LytR-CpsA-Psr Enzymes as Determinants of *Bacillus anthracis* Secondary Cell Wall Polysaccharide Assembly

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*Bacillus anthracis*, the causative agent of anthrax, replicates as chains of vegetative cells by regulating the separation of septal peptidoglycan. Surface (S)-layer proteins and associated proteins (BSLs) function as chain length determinants and bind to the secondary cell wall polysaccharide (SCWP). In this study, we identified the *B. anthracis* *lcpD* mutant, which displays increased chain length and S-layer assembly defects due to diminished SCWP attachment to peptidoglycan. In contrast, the *B. anthracis* *lcpB3* variant displayed reduced cell size and chain length, which could be attributed to increased deposition of BSLs. In other bacteria, LytR-CpsA-Psr (LCP) proteins attach wall teichoic acid (WTA) and polysaccharide capsule to peptidoglycan. *B. anthracis* does not synthesize these polymers, yet its genome encodes six LCP homologues, which, when expressed in *S. aureus*, promote WTA attachment. We propose a model whereby *B. anthracis* LCPs promote attachment of SCWP precursors to discrete locations in the peptidoglycan, enabling BSL assembly and regulated separation of septal peptidoglycan.

*Bacillus anthracis*, the causative agent of anthrax, is a spore-forming, Gram-positive bacterium that germinates in host infected tissues and replicates as elongated chains of vegetative forms (1, 2). This unique growth pattern appears to be caused by the regulated separation of septal peptidoglycan, which generates bacterial chains whose mere size precludes clearance by host phagocytes (3–5). The genetic determinants for *B. anthracis* chain formation are conserved among pathogenic species of the *Bacillus cereus* group and, as demonstrated with *B. cereus* G9241, likely contribute to disease pathogenesis (6, 7). Hallmarks of pathogenic *Bacillus* species are virulence plasmids, pXO-1 and pXO-2 in *B. anthracis* (8, 9), providing for toxin production and capsulation (10, 11) as well as the chromosomally encoded surface (S)-layer locus (12). Two S-layer proteins of *B. anthracis*, Sap and EA1, are endowed with S-layer domain (SLH) domains, which retain these proteins in the bacterial envelope by binding to the secondary cell wall polysaccharide (SCWP) (13–15). S-layer protein crystallization domains, responsible for the spontaneous assembly of these polypeptides into a paracrystalline array (16), form a two-dimensional lattice that can be thought of as bacterial integument (17–19).

The structural genes of S-layer proteins, *sap* and *eag*, are flanked by genes encoding determinants for S-layer protein secretion (*secA2* and *slAP*) (12) or pyruvlation (*csA/B*) (14) as well as acetylation of the SCWP (*patA1/2* and *patB1/2*) (20). CsA/B-mediated pyruvlation of the terminal N-acetylmuraminoamine (ManNAc) of the SCWP (21), with the repeat structure \(-\rightarrow \)-\(\beta-MANAC-(1 \rightarrow 4)-\beta-GLC\text{NaC}-(1 \rightarrow 6)-\alpha-\text{GlcNAc}-(1 \rightarrow 2)\), is a prerequisite for the assembly of S-layer proteins and 22 *B. anthracis* S-layer-associated proteins (BSLs) (14). PatAB1/2-mediated acetylation of SCWP molecules affects the assembly of EA1 and of some but not all BSLs (20). Recent studies have begun to identify genes for SCWP synthesis, which, unlike pyruvyl or acetyl modifications of SCWP, appears to be essential for *B. anthracis* growth (23, 24). Nevertheless, most of the genetic determinants for SCWP synthesis, including genes whose products catalyze attachment of the polysaccharide to the peptidoglycan scaffold of *B. anthracis*, remain unknown (19).

LytR-CpsA-Psr (LCP) enzymes are found in many Gram-positive bacteria, with many species possessing more than one *lcp* gene (25). LCP proteins typically encompass a short N-terminal cytoplasmic domain, followed by 1 to 5 transmembrane helices and a C-terminal LCP domain of approximately 150 residues, which is thought to be exposed on the trans side of the membrane (25) (Fig. 1). Mutations in *Streptococcus agalactiae* *cpsA* and *Streptococcus pneumoniae* *cspa*, the first *lcp* genes to be characterized, abrogate capsule synthesis, a phenotype initially attributed to transcriptional regulation (26–28). *Bacillus subtilis* harbors three *lcp* genes, designated *tagTUV*, which are located within a 50-kb region on the chromosome that also contains genes for the biosynthesis of teichuronic acid (TU), minor teichoic acid (TA), and wall teichoic acid (WTA) (29). Expression of at least one *lcp* gene—*tagT*, *tagU*, or *tagV*—is essential for *B. subtilis* growth and teichoic acid synthesis, which prompted Kawai and colleagues to propose a model whereby LCP proteins catalyze transfer of bac-toprenol phosphate-linked capsular polysaccharide or teichoic acid precursors to peptidoglycan (29). In agreement with this model, the crystal structure of recombinant Cps2A and TagT revealed polyprenyl lipids bound to the active site as well as intrinsic pyrophosphatase activity for both proteins (29, 30). The genome of *Staphylococcus aureus* contains three *lcp* genes, designated *lcpA*, *lcpB*, and *lcpC* (31, 32). Staphylococcal mutants lacking all three *lcp* genes (Δ*lcp*), although viable, display growth defects, are unable to attach either WTA or capsular polysaccharide to pepti-
doglycan, and release small amounts of these polymers into the extracellular medium (32–34). In *Actinomyces oris*, depletion of sortase A prevents bacterial growth and this phenotype can be suppressed by transposon insertions in a gene cluster encoding a LytR-CpsA-Psr (LCP)-like protein as well as with GspA (AcaA), a glycosylated surface protein substrate of sortase (35). Of note, GspA accumulated in the membrane of the sortase A mutant but not in wild-type *Actinomyces* (35). Thus, an LCP-like protein appears to promote the glycosylation of the GspA surface proteins in *A. oris* (35).

*B. anthracis* lacks the genes for ribitol phosphate (RboP) teichoic acid synthesis and elaborates poly-D-γ-glutamic acid cap-
sule, which is tethered via amide bonds to bacterial peptidoglycan (36, 37). Screening transposon mutants of \( B. \textit{anthracis} \) for chain length phenotypes, we identified the \( lcpD \) mutant, which displays a chain length phenotype and assembly defect for the \( S \)-layer protein Sap. \( B. \textit{anthracis} \) encodes six LCP homologues (the products of \( lcpB1, lcpB2, lcpB3, lcpB4, lcpC, \) and \( lcpD \)), whose functions were validated by heterologous expression, measuring WTA synthesis in the \( S. \textit{aureus} \Delta lcp \) mutant. We also report reduced cell size and diminished chain length in the \( lcpB3 \) variant, a phenotype that may be attributable to the altered deposition of two murein hydrolases, BsU and BsR, in the envelope of \( B. \textit{anthracis} \).

**MATERIALS AND METHODS**

**Bacterial growth and reagents.** \( B. \textit{anthracis} \) strain Sterne 34F2 (38) and its variants were grown in brain heart infusion (BHI) broth or agar at 37°C. \( S. \textit{aureus} \) Newman (39) and its variants were grown in tryptic soy broth (TSB) or on tryptic soy agar (TSA) at 37°C. \( E. \textit{coli} \) was grown in Luria-Bertani broth (LB) or agar at 37°C. Where necessary, ampicillin (Amp; 100 \( \mu \)g/ml), chloramphenicol (Cm; 10 \( \mu \)g/ml), spectinomycin (Sp; 200 \( \mu \)g/ml), erythromycin (Erm; 10 \( \mu \)g/ml), or kanamycin (Kan; 20 \( \mu \)g/ml) was added to cultures. \( S. \textit{aureus} \) WTA synthesis was inhibited with tunicamycin (Tun; 1 \( \mu \)g/ml) (14). \( B. \textit{anthracis} \) strains were sporulated in modified G medium (modG) (41). Spore preparations were heat treated at 68°C for 2 h to kill vegetative bacilli and then stored at 4°C. Spore titer were determined by plating aliquots of serially diluted spore preparations on BHI agar; CFU were enumerated after overnight incubation at 37°C. Spores were germinated by incubation into BHI and grown at 37°C. To examine growth, overnight cultures were normalized to an absorbance at 600 nm of 5 and diluted 1:100 into 100 g/ml of fresh medium (BHI or TSB) with appropriate antibiotics, and growth at 37°C was monitored every 30 min for 12 h in a Synergy HT plate reader (BioTek) by recording the absorbance.

**Bacterial strains and plasmids.** Bacterial strains and plasmids used in this study are listed in Table S1 in the supplemental material. The alleles \( lcpB2:apha3 \) and \( lcpD:aad9 \) were obtained by transposon mutagenesis with \( B. \textit{aureus} \) (42). The alleles were transduced into wild-type \( B. \textit{anthracis} \) Sterne with bacteriophage CP-51 (32). The location of insertional lesions was determined by sequence analyses of inverse PCR products of the transposon/chromosome junction using the primers Martin-F and SpecR (42). Deletion of \( lcpB1, lcpB3, lcpB4, \) and \( lcpC \) was achieved by allelic replacement with the temperature-sensitive vector pLM4 (43). Briefly, 1-kbp flanking DNA sequences upstream and downstream of target genes were ligated to generate 2 kbp of recombinant DNA that was cloned into pLM4. All cloning steps were performed in \( E. \textit{coli} \) K1077 (with \( dam \) and \( dcm \) mutations), recombinant plasmids were electroplated into \( B. \textit{anthracis} \) (44), and allelic replacement was induced via changes in incubation temperature (43). Mutant alleles were verified by PCR using chromosomal DNA as a template and primers flanking the cloning sites. \( S. \textit{aureus} \) MSSA112 and its isogenic variant lacking \( lcpA, lcpB, \) and \( lcpC \) (\( \Delta lcp \)) have been described previously (31). For complementation studies, \( B. \textit{anthracis} \) \( lcp \) genes were cloned into pWWW412 and expressed via the constitutive \( hprK \) promoter of \( S. \textit{aureus} \) (45). All \( lcp \) genes were amplified from \( B. \textit{anthracis} \) Sterne chromosomal DNA by PCR using AccuPrime Pfx DNA polymerase (Invitrogen). PCR products were ligated into pWWW412, and recombinant plasmids were analyzed by DNA sequencing. Plasmid \( pLcpA \) was described earlier (32). The names and sequences of primers are listed in Table S2 in the supplemental material.

**Microscopy, chain length measurements, and immunofluorescence.** To analyze chain lengths, \( B. \textit{anthracis} \) spores were germinated by suspension in BHI broth at 37°C. At defined time points, aliquots of cells were removed and fixed with 4% formalin, and images were captured with a charge-coupled-device (CCD) camera on an Olympus IX81 microscope using 20 \( \times \) or 40 \( \times \) objectives. Chain lengths of bacilli were measured from acquired differential interference contrast (DIC) images using ImageJ and converted to lengths (in micrometers) using reference images of an objective micrometer (4). Statistical significance of chain length distributions between different strains was analyzed using one-way analysis of variance (ANOVA) (4).

For fluorescence microscopy, germinated spores of \( B. \textit{anthracis} \) strains were fixed in 4% formalin at either 3 or 8 h postgermination. After a washing to remove fixative, samples were incubated with rabbit antiserum raised against purified Sap, EAI, BsLr, or BsU, and bound antibody was detected with DyLight594-conjugated anti-rabbit secondary antibody. Cells were counterstained with boron dipyrromethene (BODIPY)-vancocin. A Leica SP5 STED-CW Superresolution laser scanning confocal microscope with a 63× objective was used to analyze the fluorescence of \( B. \textit{anthracis} \) cells (12).

**Preparation of murein sacculi and analysis of secondary cell wall polymers.** Purification of SCWP from \( B. \textit{anthracis} \) Sterne and its variants followed a previously described protocol (14, 20). Briefly, \( B. \textit{anthracis} \) strains were grown overnight at 37°C on BHI agar plates, scraped and suspended in 4% SDS, and boiled for 30 min. Bacteria were mechanically lysed with 0.1-mm glass beads and murein sacculi sedimented by centrifugation. The pellet was suspended in 100 mM Tris-HCl (pH 7.5) and incubated for 4 h at 37°C with 10 \( \mu \)g/ml of DNase and 10 \( \mu \)g/ml of RNase supplemented with 20 mM MgSO\(_4\). Samples were then incubated for 16 h at 37°C with 10 \( \mu \)M trypsin supplemented with 10 mM CaCl\(_2\). Enzymes were inactivated by boiling for 30 min in 1% SDS. Murein sacculi were washed sequentially with water, 100 mM Tris-HCl (pH 8.0), water, 0.1 M EDTA (pH 8.0), water, and acetone and then washed twice with water. Murein sacculi were suspended in 5 ml of water, and 25 ml of 48% hydrochloric acid (HCl) was added. Samples were incubated overnight on ice with shaking. Acid-extracted murein sacculi were sedimented and SCWP in the supernatant fractions was precipitated with ice-cold ethanol. The polysaccharide was sedimented by centrifugation, washed extensively with ice-cold ethanol, and suspended in 50 mM phosphate buffer for size exclusion high-performance liquid chromatography (SEC-HPLC) analysis.

\( S. \textit{aureus} \) WTA was extracted following NaOH treatment of murein sacculi prepared from staphylococci and analyzed by polyacrylamide gel electrophoresis (PAGE) as described earlier (32).

**RESULTS**

**\( B. \textit{anthracis} \) LCP genes.** A library of \( B. \textit{anthracis} \) Sterne mutants with insertional lesions was screened for chain length variants (12). This search identified a variant with \( B. \textit{aureus} \) insertion in BAS5047, which displays increased chain length of vegetative forms compared to its wild-type parent (see Table S1 in the supplemental material). The bas5047-encoded protein product is a homologue of \( B. \textit{subtilis} \) TagV (TagV\(_{Bs}\)), a member of the LytR-CpsA-Psr (LCP) family in Gram-positive bacteria (29) (Fig. 1). This family of proteins is involved in anchoring WTA and anionic polysaccharides to peptidoglycan. This was a surprising result, as \( B. \textit{anthracis} \) lacks the genes for WTA synthesis and is not known to elaborate anionic polysaccharides (36).

Amino acid sequences of \( S. \textit{aureus} \) LcpA\(_{Sa}\), LcpB\(_{Sa}\), and LcpC\(_{Sa}\) were used as queries for BLAST searches, which identified six LCP homologues in the genome of \( B. \textit{anthracis} \) Sterne: BAS5047, BAS50572, BAS50746, BAS1830, BAS33581, and BAS5115. When queried for the presence of hydrophobic and functional domains, a conserved topology was observed with the N-terminal end inside the cell and the C-terminal end outside, resulting in the surface display of the LCP domain (Fig. 1A). The Clustal W program was used to perform a multiple-sequence alignment and derive a dendrogram comparing LCPs of \( B. \textit{anthracis} \), \( B. \textit{subtilis} \), and \( S. \textit{aureus} \) (Fig. 1B and C). \( B. \textit{anthracis} \) and \( B. \textit{subtilis} \) LCPs segregated into three groups, one corresponding to \( S. \textit{aureus} \) LcpB\(_{Sa}\) (B family), one corresponding to \( S. \textit{aureus} \) LcpC\(_{Sa}\) (C family), and one with...
out a corresponding S. aureus LCP (D family). S. aureus LcpA_{Sa} (A family) did not segregate with LCPs from B. anthracis and B. subtilis (Fig. 1B). Amino acid conservation across all LCPs occurs almost exclusively within the LCP domains, and families B, C, and D are distinguished by key amino acid identities (Fig. 1C). From this analysis, we derived the following nomenclature for B. anthracis LCPs: LcpB1_{Ba} (BAS1820), LcpB2_{Ba} (BAS0572), LcpB3_{Ba} (BAS0746), LcpB4_{Ba} (BAS3381), LcpC_{Ba} (BAS5115), and LcpD_{Ba} (BAS5047). The data suggest further that B. anthracis lcpB genes likely arose via gene duplication events.

**Mutations in lcpD and lcpB3 impact Bacillus anthracis chain length.** Screening transposon mutants by DNA sequencing, we identified a B. anthracis Sterne variant with a Bursa aurealis insertion in lcpB2. Alleric replacement was used to generate B. anthracis Sterne variants with deletions of BAS1820 (lcpB1), BAS0746 (lcpB3), BAS3381 (lcpB4), and BAS5115 (lcpC). By measuring increased absorbance (A_{oo}) of cultures incubated at 37°C, it was determined that B. anthracis variants lacking any one of the six lcp genes grew at a rate similar to that of their wild-type parent (Fig. 2A). Further, following incubation in sporulation medium, quantification of the abundance of heat-resistant spores showed that B. anthracis variants lacking any one of the six lcp genes sporulated with an efficiency similar to that of the wild-type (data not shown). To assess chain formation, equal numbers of spores were diluted into fresh broth for germination of vegetative forms and synchronized growth. At timed intervals (3, 3.5, and 4 h), culture aliquots were fixed, images of 100 chains of vegetative bacilli were captured by DIC microscopy, and their lengths and widths were quantified (Fig. 2B). The chain lengths of lcpB1, lcpB2, lcpB4, or lcpC mutant bacilli were not significantly different from those of wild-type B. anthracis. In contrast, disruption of lcpD resulted in increased chain lengths at each observation period, whereas disruption of lcpB3 produced shortened chains (Fig. 2B). The increased chain length phenotype of the lcpD mutant was caused by the insertion lesion in lcpD, as transformation of the mutant with a plasmid carrying wild-type lcpD reversed the phenotype to wild-type levels (Fig. 3A). Of note, the width and volume of individual lcpD cells did not differ from those of wild-type bacilli (Fig. 3C to E).

Plasmid-borne lcpB3 (p{lcpB3}) increased the otherwise shortened chain length of the lcpB3 mutant to a level that exceeded even the chain length of wild-type bacilli (Fig. 3B). Microscopy images suggested that individual cells lacking lcpB3, unlike lcpD mutant bacilli, are smaller than the vegetative forms of wild-type bacilli (Fig. 3C). Using fluorescence microscopy and BODIPY-vancomycin staining of cell septa, we quantified the lengths of >200 bacilli, which revealed a reduced cell length for the lcpB3 mutant compared to wild-type B. anthracis (Fig. 3C and D). Further, when measuring cell widths and deriving volume, it was found that lcpB3 vegetative forms were smaller than wild-type B. anthracis (Fig. 3E). Both phenotypes, diminished cell length and volume, were restored when p{lcpB3} was transformed into the lcpB3 mutant strain (Fig. 3B). Thus, the lcpB3 mutation diminishes B. anthracis cell size and chain length, whereas the lcpD mutation increases the chain length of its vegetative forms.

**Bacillus anthracis LCP enzymes attach teichoic acid to Staphylococcus aureus peptidoglycan.** The S. aureus Δlcp mutant lacks LCP enzymes and cannot attach ribitol phosphate (RboP) WTA to peptidoglycan, a phenotype that is restored by plasmid-borne expression of any one of its three lcp genes, lcpA_{Sa}, lcpB_{Sa}, or lcpC_{Sa} (32). We asked whether the B. anthracis lcp genes can complement the WTA attachment phenotype of the S. aureus Δlcp mutant. Plasmids expressing any one of the six B. anthracis LCP enzymes or the lcpA_{Sa} control were transformed into S. aureus MSSA1112 Δlcp. The murein sacculi from each strain were isolated and treated with dilute sodium hydroxide, which releases RboP polymers via alkaline hydrolysis from peptidoglycan. Extracts were separated by PAGE and stained with alcian blue, which releases RboP polymers via alkaline hydrolysis from peptidoglycan. Extracts were separated by PAGE and stained with alcian blue and silver. As controls, alkaline hydrolysis liberated RboP from wild-type S. aureus but not from tunicamycin-treated cultures, in which TagO-mediated WTA synthesis was inhibited (Fig. 4A). As expected, alkaline hydrolysis did not release RboP from S. aureus Δlcp mutant murein sacculi unless the strain carried a complementing plasmid (p{lcp}_{Sa}) (Fig. 4A). Plasmids harboring B. anthra-
cis lcpB2 (pCPB2_{mut}), lcpC (pCPB1_{mut}), or lcpD (pCPB1_{mut}) restored S. aureus Δlcp mutant WTA assembly to the same level as pCPB1_{mut} (Fig. 4A). Plasmids harboring B. anthracis lcpB3 (pCPB3_{mut}) or lcpB4 (pCPB4_{mut}) supported attachment of reduced amounts of WTA, whereas lcpB1 (pCPB1_{mut}) failed to anchor detectable levels of RboP polymer (Fig. 4A). Because plasmid pCPB1_{mut} restored the chain length defect of the lcpD mutant (data not shown), we surmise that failure to restore WTA assembly in S. aureus is not merely due to a lack of lcpB1_{mut} expression.

Compared to wild-type S. aureus, the Δlcp mutant strain displays a growth defect, presumably because bactoprenol, a carrier required for peptidoglycan synthesis, is sequestered in the WTA synthesis pathway (32). In agreement with this model, transformation of the S. aureus Δlcp mutant with pCPB1_{mut} but not empty vector promoted growth of the mutant strain (32) (Fig. 4B). Similar to the case with pCPB1_{mut}, transformation of the Δlcp mutant with pCPB2_{mut}, pCPB3_{mut}, or pCPB4_{mut} also promoted growth, whereas pCPB1_{mut}, pCPB4_{mut}, or pCPB1_{mut} did not (Fig. 4B). Together these data indicate that lcpB2, lcpB3, lcpB4, lcpC, and lcpD encode functional LCP enzymes that can catalyze transfer of bactoprenol-linked RboP to peptidoglycan in a heterologous host.

LcpD promotes SCWP attachment to Bacillus anthracis peptidoglycan. Similar to S. aureus WTA, the SCWP of B. anthracis can be released from murein sacculi via hydrofluoric acid (HF) treatment (14, 20). HF-extracted SCWP can be purified and quantified by size exclusion high-performance liquid chromatography (SEC-HPLC), resulting in a homogenous absorbance peak at 206 nm (20) (Fig. 5A). To quantify the abundance of SCWP attached to peptidoglycan, equal amounts of murein sacculi from wild-type B. anthracis Sterne and its lcpD mutant carrying either vector alone or plasmid-encoded lcpD (pCPB1) were extracted with HF and analyzed by SEC-HPLC. As determined by quantifying the absorbance peak at 206 nm, significantly less SCWP was extracted by HF from the envelope of the lcpD mutant than from that of wild-type bacilli (Fig. 5A and B). These data suggest that plasmid-encoded LcpD is overproduced (a phenomenon described before [46]) and that the LcpD enzyme promotes the attachment of SCWP to B. anthracis peptidoglycan.

Assembly of S-layer and S-layer-associated proteins in Bacillus anthracis lcp mutants. B. anthracis chain length is impacted by genes that affect SCWP synthesis (gmeZ) (23) or modification (csaB, patA1/2, and patB1/2) (14, 20) as well as the S-layer (sap) and S-layer-associated proteins that are anchored to it (bslO) or...
that promote the secretion of their precursors (secA2 and slaP) (4, 5, 12). We asked whether lcp mutations, which appear to affect SCWP attachment to peptidoglycan, also impact the assembly of S-layer and S-layer-associated proteins. Spores from individual B. anthracis lcp mutants were inoculated into broth and incubated. At timed intervals, culture aliquots were examined for the deposition of Sap, EA1, BslR, and BslU on the surface of vegetative cells by immunofluorescence microscopy. The lcpB3 and lcpD mutants displayed S-layer assembly phenotypes, and our data analysis was limited to these strains. Compared to the case with wild-type bacilli, the abundance of Sap, the predominant S-layer protein of B. anthracis, was diminished in both lcpD and lcpB3 variants (Fig. 6A). This defect was rescued for each of these mutants by plasmid-borne expression of the corresponding wild-type lcp gene (Fig. 6A). As a control, sap mutant bacilli did not generate immunofluorescent signals during staining with Sap antiserum, demonstrating the specificity of these antibodies (Fig. 6A). The abundance and localized deposition of the EA1 S-layer protein at the septa and cell poles were not affected in the lcpD and lcpB3 mutants (Fig. 6B). BODIPY-vancomycin labeling was also unchanged, suggesting that the loss of lcp genes or their overexpression did not alter early cell division events.

BslR and BslU, two S-layer-associated proteins with predicted murein hydrolase activity (N-acetylmuramyl-l-alanine-amidase domains), attach to the SCWP and the envelope of B. anthracis in a manner requiring N-terminal S-layer homology domains, similar to Sap and EA1 (7). Unlike the two highly abundant S-layer proteins, BSLs are deposited with lesser abundance and at discrete locations in the envelope of B. anthracis (5). Immunofluorescence microscopy of wild-type bacteria revealed that BslR localizes in uneven patches throughout the envelope, whereas BslU is deposited in a few discrete puncta (Fig. 7). The lcpD mutation did not affect the distribution or abundance of BslR and BslU, as these proteins were detected in similar abundances and locations in the envelope of the lcpD mutant (Fig. 7). In contrast, in the lcpB3 mutant, BslR- and BslU-specific staining was increased in the lcpB3 mutant, where these proteins appear to accumulate at the most distal poles of chains formed from vegetative cells (Fig. 7). Plasmid-borne expression of wild-type lcpB3 (plpB3) somewhat improved the distribution of these proteins along the elongated chains of vegetative forms; however, the intensity of BslU and BslR staining remained increased compared to that for wild-type bacilli (Fig. 7). We surmise that the phenotypes of B. anthracis lcpB3 (plpb3) cells are caused by the overexpression of lpB3 from the constitutive hrpK promoter in the high-copy-number plasmid pWWW412 (see Table S1 in the supplemental material). As noted

![FIG 4 Bacillus anthracis LCP enzymes attach teichoic acid to Staphylococcus aureus peptidoglycan. (A) Purified murein sacculi from S. aureus MSSA1112 (WT with empty vector) grown in the absence (lane 1) or presence (lane 2) of tunicamycin (tun) were treated with dilute sodium hydroxide and the released wall teichoic acids (WTA) were separated by PAGE and stained with alcian blue and silver. Lanes 3 to 10, WTA extracts of murein sacculi from the S. aureus Δlcp mutant carrying either empty plasmid (vector), control plasmid with S. aureus lcpA (plcpA), or plasmids with any one of the six lcp genes of B. anthracis: plcpB1, plcpB2, plcpB3, plcpB4, plcpC, and plcpD. The bottom portion shows an immunoblot for staphylococcal sortase (SrtA) using bacterial lysate samples for the corresponding WTA extracts. (B) Growth of the S. aureus Δlcp mutant carrying vector, plcpA, or plasmids with the six lcp genes of B. anthracis was monitored by first normalizing stationary-phase cultures to an A600 of 5, next diluting these cultures (1:100) into fresh brain heart infusion broth, and then recording the A600 during the 12-h incubation at 37°C with rotation.

![FIG 5 The lcpD mutation diminishes the attachment of secondary cell wall polysaccharide (SCWP) to Bacillus anthracis peptidoglycan. (A) Equal amounts of purified murein sacculi from B. anthracis Sterne (wild-type [WT] with empty vector) or its lcpD mutant carrying empty vector or complementing plasmid (plcpD) were treated with hydrofluoric acid, released SCWP was separated by SEC-HPLC, and absorbance was recorded at 206 nm (mAu). (B) SEC-HPLC peak sizes were measured (mAu) in triplicate from independent samples, and the average peak size and standard deviation were calculated. Comparison of SCWP abundance between strains with the two-tailed Student’s t test was used to analyze statistically significant differences (*, P < 0.05).](http://jb.asm.org/.../fig5.png)
above, lcpB3 overexpression also resulted in elongated chains of vegetative bacilli. We presume that the increased abundance of BslU and BslR near the poles and cell septa of the B. anthracis lcpB3 mutant may be responsible for the reduced chain length phenotype of this variant. On the other hand, the reduced abundance of Sap is likely responsible for the increased chain length phenotype of the B. anthracis lcpD mutant (5).

**DISCUSSION**

In this study, we characterized lcp genes as determinants of B. anthracis cell size and chain length, a unique growth pattern that involves assembly of the SCWP and attached S-layer and BSL proteins (19). Earlier work had identified LCP proteins as catalysts of WTA and capsular polysaccharide attachment to peptidoglycan (29), specifically the formation of HF-sensitive phosphodiester...
bonds between the C-6 hydroxyl of N-acetylmuramic acid and WTA or capsular polysaccharide (47–49). Of note, recent work implicated an LCP-like protein from A. orsi in the glycosylation of a sortase-anchored surface protein, although the chemical nature of the glycosyl modification is still unknown (35). S. aureus Newman variants lacking all three lcp genes (Δlcp) cannot attach either WTA or the type 5 capsule, a polysaccharide with the repeat structure \( \alpha(4)\beta-M\text{AnNAc}\alpha\text{-}(1\rightarrow4)\alpha\text{-FucNAc(OAc)}\alpha\text{-}(1\rightarrow3)\beta\text{-FucNAc(1\rightarrow}\text{n}) \) (50), to peptidoglycan (32,33). Staphylococcal WTA synthesis requires tagO and tagA, whose products promote synthesis of murein linkage units (GlcNAc-ManNAc) on bactoprenol carriers (49,51,52). The tagO murein linkage pathway is dispensable for the synthesis of capsular polysaccharide, which also involves bactoprenol as the lipid carrier (33). When examined with single and double lcp mutants, S. aureus LcpASa, LcpBSa, and LcpCSa have been found to exert substrate preferences for WTA or

FIG 7 Deposition of S-layer-associated proteins in the envelope of wild-type and lcpB3 and lcpD mutant Bacillus anthracis. Spores from wild-type B. anthracis Sterne (WT with empty vector) or lcpB3 or lcpD mutant strains containing empty vector or complementing plasmid were germinated in BHI broth, incubated for 8 h, fixed in formalin, and stained with polyclonal anti-BslR (A) or anti-BslU (B) rabbit antibodies. Antibody binding and bacterial architecture were revealed with DyLight594-conjugated anti-rabbit secondary antibody and BODIPY-vancomycin, respectively. Differential interference contrast (DIC) and fluorescence microscopy images were acquired. Data sets were merged to reveal the location of the proteins in relation to the cell septa (BODIPY-vancomycin). As controls, bslR and bslU mutants were also stained with their respective antisera. Scale bar = 10 μm.
capsular polysaccharide. Nevertheless, each enzyme displays functional redundancy and catalyzes attachment of both WTA and polysaccharide to peptidoglycan (33). Testing all six *lcp* genes of *B. anthracis*, we showed in this study for the first time that heterologous expression of LCPs, specifically encoded by *lcp* genes from *B. anthracis* (which does not synthesize RboP), restores WTA synthesis in the *S. aureus* Δ*lcp* mutant. Full complementation and growth restoration were achieved with plasmids harboring *lcpB2*, *lcpC*, and *lcpD*, whereas *lcpB1* and *lcpB4* plasmids achieved partial complementation. Only the *lcpB1* plasmid failed to complement. As *lcpB1* may not be expressed in staphylococci, it seems plausible that all LCP enzymes of *B. anthracis* are capable of catalyzing WTA attachment in *S. aureus*. We showed further that *lcpD* is required for efficient attachment of SCWP to *B. anthracis* peptidoglycan. Together, these observations suggest that LCP enzymes recognize bactoprenol-(PO4)2-linked amino sugars but not RboP as the substrate for transfer to peptidoglycan. Lipid-linked LCP substrates include the GlcNAc-ManNAc moiety of murein linkage units, the ManNAc-FucNAc-FucNAc repeats of capsular polysaccharide, and the ManNAc-GlcNAc-GlcNAc repeats of SCWP.

Mutations in *B. anthracis* *lcpD* or *lcpB3* are associated with cell size and chain length phenotypes that have their underpinnings in SCWP assembly defects. In this regard, *B. anthracis* more closely resembles *B. subtilis*, in which mutations in individual *lcp* genes (*tagT*, *tagU*, and *tagV*) display discrete defects in teichoic acid attachment to peptidoglycan and in which the LCP enzymes TagT, TagU, and TagV assemble along MreB scaffolds (29, 53), cytoskeletal elements that inform assembly and disassembly of peptidoglycan and thereby determine bacterial cell shape (54, 55). The phenotypic differences between bacilli and staphylococci may be explained through differences in cell structure and cytokinesis. *S. aureus* forms spherical cells that replicate without MreB by forming cross wall structures for cytokinesis, peptidoglycan synthesis, and cell separation (56, 57). Rod-shaped bacilli, on the other hand, require MreB proteins for maintenance of cell shape and discrete mechanisms that support peptidoglycan and teichoic acid synthesis at the cylindrical and polar portions of their envelope (29). In agreement with these models, mreB, *tagT*, *tagU*, and *tagO* mutations each abrogate the rod-shaped character of *B. subtilis* (29). We presume that mutations in the corresponding genes of *B. anthracis* would cause similar phenotypes, as occurs when TagO is inhibited by tunicamycin (14).

Why has *B. anthracis* evolved six LCP enzymes when *B. subtilis* makes do with three? Unlike *B. subtilis* (58), *B. anthracis*, although unable to synthesize RboP, appears to require *tagO* for growth (14). The *tagO* gene is part of a 50-kb genomic region of *B. subtilis* that harbors virtually all genes required for WTA synthesis, including *tagTUV* (29). On the *B. anthracis* chromosome, *lcpD* (BAS5047) is also located in close proximity to *tagO* (BAS0506) and *gneY* (BAS0548) yet at a distance from *gnez* (BAS1177) (23); of note, only *gnez* is required for SCWP synthesis and is located within the *sps* gene cluster, which has been proposed to support SCWP synthesis in *B. cereus* group members (23, 24). Thus, the chromosomal region of *B. anthracis* associated with SCWP synthesis, which is currently thought to encompass the *tagO* and the *sps* gene clusters, harbors only two of the six *lcp* genes, *lcpD* and *lcpC* (BAS5115). The remaining genes—*lcpB1*, *lcpB2*, *lcpB3*, and *lcpB4*—are distributed elsewhere in the chromosome and are not associated with genes that are known to contribute to the synthesis of murein linkage units or polysaccharides. We presume that the expanded repertoire of *B. anthracis* LCP enzymes may attach different types of secondary cell wall polymers to peptidoglycan. To explore this possibility, future work could seek to identify peptidoglycan-attached polymers in variants that express only one of the six LCP enzymes.

Interestingly, *lcpB3* mutant bacilli form smaller cells as well as shortened chains and, conversely, plasmid overexpression of *lcpB3* results in chains that are longer than wild type. These phenotypes are unique to *lcpB3* and suggest a direct contribution to the bacterial cell cycle compared, for example, to the small cell phenotype observed upon overexpression of the virulence gene set (59). Both the *lcpD* and *lcpB3* mutants assembled reduced amounts of Sap, the predominant S-layer protein, in their envelope, although the spatial distribution of Sap was not affected. A clear difference between the *lcpD* and *lcpB3* mutants was observed with BslU and BslR, two predicted murein hydrolases with SLH domains (60). In wild-type bacilli, BslR is found in small patches along the cylindrical surface, whereas BslU appears to be deposited as small puncta in the immediate vicinity of septal peptidoglycan. This distribution was not affected in *lcpD* bacilli. In contrast, the deposition of BslR and BslU into the envelope of *lcpB3* mutant bacilli was increased and both proteins accumulated at the poles of the most distal cells within *lcpB3* chains. BslR and BslU are members of the family of *B. anthracis* N-acetylmuramoyl-l-alanine amidases with S-layer homology domains, which exist neither in *B. subtilis* nor in *S. aureus* (7). Plasmid pXO-2, which has been lost in the *B. anthracis* Sterne strain, encodes AmiA, one of the seven members of this family of amidases, while the remaining six genes are located on the chromosome (60). Future work must reveal the contributions of BslR and BslU toward regulated chain length of *B. anthracis* and the specific requirements for the unique distribution pattern of these BSls in the bacterial envelope.

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