occurred when PBP1 TP was inactive, implying that TP activity is necessary for correct septal architecture during cell division.

PBP1 is important for the septal surface PG ring structure (Fig. 2F and Fig. S5B), where it has been proposed that mature WTA is not present throughout the septum (23, 36). Loss of WTA also results in a proportion of cells with aberrant septa (21), suggesting a potential link with PBP1 function. Loss of *tarO* (leading to a lack of WTA) caused minor cell division defects in SH1000 (Fig. S7A, E, and F). Combining *tarO* with the mutations in *pbp1* exacerbated the observed morphological defects, with the appearance of distinct septal and off-septal PG foci appearing (marked with HADA) in $\Delta pbp1$ *tarO* and $pbp1_{\Delta PASTA}$ *tarO* (Fig. S7B to F), demonstrating that both WTA and PBP1 are involved in cell cycle progression in parallel.

As PBP1 PASTA has a role in the regulation of septal plate formation, this may be determined by interacting with other protein components. In order to examine this hypothesis, we performed a bacterial two-hybrid assay, in which PBP1 has previously been found to have apparent, multiple interactions (35). Truncation of the PASTA domains reduced *S. aureus* PBP1 interaction not only with DivIB but also with FtsW, while recognition of other known interacting partners of PBP1 (EzrA, PBP2, and DivIC) were unaffected by the PASTA truncation (Fig. S8A and B), suggesting that these potential, wider interactions involve the N-terminal domain of PBP1.

PBP1 PASTA domains bind peptidoglycan. Impaired interaction with DivIB could be one explanation for why cells depleted of PBP1 PASTA domains initiate irregular piecrusts and septation defects accrue as a consequence. PASTA domains have long been associated with PG binding because of work performed mainly on serine/threonine protein kinases (STPK) (26, 37–39). Very recently, PBP1 PASTA domains have been shown to bind isolated, small fragments of PG (27). Therefore, we assessed whether *S. aureus* PBP1 and its PASTA domains could recognize PG by measuring their affinities for *S. aureus* cell wall PG with or without WTA (\pm WTA) with a semiquantitative fluorescence-binding assay and *S. aureus* PBP1 derivatives produced in *Escherichia coli* (Fig. 5A and Fig. S8C). Cytochrome *c* (34) was used as a negative control to rule out nonspecific binding (dissociation constant [K_d], $1,126 \pm 37$ nM [+WTA] and $1,171 \pm 363$ nM [–WTA]) (Fig. S8D). Both SaPBP1 (K_d , 19 ± 4 nM [+WTA] and 115 ± 21 nM [–WTA]) and its PASTA domains (SaPASTA_{PBP1}; K_d , 198 ± 42 nM [+WTA] and 109 ± 23 nM [–WTA]) bound PG (Fig. 5B). Inactive SaPBP1* was still able to bind PG with a preference for PG with WTA present (K_d , 53 ± 8 nM [+WTA] and 227 ± 46 nM [–WTA]; Fig. 5B), similar to active SaPBP1. Although removal of the PASTA domains did not abolish BocillinFL binding (Fig. S8C), it considerably reduced the ability of SaPBP1 Δ PASTA to bind PG, and binding was abolished in the presence of WTA (K_d , $>2,000$ nM [+WTA] and 440 ± 57 nM [–WTA]; Fig. 5B). In contrast, the PASTA domains (SaPASTA_{PBP1}) on their own bind to *S. aureus* PG but are incapable of binding BocillinFL (Fig. 5B and Fig. S8C). These results demonstrate that PBP1 is a PG-binding protein, and the PASTA domains have a dominant role in this interaction. Sequence conservation analysis of PASTA domains revealed the presence of either Arg or Glu residues in classifying a PASTA domain as a PG-binder (40). The PASTA domains of *S. aureus* PBP1 each have proline at the equivalent positions (residues Pro603 and Pro661), and thus PBP1 would be predicted as a non-PG-binder. Our data suggest that the predicted significance of conserved Arg or Glu residues with regard to PG binding is either only relevant to PASTA domains found in STPKs, linear arrangements of tandem PASTA repeats, or is not suitable for proteins with multiple and complex functions like PBPs.

To gain a better understanding of the role of the PASTA domains in *S. aureus* PBP1 (SaPASTA_{PBP1}), we determined their structure by X-ray crystallography. Soluble recombinant protein was obtained in high yield from the cytoplasm of *E. coli* cells, and well-ordered crystals were subsequently produced that diffracted to a maximum resolution of 1.78 Å (Table S1). The structure was solved by molecular replacement using the corresponding PASTA domains present in SpPBP2x from PDB entry 5OAU (41), which shares 26% sequence identity with SaPASTA_{PBP1}. The asymmetric unit contains two monomers (labeled A and B), each forming a 2-layer sandwich comprising an α -helix and a three-stranded antiparallel β -sheet, distinct from the TP domain (Fig. 5C). Clear and continuous electron density allowed the modeling and unambiguous assignment of both PASTA domains (Fig. 5C). When SaPASTA_{PBP1} is compared with other structures deposited in the PDB using DALI (42), the top hit identified was *S. pneumoniae* PBP2x (Z-score, 15.7), showing a significant conservation of the PASTA fold despite low sequence identity (Fig. 5C). Unlike the linear arrangement observed for PASTA domains in serine/threonine kinases (43, 44), SaPASTA_{PBP1} adopts a compact upside-down globular arrangement (Fig. 5C). The arrangement of the two PASTA domains solved here, in isolation from the TP domain in comparison to structural analyses of SpPBP2x, is entirely consistent with a nonlinear PASTA domain arrangement. First, the structures of SaPASTA_{PBP1} and the PASTA domains

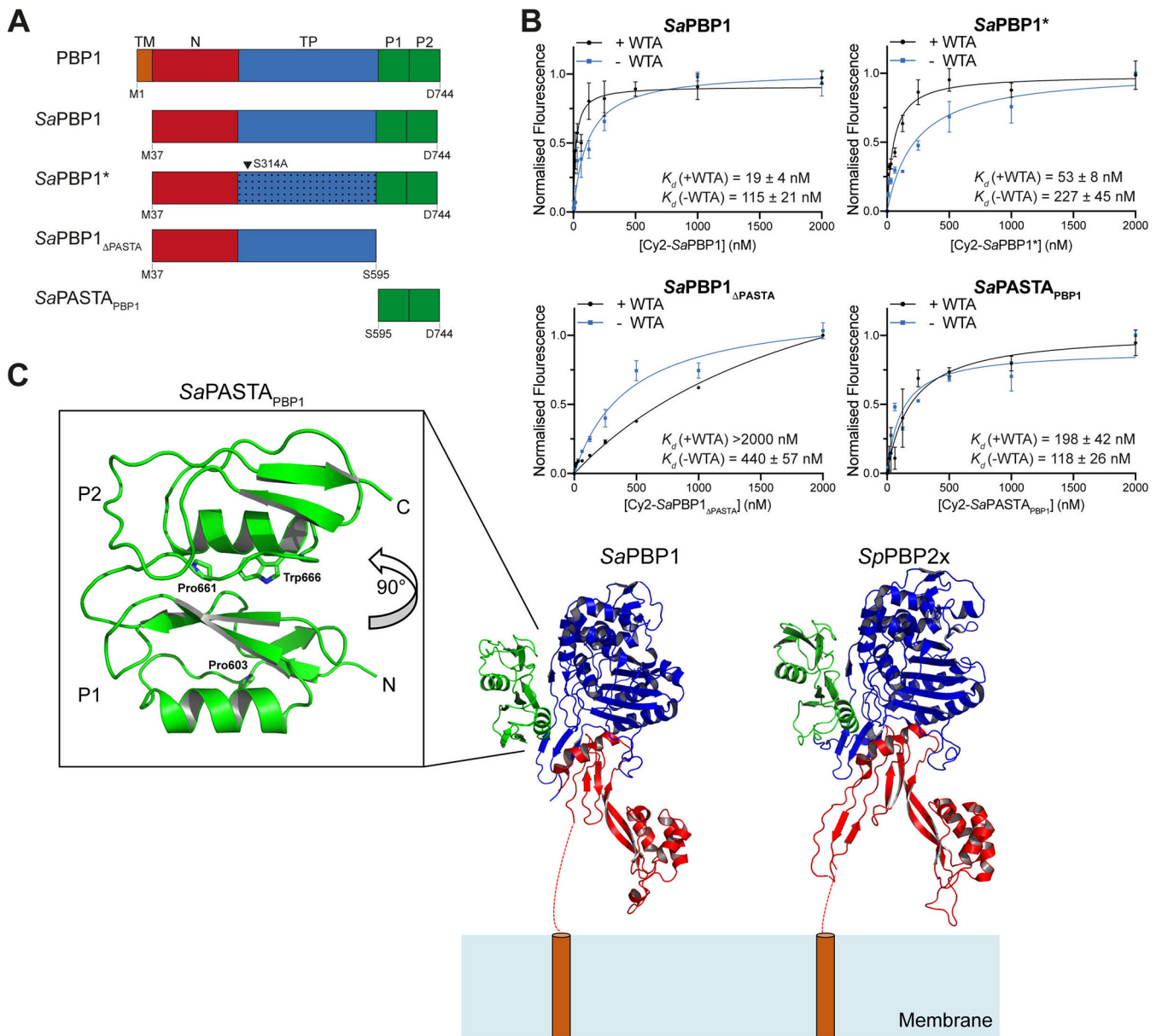


FIG 5 PBP1 PASTA domains bind cell walls. (A) Schematic representation of structural domain organization of *S. aureus* PBP1 (top) and recombinant proteins (*SaPBP1*, *SaPBP1*_{ΔPASTA}, *SaPBP1*^{*}, and *SaPASTA*_{PBP1}) used in this study. TM, short cytoplasmic fragment and transmembrane helix (orange); N, N-terminal pedestal domain (red); TP, transpeptidase domain (blue); P, PASTA domains (green). The arrowhead indicates the inactivation substitution in the TP domain of *SaPBP1*^{*}. The first and last amino acids of constructs are indicated. (B) Fluorescence cell wall sedimentation assay. A Wilcoxon signed rank test ($P < 0.05$) was carried out to assess the significance of difference in +WTA/-WTA PG binding: *SaPBP1*, $P = 0.0273$; *SaPBP1*^{*}, $P = 0.0078$; *SaPBP1*_{ΔPASTA}, $P = 0.0195$; *SaPASTA*_{PBP1}, $P \geq 0.9999$, not significant. Data represent the mean \pm SD. Error bars that are smaller than the symbols are not shown. (C) Structure of *SaPASTA*_{PBP1}. The crystal structure of *S. aureus* PBP1 lacking the PASTA domains (*SaPBP1*, PDB 5TRO) and our *SaPASTA*_{PBP1} structure were superimposed on *S. pneumoniae* PBP2x (*SpPBP2x*, PDB 5OAU) and displayed as cartoons. Their N termini are orientated close to the representative cell membrane, as if anchored there by their respective transmembrane helices (dashed red line/orange cylinder). Individual domains are colored as follows: N-terminal pedestal domain (red), transpeptidase domain (blue), and PASTA domain (green). The individual PASTA domains are labeled P1 and P2, respectively. Residues Pro603, Pro661, and Trp666 are displayed as sticks, with nitrogen atoms colored blue.

of *SpPBP2x* share a pairwise root mean square deviation (RMSD) of 2.2 Å over 114 C α , and when *SaPASTA*_{PBP1} is superimposed on the PASTA domains of *SpPBP2x*, there are no steric clashes with the TP domain. Second, the linker between PASTAs in *SaPASTA*_{PBP1} has a sequence of DGLTMPDMSGW, is neither glycine- nor alanine-rich, is not predicted to be disordered using the IUPred2 or ANCHOR2 web servers, and has a mean B factor of 44 Å² in comparison to a mean B factor of 42 Å² for the entire chain. Third, the interface between the PASTA domains is more reminiscent of the hydrophobic core of a globular

protein than the more polar interface observed between molecules in crystal packing. Finally, the two proline residues that apparently define PBP1 as a nonbinder of PG are found buried from solvent either at the interface of PASTA domain 1 with the TP domain (Pro603) or at the interface between the TP domain and PASTA domains 1 and 2 (Pro661). The latter interface includes the only tryptophan (Trp666) in the sequence of SaPASTA_{PBP1}; tryptophan residues are frequent markers of carbohydrate binding sites in proteins (45), and in the absence of any obvious grooves or surface features associated with conserved sequence distributions and/or electrostatics, it remains unclear how the PASTA domains of SaPBP1 recognize PG.

DISCUSSION

S. aureus has just two essential PBPs (46) and so forms an apparently simple system to understand cell wall growth and division. Even the transpeptidase activity of these two enzymes can be substituted by a single enzyme in the presence of β -lactam antibiotics via the acquisition of PBP2A, encoded by *mecA*, in MRSA strains. Our recent study has revealed that the presence of *mecA* and associated genetic lesions has a profound effect on *S. aureus*, even in the absence of antibiotics (25), leading to the discovery here that the PG biosynthetic activity of PBP1 is essential in MSSA but not in MRSA (Fig. 1D). This observation has important ramifications for many studies in *S. aureus*, where the use of an MRSA background can complicate phenotype interpretation. To understand the fundamental role of PBP1 activity in basic cell physiology, we have thus used an MSSA strain with a defined genetic background.

The essential function of PBP1 is associated with its crucial role in septal PG synthesis (14, 47). Here, we show that PBP1, in MSSA, has roles in both early and later stages of septum synthesis and can interact with other cell division components and make and bind to PG. PG binding is primarily mediated by the PASTA domains that are essential for cell division. There is clear overall structural similarity between *S. aureus* PBP1 and *S. pneumoniae* PBP2x PASTA domains in the way that the two tandem PASTA domains associate into an antiparallel bundle (Fig. 5C); this is in marked contrast to the head-to-tail linear PASTA domain repeats more typically found in STPKs. The highly hydrophobic interface between the two PASTA domains means it is unlikely to open up like butterfly wings to bind to PG; similarly, an extensive, linear interaction with PG, which is likely to occur with the head-to-tail PASTA domain arrangements seen in STPKs and which may require their dimerization (44), does not occur in SaPBP1. Despite the successful production of diffracting crystals of SaPASTA_{PBP1} grown in the presence of PG fragments (including an *N*-acetylglucosamine:*N*-acetylmuramic acid disaccharide), none of the structures yielded electron density features consistent with the stable binding of PG fragments. There are several potential explanations, including a lack of affinity of PASTA domains for small PG fragments, unrepresentative of the sacculus of *S. aureus*; our sedimentation assay does not permit the analysis of the binding of PASTA domains to small, soluble PG precursors. Consequently, and in common with all other PASTA domain structural analyses, the molecular details of PG recognition by SaPBP1 remain elusive. During the preparation of the manuscript, Martínez-Caballero et al. (27) published a crystal structure of the two PASTA domains of PBP1, also in the absence of endogenous ligand, which is indistinguishable (RMSD, 0.7 Å over 204 superimposed residues) from the structure that we report here. The same authors also solved structures of SaPBP1 in the presence and absence of β -lactams and pentaglycine (in which the PASTA domains were disordered). The latter structural analysis revealed that the pentaglycine substrate mimetic is not long enough to span between the transpeptidase active site and the PASTA domains, suggesting that the PG feature(s) recognized by the PASTA domain is/are chemically more complex than a simple short polypeptide.

S. aureus is a spheroid coccus that can divide successively in orthogonal planes (30, 48). Septation is first observed as the formation of a band of PG known as the piecrust (30). This then transitions to the production of the septal plate itself, an initially V-

shaped structure with a narrower leading edge (9). After closure of the septal annulus, the now bowed septum fills out to yield the mature structure prior to septal scission. The septal plate has two distinct PG architectures with a ring-like pattern at its core, which is exposed upon scission, and a subsequently synthesized fine mesh, akin to the rest of the peripheral cell wall (5). Loss of the entire PBP1, or just its PASTA domains, does not prevent piecrust formation but does result in multi- and/or off-center piecrusts without the ability to produce the septal plate (Fig. 2F). Thus, piecrust formation does not require PBP1 but is likely the result of the activity of the essential PBP2. PBP1 may regulate division site selection through PG cell wall recognition via its PASTA domains. Alternatively, as the division apparatus is unable to progress effectively to septal plate formation due to the lack of PBP1, this may lead to further rounds of initiation and piecrust formation. PBP1 has a clear role in septal plate formation where in the absence of PBP1 or the PASTA domains, cells form aberrantly shaped septa that do not close their annuli (Fig. 2A to E). In stark contrast, inactivation of PBP1 TP activity (*pbp1**) does not stop inward septum progression, as observed with loss of PBP1 or the PASTA domains. However, such septa are misshapen, curved, and abnormally thick (Fig. 2A to E and Fig. 3). The use of the PBP1-specific antibiotic MEM at $1 \times$ MIC led to the similar morphology of thickened and misshapen septa. Two independent avenues of research both led to the conclusion that PBP1 TP activity is essential for continued division and colony formation, and while septum formation is disturbed, it is not entirely prevented. Therefore, PBP1 retains its regulatory function(s) regardless of activity loss. Loss of PBP1 activity may result in futile glycan strand synthesis (49) by its partner transglycosylase FtsW (14) and/or the continued activity of PBP2, resulting in the observed aberrant septa and stasis. FtsW in *S. aureus* is essential and required for septum progression (14). In *Bacillus subtilis* the cell division-associated PBP2B is essential, but its enzyme activity is not and can be compensated for by PBP3 (50). Here, in the *pbp3 pbp4* background loss of PBP1 activity did not lead to death of the cells (Fig. S4E and F), suggesting that the nonessential enzymes do not support survival in the absence of PBP1 TP activity, whereas deletion of the PASTA domains leads to rapid death of the cells (Fig. 1D) due to loss of protein functionality not observed in the *pbp1**. Differences in plating efficiency and rate of loss of cellular viability between $\Delta pbp1$ and *pbp1* _{Δ PASTA} may reflect aberrant function of the truncated protein. As well as binding to the cell wall, PBP1 also apparently interacts with multiple protein partners, including EzcA, DivIB/C, PBP2, and FtsW (Fig. S8A and B) (14, 35). Recently, the PASTA domains from *B. subtilis* PBP2B were shown to regulate PBP2B interaction with DivIB (51). *S. aureus* DivIB is a PG-binding protein essential for division, the depletion of which leads to septal plate formation loss (34, 35). Here, the PBP1 PASTA domains were found to be involved in binding to DivIB and FtsW, alluding to their essential role in cell division. This could be a direct interaction, or loss of PBP1 PASTA may cause a conformational change in the remaining protein, as other's data suggest the FtsW-PBP1 interaction occurs via the PBP1 stalk domain (52). FtsW is a SEDS protein whose TG activity requires the presence of PBP1. Bifunctional aPBP (including PBP2) and bPBP-SEDS (including PBP1-FtsW) pairs share similar activities, but the fact that they coexist in many bacterial species implies there is a division of responsibilities between them. Indeed, it has been proposed lately that bPBP-SEDS pairs likely lay the primary PG matrix, while aPBPs support the initial PG by modifying, filling in, and adding PG to it (53, 54). The *S. aureus* septal plate PG has two distinct architectures, a disordered mesh present on its cytoplasm facing side and a ring structure at its core, which is revealed after the cells have split (5, 30) (Fig. 6). Recent AFM analysis from *Staphylococcus warneri* also describes the distinct PG architectures during septation as piecrust and septal plate rings/mesh (55). When sacculi are purified from *S. warneri*, the septum can split apart, revealing the rings, even in septa that have not closed their annulus, showing that the rings are not a likely result of PG hydrolysis during cell scission. We hypothesize (Fig. 6) that once the piecrust has been produced, PBP1 and FtsW use this as a foundation to initiate septal plate formation. Together they make the rings of material

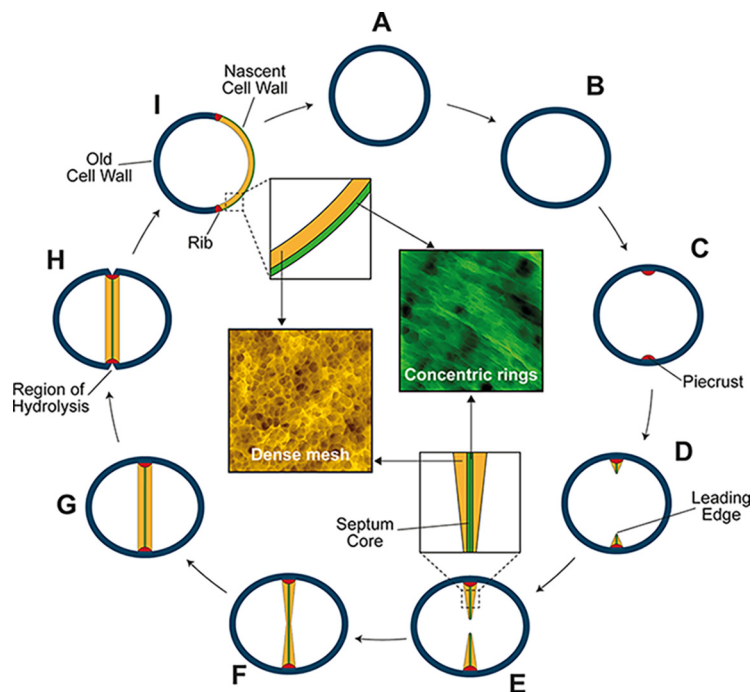


FIG 6 Conceptual model of septum formation in *S. aureus*, where PBP1 is required for septum formation and its characteristic ring-like PG architecture. (A and B) The growing *S. aureus* cell increases in volume (63). (C) Septal synthesis starts by formation of the piecrust (red) (30). This occurs as the result of the activity of PBP2 and forms the foundation for the septal plate. (D and E) The V-shaped septal plate (9) progresses inward by insertion of an initial, concentric ring-like structured PG synthesized by PBP1-FtsW at its core. Without PBP1 or PBP1 PASTA, the septal plate cannot be initiated, but in PBP1* it progresses but is aberrant. The PBP1-derived ring structure acts as a framework for the ensuing mesh-structured PG produced by PBP2. (F) The annulus closes, resulting in a bowed septum. (G) The septum is filled out by peptidoglycan insertion executed by PBP2, and this continues until the cross-wall is of uniform thickness (9). (H) The cell wall is hydrolyzed at the plane of septation. (I) Daughter cells separate. The cell wall of the daughter cell (colored insets) is a chimera of the old cell wall with both internally and externally mesh-structured PG and a nascent cell wall with the external ring-structured PG and the mesh-like cytoplasmic facing PG (5).

that become the core of the developing septum, providing the framework for PBP2 to make the bulk of the septal plate as a tight mesh alongside PBP4 and the insertion of WTA via the *tar* pathway. Loss of PBP1 TP activity in the presence of active PBP2 leads to the lack of the ring framework and aberrant, unproductive septum formation. The rings that form the center of the developing septum also provide the cleavage plane during scission.

Cell division is a fundamental requirement for life. A central question in bacteria is how is the division septum synthesized and then split to yield two daughter cells while maintaining cellular integrity in the face of internal turgor? Here, we have begun to answer this question by revealing the complex synthesis coordination mechanisms that allow this biological engineering feat to be accomplished.

MATERIALS AND METHODS

Bacterial growth conditions. The strains used in this study are listed in Table S1A.

All *Staphylococcus aureus* strains were grown in tryptic soy broth (TSB) containing appropriate antibiotics at 37°C, unless otherwise indicated, with aeration. All *Escherichia coli* strains, unless otherwise stated, were grown in Lysogeny broth (LB) containing appropriate antibiotics at temperatures ranging from 20°C to 37°C with aeration. For solid medium, 1.5% (wt/vol) agar was added. When necessary, growth medium was supplemented with kanamycin (50 $\mu\text{g mL}^{-1}$), tetracycline (1 $\mu\text{g mL}^{-1}$), chloramphenicol (10 $\mu\text{g mL}^{-1}$, *S. aureus*; 30 $\mu\text{g mL}^{-1}$, *E. coli*), erythromycin (5 $\mu\text{g mL}^{-1}$), spectinomycin (250 $\mu\text{g mL}^{-1}$), ampicillin (100 $\mu\text{g mL}^{-1}$), meropenem (0.4 $\mu\text{g mL}^{-1}$, 1 \times MIC for SH1000 WT; 0.2 $\mu\text{g mL}^{-1}$, 1 \times MIC for *pbp3 pbp4*), 5-bromo-4-chloro-3-indolyl β -D-thiogalactopyranoside (X-Gal; 80 $\mu\text{g mL}^{-1}$, *S. aureus*; 40 $\mu\text{g mL}^{-1}$, *E. coli*), or isopropyl β -D-thiogalactopyranoside (IPTG; 50 μM or 1 mM).

Plasmid construction. The plasmids and oligonucleotides used in this study are listed in Table S1 parts B and C, respectively.

Plasmids were cloned using *E. coli* NEB5 α following previously described methods (56, 57).

pKB-Pspac-pbp1. A fragment containing the ribosome-binding site (RBS) and coding region of *S. aureus* *pbp1* was PCR amplified from the genomic DNA of *S. aureus* SH1000 using pCQ-pbp1-F/-R primers and cloned into pCQ11-FtsZ-SNAP (9) cut with NheI and AclI by Gibson assembly, resulting in pCQ11-Pspac-pbp1. Next, the region containing *Pspac*, RBS, and *pbp1* was PCR amplified from pCQ11-Pspac-pbp1 using pKB-Pspac-pbp1-F/-R primers and cloned into BamHI and EcoRI cut pKASBAR (34) by Gibson assembly, giving pKB-Pspac-pbp1.

pMAD- Δ pbp1. Fragments encompassing 1-kb regions flanking upstream (from -980 bp upstream of *pbp1* to the first 20 bp of *pbp1*) and downstream (from 2,214 bp of *pbp1* to 970 bp downstream of *pbp1*) of *pbp1* were PCR amplified from *S. aureus* SH1000 genomic DNA using primer pairs *pbp1*-A/-B and *pbp1*-C/-D, respectively, and cloned into BamHI and EcoRI cut pMAD by Gibson assembly, creating deletion vector pMAD- Δ pbp1.

pMAD-pbp1 $_{\Delta$ PASTA}. Fragments encompassing 1.5-kb regions flanking the region encoding *pbp1* PASTA domains (upstream, from 286 bp to 1,785 bp of *pbp1*; downstream, from 2,214 bp of *pbp1* to 970 bp downstream of *pbp1*) were PCR amplified from *S. aureus* SH1000 genomic DNA using *pbp1*-E/-F and *pbp1*-G/-H primers and cloned into BamHI and EcoRI cut pMAD by Gibson assembly, resulting in deletion vector pMAD-pbp1 $_{\Delta$ PASTA}.

pMAD-pbp1*. An ~1.3-kb fragment covering an upstream region of the active site of *pbp1* (from -334 bp upstream of *pbp1* to the first 950 bp of the *pbp1* coding sequence) and an ~1.3-kb fragment comprising the 3' fragment of *pbp1* (930- to 2,235-bp region of *pbp1*) were PCR amplified from *S. aureus* SH1000 genomic DNA using primer pairs *pbp1**5'-F/-R and *pbp1**3'-F/-R, respectively. Primers *pbp1**5'-R and *pbp1**3'-F were designed to introduce a T to G point mutation resulting in a Ser314Ala substitution. The PCR products were ligated with pMAD cut with EcoRI and BamHI by Gibson assembly, resulting in pMAD-pbp1*.

T25-PBP1 $_{\Delta$ PASTA}. A fragment carrying *S. aureus* *pbp1* without the PASTA domains (M1-S595) was PCR amplified from *S. aureus* SH1000 genomic DNA using T25-pbp1-F and T25-pbp1pasta-R and cloned into BamHI and EcoRI cut pKT25, resulting in T25-PBP1 $_{\Delta$ PASTA}.

pVR plasmids. Full-length *pbp1* (M1-D744) was *E. coli* codon optimized, synthesized with GenScript, PCR amplified using VR47F/R, and cloned into KpnI and HindIII cut pOPINRSF using In-Fusion cloning (TaKaRa Bio), resulting in pVR01. Construction of pVR02 (*Sa*PBP1, M37-D744) and pVR06 (*Sa*PASTA $_{PBP1}$, S595-D744) was performed using inverse PCR (iPCR) (58), with pVR01 as a template and primer pairs VR49F/VR49R and VR57F/VR49R, respectively. pVR03 (*Sa*PBP1*, M37-D744, S314A) and pVR04 (*Sa*PBP1 $_{\Delta$ PASTA}, M37-S595) were constructed with QuikChange site-directed mutagenesis of pVR02 using VR51 and VR53, respectively.

pSA50. In order to construct an overexpression plasmid for sPBP1-BAP, the A51-D744 fragment of *E. coli* codon optimized *pbp1* was PCR amplified using primers OPPF20018F/OPPF20018R and cloned into KpnI and SfoI cut pOPINJB by In-Fusion cloning (TaKaRa Bio). The resulting construct, pSA50 contains an N-terminal hexahistidine tag fused to glutathione-S-transferase followed by a human rhinovirus 3C protease site, while the PBP1 (A51-D744) C-terminal end is fused to a biotin acceptor peptide (BAP) sequence.

Construction of *S. aureus* mutants. All vectors were passed through a restriction-deficient *S. aureus* RN4220 before being transduced into a final *S. aureus* SH1000 strain. Transformation and phage transduction of *S. aureus* were carried out as described previously (59, 60).

Δ pbp1, *pbp1* $_{\Delta$ PASTA} and *pbp1**. For construction of *pbp1* mutation strains, first, an ectopic copy of *pbp1* under the control of the *Pspac* promoter was introduced at the lipase (*geh*) locus. Electrocompetent CVL316 was transformed with pKB-Pspac-pbp1. The chromosomal fragment containing the integrated plasmid was moved into *S. aureus* SH1000 by phage transduction, resulting in SJF4588 (*S. aureus* SH1000 *geh*::*Pspac-pbp1*). Next, electrocompetent RN4220 was transformed with pMAD- Δ pbp1, pMAD-pbp1 $_{\Delta$ PASTA}, or pMAD-pbp1*, and the plasmids were moved to SJF4588 by phage transduction. Integration at 42°C and excision at 28°C of pMAD- Δ pbp1, pMAD-pbp1 $_{\Delta$ PASTA} or pMAD-pbp1* resulted in strains SJF5116, SJF5275, and SJF4590, respectively. To allow controlled expression of *pbp1* from *Pspac*, pGL485, a multicopy plasmid carrying *lacI* was introduced, creating strains Δ pbp1 (*S. aureus* SH1000 *geh*::*Pspac-pbp1* Δ pbp1 *lacI*), *pbp1* $_{\Delta$ PASTA} (*S. aureus* SH1000 *geh*::*Pspac-pbp1* *pbp1* $_{\Delta$ PASTA} *lacI*), and *pbp1** (*S. aureus* SH1000 *geh*::*Pspac-pbp1* *pbp1** *lacI*). On all occasions, consistent colony size and growth kinetics were monitored to prevent the selection of suppressor mutations.

MRSA Δ pbp1 and MRSA *pbp1**. In order to construct high-level β -lactam-resistant mutants, Δ pbp1 and *pbp1** were transformed with a phage lysate from SJF5046 (*S. aureus* SH1000 *lysA*::*pmecA* *rpoB*^{H929Q}) with selection for erythromycin resistance, resulting in low-level β -lactam-resistant Δ pbp1 *pmecA* and *pbp1** *pmecA*. The low-level-resistant mutants were transduced again with the phage lysate from SJF5046 and selected for kanamycin resistance, resulting in MRSA Δ pbp1 (*S. aureus* SH1000 *geh*::*Pspac-pbp1* Δ pbp1 *lacI* *lysA*::*pmecA* *rpoB*^{H929Q}) and MRSA *pbp1** *pbp1* (*S. aureus* SH1000 *geh*::*Pspac-pbp1* *pbp1** *lacI* *lysA*::*pmecA* *rpoB*^{H929Q}). MIC values were determined using antibiotic susceptibility tests using Etest MIC evaluator (Oxoid) strips.

pbp3 pbp4. SH1000 was transduced with a phage lysate from NE420 (*S. aureus* JE2 *pbp3*::*Tn*), resulting in SH4421 (*S. aureus* SH1000 *pbp3*::*Tn*). To swap the erythromycin resistance cassette to a kanamycin cassette, SH4425 (*S. aureus* SH1000 *pbp4*::*Tn*) was transduced with a phage lysate from NE3004 (*S. aureus* RN4220 pKAN). Integration at 42°C and excision at 28°C of pKAN resulted in strain SH5115 (*S. aureus*

SH1000 *pbp4::kan*). SH4421 was subsequently transduced with a phage lysate from SH5115 (*S. aureus* SH1000 *pbp4::Tn*), resulting in *pbp3 pbp4* (SH5483; *S. aureus* SH1000 *pbp3::Tn pbp4::kan*).

Δ pbp1 *pbp4*, *pbp1* _{Δ PASTA} *pbp4*, and *pbp1 *pbp4*.** Δ *pbp1*, *pbp1* _{Δ PASTA} and *pbp1** were transduced with a phage lysate from SH5115 (*S. aureus* SH1000 *pbp4::kan*), resulting in Δ *pbp1 pbp4* (*S. aureus* SH1000 *geh::Pspac-pbp1* Δ *pbp1 lacl pbp4::kan*), *pbp1* _{Δ PASTA} *pbp4* (*S. aureus* SH1000 *geh::Pspac-pbp1 pbp1* _{Δ PASTA} *lacl pbp4::kan*), and *pbp1** *pbp4* (*S. aureus* SH1000 *geh::Pspac-pbp1 pbp1** *lacl pbp4::kan*), respectively.

***pbp1** *pbp3 pbp4*.** *pbp1** *pbp4* (*S. aureus* SH1000 *geh::Pspac-pbp1 pbp1** *lacl pbp4::kan*) was transduced with a phage lysate from SH4421 (*S. aureus* SH1000 *pbp3::Tn*), resulting in *pbp1** *pbp3 pbp4* (*S. aureus* SH1000 *geh::Pspac-pbp1 pbp1** *lacl pbp3::Tn pbp4::kan*).

Δ pbp1 *ezrA-gfp*, *pbp1* _{Δ PASTA} *ezrA-gfp*, and *pbp1 *ezrA-gfp*.** Δ *pbp1*, *pbp1* _{Δ PASTA} and *pbp1** were transduced with a phage lysate from JGL227 (*S. aureus* SH1000 *ezrA-gfp*+) (35), resulting in Δ *pbp1 ezrA-gfp* (*S. aureus* SH1000 *geh::Pspac-pbp1* Δ *pbp1 lacl ezrA-gfp*), *pbp1* _{Δ PASTA} *ezrA-gfp* (*S. aureus* SH1000 *geh::Pspac-pbp1 pbp1* _{Δ PASTA} *lacl ezrA-gfp*), and *pbp1** *ezrA-gfp* (*S. aureus* SH1000 *geh::Pspac-pbp1 pbp1** *lacl ezrA-gfp*), respectively.

Δ pbp1 *tarO*, *pbp1* _{Δ PASTA} *tarO*, and *pbp1 *tarO*.** Δ *pbp1*, *pbp1* _{Δ PASTA} and *pbp1** were transduced with a phage lysate from *tarO* (*S. aureus* SA113 Δ *tarO::ery pUC1-tarO*) (31), resulting in Δ *pbp1 tarO* (*S. aureus* SH1000 *geh::Pspac-pbp1* Δ *pbp1 lacl* Δ *tarO::ery*), *pbp1* _{Δ PASTA} *tarO* (*S. aureus* SH1000 *geh::Pspac-pbp1 pbp1* _{Δ PASTA} *lacl* Δ *tarO::ery*), and *pbp1** *tarO* (*S. aureus* SH1000 *geh::Pspac-pbp1 pbp1** *lacl* Δ *tarO::ery*), respectively.

PBP1 depletion. *Pspac-pbp1* strains were grown from an optical density at 600 nm (OD₆₀₀) of 0.1 to the exponential phase (OD₆₀₀ ~0.5) in TSB containing 10 μ g mL⁻¹ chloramphenicol and 50 μ M IPTG. Cells were washed three times by centrifugation and resuspension in TSB. Washed cells were then used to inoculate TSB containing 10 μ g mL⁻¹ chloramphenicol. Cultures were inoculated to an OD₆₀₀ of 0.05 for phenotypic studies and an OD₆₀₀ of 0.005 for growth studies. For phenotypic analysis, cultures were incubated for 2 h to allow depletion of PBP1 before microscopy imaging. Control samples were grown in TSB supplemented with 10 μ g mL⁻¹ chloramphenicol and 1 mM (50 μ M, *ezrA-gfp* mutants) IPTG.

For the plating efficiency test, cells grown in the presence of 10 μ g mL⁻¹ chloramphenicol and 50 μ M IPTG to the exponential phase (OD₆₀₀ ~0.5) were washed three times in phosphate-buffered saline (PBS). Serial dilutions of washed cells were plated on TSB containing 10 μ g mL⁻¹ chloramphenicol, with or without 1 mM IPTG. Relative plating efficiency (% CFU) is expressed as the number of cells that grow on plates without IPTG (CFU_{no IPTG}) to cells that grow in the presence of IPTG (CFU_{IPTG}) multiplied by 100%:

$$\%CFU = \frac{CFU_{no\ IPTG}}{CFU_{IPTG}} \times 100\%$$

Meropenem activity assays. *S. aureus* strains were grown overnight in TSB. The overnight cultures were used to inoculate fresh TSB medium to an OD₆₀₀ of 0.05. When cells reached an OD₆₀₀ of 0.2 to 0.4, meropenem was added, and the change of bacterial count was monitored. The CFU per mL of culture measures were normalized to the initial CFU/mL at the time of the antibiotic addition, at time zero (*t*₀).

$$Relative\ CFU/mL = \frac{CFU/mL_{t_n}}{CFU/mL_{t_0}}$$

For phenotypic analysis, cells were treated for 1 h with 1 \times MIC meropenem before microscopy imaging.

Fractionation of *S. aureus* membranes. The membrane fraction of *S. aureus* was prepared as previously described (61) with the following modifications. *S. aureus* cells grown to the appropriate growth phase were recovered by centrifugation (5,000 \times *g*, 10 min, 4°C) and washed three times by resuspension and centrifugation (5,000 \times *g*, 10 min, 4°C) in PBS. Cells were resuspended in 50 mM Tris, 100 mM NaCl, pH 8.0 containing Complete Protease Inhibitor (Roche) and broken using 0.1-mm silica spheres (lysing matrix B) and FastPrep homogenizer (MP Biomedicals) in 12 cycles of 30 s, at maximum speed (6.5 m s⁻¹), with 5 min of incubation on ice between cycles. Cell lysates were centrifuged (8,000 \times *g*, 10 min, 4°C) to remove unbroken cells. The supernatant was then spun (8,000 \times *g*, 10 min, 4°C) to sediment cell wall material. The membrane fraction was recovered from the supernatant by centrifugation (35,000 \times *g*, 20 min, 4°C), and the pellet (membranes) was resuspended in PBS. The total protein concentration was estimated by Bradford assay.

In vitro labeling of *S. aureus* PBPs with BocillinFL. This method was adopted from a published protocol (62) with minor modifications. Membrane proteome samples (25 μ g in 20 μ L PBS) and purified proteins (2.5 μ g in 25 μ L HEPES pH 7.5 150 mM NaCl) were incubated with 25 μ M BocillinFL (Invitrogen) for 20 min at 37°C. Additionally, for the competition assay, purified 5aPBP1 was mixed with 2.5 μ g (~286 μ M final concentration) ampicillin and incubated at 37°C for 10 min prior to the addition of BocillinFL. The reaction was stopped by the addition of 5 \times SDS-PAGE loading buffer. Membrane proteome was additionally incubated for 10 min at 90°C. The samples were run on a 6 to 20% (wt/vol) SDS-PAGE gradient or 10% (wt/vol) SDS-PAGE gel and visualized using a Bio-Rad ChemiDoc MP imaging system or a GE Typhoon FLA 9500.

Labeling *S. aureus* D-amino acids. *S. aureus* cells were incubated with 500 μ M (2 mM for *pbp4* mutants) HADA or 1 mM ADA-DA at 37°C for 5 min. Cells were then washed by centrifugation and resuspension in PBS.

Click chemistry. ADA-DA containing an azide functional group was fluorescently labeled with Atto 488 alkyne at $5 \mu\text{g mL}^{-1}$ via the click reaction [copper (I)-catalyzed alkyne-azide cycloaddition]. This was carried out using the Click-iT cell reaction buffer kit (Thermo Fisher) according to the manufacturer's protocol.

Labeling *S. aureus* with fluorescent NHS-ester. Fixed cell wells were resuspended in PBS containing $8 \mu\text{g mL}^{-1}$ Alexa Fluor 555 NHS-ester (Invitrogen) and incubated at room temperature for 30 min. Cells were washed twice by centrifugation and resuspension in PBS.

Fixing for fluorescence microscopy. Cells were fixed by incubation in 1.6% (wt/vol) paraformaldehyde at room temperature for 30 min.

Fluorescence microscopy. Fixed cells were dried onto a poly L-lysine-coated slide, mounted in PBS, and imaged on a Nikon Ti inverted microscope fitted with a Lumencor Spectra X light engine. Images were taken using a $100\times$ PlanApo (1.4 NA) oil objective using 1.518 RI oil and detected by an Andor Zyla sCMOS camera.

Cell volume estimation. Cell volume calculations were carried out as previously described (63). The long and short axes of cells were measured using Fiji. The volume was then calculated based on a prolate spheroid shape with volume

$$V = 4/3\pi ab^2,$$

where a and b are the radii along the long and short axes, respectively.

Transmission electron microscopy. *S. aureus* strains were prepared for electron microscopy as previously described (64).

Preparation of *S. aureus* sacculi. Peptidoglycan from *S. aureus* cells was extracted and, if required, hydrofluoric acid (HF)-treated to remove cell wall accessory polymers as previously described (64).

Sacculi immobilization for AFM imaging. The immobilization surface was prepared by adding the solution mixed with $171 \mu\text{L}$ of 100 mM NaHCO_3 , $3 \mu\text{L}$ of 1 M NaOH, and $6 \mu\text{L}$ of Cell-Tak (Corning, 5% [wt/vol] in acetic acid) on freshly cleaved mica. After 30 min of incubation, the surface was washed with $5 \times 200 \mu\text{L}$ high-pressure liquid chromatography (HPLC)-grade water. Sacculi stocks were 10 times diluted in HPLC-grade water and briefly tip-sonicated to resuspend them prior to immobilization. Then, $10 \mu\text{L}$ of the sacculi suspension was added to $40 \mu\text{L}$ of HPLC-grade water on the Cell-Tak immobilization surface and incubated for 1 h. The surface was then thoroughly rinsed with HPLC-grade water, blow-dried with nitrogen, and stored in a petri dish at room temperature before AFM imaging.

AFM Imaging and image analysis. AFM imaging was carried out on a Nanowizard III ULTRA Speed system (JPK, Germany). Rectangular cantilevers with a nominal spring constant of 0.3 N/m and resonant frequency (in liquid) of ~ 150 kHz (USC-F0.3-k0.3; NanoWorld, Switzerland) were used. The spring constant and deflection sensitivity of each cantilever were calibrated prior to each measurement (65, 66) Measurements were carried out in quantitative imaging mode at room temperature in the buffer composed of 200 mM KCl and 10 mM Tris. Scans were driven at a line rate of ~ 0.78 Hz, with a typical Z length of 300 nm and trigger force of 20 nN. The resultant topographic images were processed using JPK Data Processing. No flattening or surface subtraction was applied. A high-pass filter (scale: 100% to 500%, degree of smoothing: 5 px, horizontal) was applied to the higher magnification images to enhance the contrast without modifying the morphological features. The morphological features of sacculi were summarized from images obtained on abundant technical repeats of 2 biological replicates.

Recombinant protein production and purification. sPBP1-BAP. *E. coli* BL21(DE3) cells containing plasmid pSA50 were grown in LB medium supplemented with $100 \mu\text{g mL}^{-1}$ ampicillin at 37°C to an OD_{578} of 0.5. Protein overproduction was induced by addition of 0.5 mM IPTG to the cell culture and further incubation for 4 h at 30°C . Cells were harvested by centrifugation ($6,200 \times g$, 15 min, 4°C), and the pellet was resuspended in basic buffer (25 mM Tris-HCl, 100 mM NaCl, pH 7.5). After addition of 1 mM phenylmethylsulfonyl fluoride (PMSF), a 1:1,000 dilution of protease inhibitor cocktail (Sigma-Aldrich), and DNase, the cells were disrupted by sonication (Branson digital sonifier). The cell lysate was centrifuged ($130,000 \times g$, 60 min, 4°C), and the supernatant was recovered. The supernatant was incubated with Ni-NTA Superflow (Qiagen) for 2 h at 4°C with gentle stirring, which had been preequilibrated in basic buffer. The resin was poured into a gravity column and washed with 20 volumes of wash buffer (25 mM Tris-HCl, 150 mM NaCl, 10% [vol/vol] glycerol, 10 mM MgCl_2 , 20 mM imidazole, pH 7.5). Bound protein was eluted with elution buffer (25 mM Tris-HCl, 150 mM NaCl, 10% [vol/vol] glycerol, 10 mM MgCl_2 , 600 mM imidazole, pH 7.5). Next, 10 U mL^{-1} of HRV-3C protease (TaKaRa) was added to the Ni-NTA-eluted protein to remove the oligohistidine-GST tag during dialysis against 3 L of dialysis buffer I (25 mM Tris-HCl, 150 mM NaCl, 10 mM EGTA, 10% [vol/vol] glycerol, pH 7.5) for 20 h at 4°C . Digested protein was dialyzed against 3 L of dialysis buffer II (25 mM Tris-HCl, 150 mM NaCl, 10 mM MgCl_2 , 10% [vol/vol] glycerol, pH 7.5), for 3 h at 4°C . The protein was incubated in the same Ni-NTA beads (preequilibrated in dialysis buffer II) for 2 h at 4°C to remove the contaminants and the His-GST tag from the sample. The flowthrough and the washes (2 volumes of wash buffer) were pooled, dialyzed against storage buffer (25 mM HEPES-NaOH, 150 mM NaCl, 10 mM MgCl_2 , 10% [vol/vol] glycerol, pH 7.5), and concentrated using a Vivaspinn Turbo 15 column (molecular weight cutoff [MWCO] of 50,000 Da).

SaPBP1, SaPBP1*, SaPBP1 $_{\Delta\text{PASTA}}$, and SaPASTA $_{\text{PBP1}}$. All recombinant proteins were produced in *E. coli* Rosetta (DE3) cells at 37°C in Terrific Broth (TB) medium supplemented with $50 \mu\text{g mL}^{-1}$ kanamycin and $30 \mu\text{g mL}^{-1}$ chloramphenicol. Once cultures had reached an OD_{600} of 0.9, protein expression was induced with 1 mM IPTG for 20 h at 20°C . Cells were harvested by centrifugation ($4,000 \times g$ at 4°C for 30 min), and the pellet was resuspended in a buffer of 50 mM Tris-HCl, pH 8.0, 500 mM NaCl, 20 mM imidazole supplemented with one EDTA-free protease inhibitor cocktail tablet (Roche), and DNase ($4 \mu\text{g mL}^{-1}$

final concentration). The cells in this resuspension were lysed by two passes through a One-Shot cell disruptor (Constant Systems) at 23 kilopounds per square inch (kpsi), and the cell debris was removed by centrifugation ($40,000 \times g$ at 4°C for 30 min). The first purification step was affinity chromatography with a 5-mL HisTrap FF column (GE Healthcare) precharged with Ni^{2+} and equilibrated in buffer A (50 mM Tris-HCl, pH 8.0, 500 mM NaCl, 20 mM imidazole). A linear concentration gradient of imidazole was applied to elute the protein using buffer B (50 mM Tris-HCl, pH 8.0, 500 mM NaCl, 800 mM imidazole). Further purification was carried out by size exclusion chromatography using a Superdex 200 Hi Load 16/60 column (GE Healthcare). Proteins were eluted with SEC buffer (25 mM Tris-HCl, pH 8.0, 150 mM NaCl) and analyzed by SDS-PAGE.

Generation of anti-PBP1 antibody. Serum against sPBP1A-BAP was produced from rabbits following a 28-day immunization program at Eurogentec (Belgium), and it was purified as previously described (67).

Immunoblot analysis. *S. aureus* cultures were washed three times by resuspension and centrifugation ($5,000 \times g$, 10 min, 4°C) in PBS. Cells were resuspended in TBSI (50 mM Tris, 100 mM NaCl, pH 8.0, plus Complete Protease Inhibitor Cocktail [Roche]) and broken using 0.1 mm silica spheres (lysing matrix B) and FastPrep homogenizer (MP Biomedicals) in 12 cycles of 30 s, at maximum speed (6.5 m s^{-1}), with a 5-min incubation on ice between cycles. Cell lysates were centrifuged ($5,000 \times g$, 10 min, 4°C) to remove unbroken cells. Approximately 60 μg of total protein was separated on a 12% (wt/vol) SDS-PAGE gel and electroblotted onto a nitrocellulose membrane and blocked in 5% (wt/vol) skimmed milk in TBST (20 mM Tris-HCl, pH 7.6, 17 mM NaCl, 0.1% [vol/vol] Tween 20). The membrane was incubated with primary polyclonal anti-PBP1 (1:1,000) overnight with gentle agitation at 4°C . Primary antibodies were detected using horseradish peroxidase-conjugated goat anti-rabbit IgG (1:10,000; Bio-Rad) and Clarity Western enhanced chemiluminescence (ECL) substrate (Bio-Rad) reagent according to the manufacturer's protocol. Chemiluminescence was detected using a Syngene G:BOX Chemi XX9 system.

Crystallization, data collection, and structure determination. Crystallization of SaPASTA_{PBP1} was carried out at 20°C by the sitting-drop vapor diffusion method in 96-well MRC plates (Molecular Dimensions) with a Mosquito crystallization robot (TTP LabTech) and commercial crystallization screens (Hampton Research and Molecular Dimensions). Orthorhombic crystals of diffraction quality, with a maximum dimension of approximately 500 μm , appeared overnight from a mixture of equal volumes of protein solution (42 mg mL^{-1} in 25 mM Tris-HCl, pH 8.0, 150 mM NaCl) and reservoir solution (0.2 M NaCl, 0.1 M sodium/potassium phosphate, pH 6.2, 50% [vol/vol] polyethylene glycol [PEG] 200). Diffraction data were indexed and integrated using XDS (68) and scaled using AIMLESS (69) from the CCP4 program suite (70). The crystals displayed space group $P2_12_12_1$ with unit cell lengths $a = 39.8 \text{ \AA}$, $b = 81.4 \text{ \AA}$, and $c = 89.6 \text{ \AA}$. The asymmetric unit consisted of two polypeptide chains with an estimated solvent content of 45% and a V_m (Matthew's coefficient) of $2.24 \text{ \AA}^3/\text{Da}$. The region corresponding to the two PASTA domains in the crystal structure of *S. pneumoniae* PBP2x (PBP 5OAU) was used as a molecular replacement search model, sharing approximately 26% sequence identity with SaPASTA_{PBP1}. The search model was generated using phenix.sculptor (71) to remove nonmacromolecular chains and prune sidechains. The structure was solved by molecular replacement using PHASER (72), and the resultant electron density map was of high quality, allowing the tracing of the main chain. Model building and refinement were carried out with Coot (73) and Phenix (74), respectively. Assessments of the geometry and validation of the final model were carried out using MolProbity (75). Analyses of surface areas and interactions were made using the PISA web service (76). The graphics program PyMOL (Schrödinger, LLC) was used to generate all molecular figures presented.

Cell wall binding assays. Cell wall binding assays of recombinant PBP1 proteins fluorescently labeled with Cy2 bis-reactive dye (GE Healthcare) were performed as previously described (34), except for the binding buffer: 25 mM HEPES (pH 7.5), 150 mM NaCl, and 10 mM MgCl_2 . Cy2-labeled cytochrome c (Cy2-CytC) was included as a negative control. Binding of fluorescently labeled proteins was determined by fluorescence measurements using a Hidex Sense plate reader (excitation at 490/20 nm and emission at 535/20 nm).

Bacterial two-hybrid assay. Competent BTH101 was cotransformed with pKT25 and pUT18 derivatives. Transformants were selected on LB agar plates containing 100 $\mu\text{g mL}^{-1}$ ampicillin, 50 $\mu\text{g mL}^{-1}$ kanamycin, and 40 $\mu\text{g mL}^{-1}$ X-Gal and incubated at 30°C . Single colonies were grown in 150 μL LB with 100 $\mu\text{g mL}^{-1}$ ampicillin, 50 $\mu\text{g mL}^{-1}$ kanamycin, and 0.5 mM IPTG at 30°C .

To qualitatively measure for pairwise interactions, 5 μL of each overnight culture was spotted onto LB agar plates containing 100 $\mu\text{g mL}^{-1}$ ampicillin, 50 $\mu\text{g mL}^{-1}$ kanamycin, 0.5 mM IPTG, and 40 $\mu\text{g mL}^{-1}$ X-Gal. The plates were incubated at 30°C for 24 to 48 h in an environment protected from light and imaged. To quantify interactions, overnight cultures were assayed for β -galactosidase activity against MUG (4-methylumbelliferyl- β -D-galactopyranoside) using an assay as previously described (35).

Data availability. All study data are included in the article and/or supporting information. The data that support the findings of this study are available in the Online Research Data Figshare from the University of Sheffield with the identifier https://figshare.shef.ac.uk/collections/Penicillin-Binding_Protein_1_PBP1_of_Staphylococcus_aureus_Has_Multiple_Essential_Functions_in_Cell_Division/5656339/1. The crystal structure of the *S. aureus* PBP1 PASTA domains (PDB ID 7O61) can be accessed at <https://www.rcsb.org/structure/7O61>.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

FIG S1, TIF file, 1.6 MB.

FIG S2, TIF file, 1.3 MB.

FIG S3, TIF file, 2.5 MB.

FIG S4, TIF file, 2 MB.

FIG S5, TIF file, 1.3 MB.

FIG S6, TIF file, 2.2 MB.

FIG S7, TIF file, 1.9 MB.

FIG S8, TIF file, 1.1 MB.

TABLE S1, DOCX file, 0.05 MB.

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We declare no competing interests.

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