

The phosphatidic acid pathway enzyme PlsX plays both catalytic and channeling roles in bacterial phospholipid synthesis

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PlsX is the first enzyme in the pathway that produces phosphatidic acid in Gram-positive bacteria. It makes acylphosphate from acyl-acyl carrier protein (acyl-ACP) and is also involved in coordinating phospholipid and fatty acid biosyntheses. PlsX is a peripheral membrane enzyme in *Bacillus subtilis*, but how it associates with the membrane remains largely unknown. In the present study, using fluorescence microscopy, liposome sedimentation, differential scanning calorimetry, and acyltransferase assays, we determined that PlsX binds directly to lipid bilayers and identified its membrane anchoring moiety, consisting of a hydrophobic loop located at the tip of two amphipathic dimerization helices. To establish the role of the membrane association of PlsX in acylphosphate synthesis and in the flux through the phosphatidic acid pathway, we then created mutations and gene fusions that prevent PlsX's interaction with the membrane. Interestingly, phospholipid synthesis was severely hampered in cells in which PlsX was detached from the membrane, and results from metabolic labeling indicated that these cells accumulated free fatty acids. Because the same mutations did not affect PlsX transacylase activity, we conclude that membrane association is required for the proper delivery of PlsX's product to PlsY, the next enzyme in the phosphatidic acid pathway. We conclude that PlsX plays a

dual role in phospholipid synthesis, acting both as a catalyst and as a chaperone protein that mediates substrate channeling into the pathway.

Most bacteria synthesize phosphatidic acid (PA),⁵ the central precursor of membrane phospholipids, using an unusual acyl-phosphate intermediate in a three-step pathway mediated by the PlsX, PlsY, and PlsC enzymes (1, 2) (Fig. 1A). PlsX is an acyl-ACP:phosphate transacylase that converts acyl-ACP made by FASII into acylphosphate (acyl-PO₄). Next, PlsY, a glycerol 3-phosphate (G3P) acyltransferase (GPAT), which uses exclusively acyl-PO₄, transfers the acyl group of acyl-PO₄ to position 1 of G3P to produce lysophosphatidic acid (LPA). The last step of the pathway is catalyzed by PlsC, an LPA acyltransferase (AGPAT), which uses acyl-ACP to convert LPA into PA (1, 3). PlsX and PlsY are found exclusively in bacteria and are essential in important human pathogens (1), therefore, these enzymes are considered promising targets for new antibacterial molecules.

Even though the enzymology of PA synthesis is well-understood, much less is known about other aspects of this pathway. One important but often unappreciated challenge of PA synthesis is the topology of the process. The amphipathic and segregated nature of the membrane bilayer makes the chemistry of its synthesis less straightforward than the metabolic reactions involving soluble molecules. Membranes grow by the *in situ* synthesis of new phospholipid molecules from soluble precursors (G3P, acyl-ACP) that are made in the cytoplasm (4). How these precursors get channeled into the membrane and how acyltransferases manage to access simultaneously substrates that are embedded within the bilayer and substrates that come

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This article contains Figs. S1–S5 and Tables S1 and S2.

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⁵ The abbreviations used are: PA, phosphatidic acid; LPA, lysophosphatidic acid; G6P, glycerol 3-phosphate; DSC, differential scanning calorimetry; LUV, large unilamellar vesicle; DMPG, 1,2-dimyristoyl-*sn*-glycero-3-phospho-(1'-rac glycerol); DMPS, 1,2-dimyristoyl-*sn*-glycero-3-phospho-L-serine; DMPC, 1,2-dimyristoyl-*sn*-glycero-3-phosphocholine; IPTG, isopropyl 1-thio- β -D-galactopyranoside; AH, amphipathic helix.

from the cytoplasm was poorly understood until the recent determination of the crystal structures of PlsY (5) and PlsC (6). PlsY is a typical integral membrane protein with seven transmembrane segments delimiting a V-shaped active site that extends deeply into the bilayer. Access of its substrate acyl-PO₄ to the active site seems to occur from within the membrane, through a putative lateral gate between transmembrane helices 2 and 5, whereas G3P enters the active site via its cytoplasmic opening (5). The crystal structure of PlsC, on the other hand, showed that the protein lacks transmembrane segments and is instead anchored to the cytoplasmic leaflet of the bilayer via a novel motif comprised of two α helices that lay parallel to the membrane (6). This mode of anchoring places the active site of PlsC above the bilayer in a way that it can simultaneously access LPA residing within the membrane and acyl-ACP from the cytoplasm (6). Despite these advances, topological information is still not available for the first reaction in the pathway, acyl-PO₄ formation by PlsX.

Based on sequence analysis and fractionation, PlsX was originally thought to be a soluble enzyme but subsequent work showed that it is a peripheral membrane protein (3, 7, 8). One of these publications reported that PlsX localization on membranes followed that of the bacterial cell division apparatus but our group has challenged this conclusion by showing that PlsX is evenly distributed on the membrane of proliferating cells, and that its localization was independent of cell division proteins (8). We also showed that PlsX association with the membrane was independent of PlsY, and that both inhibition of phospholipid synthesis and changes in membrane potential perturbed PlsX localization (8). These data suggested that PlsX associates directly with the phospholipid bilayer, but this possibility has not been tested experimentally.

Currently, two crystal structures of PlsX are available, those from *Bacillus subtilis* (9) and from the human pathogen *Enterococcus faecalis* (10). In both structures PlsX forms an S-shaped dimer. Dimerization is mainly mediated by hydrophobic contacts between a long amphipathic helix (α 9) and α 10 in each monomer, which together form a tip that protrudes away from the core of the enzyme. This dimeric structure is further stabilized by additional hydrophobic interactions with the enzyme core and several intrasubunit salt bridges on the protein surface. The core domain of the enzyme comprises a long β -strand surrounded by α -helices, generating a deep groove where the active site is located (10). Both structures suggest that this tip might participate into the association of PlsX with the membrane, however, the membrane-binding mechanism and how membrane association allows PlsX to perform its transacylase reaction are largely unknown.

In this study, we report that PlsX binds directly to lipid bilayers and identified a loop rich in hydrophobic residues localized at the tip of PlsX, which plays a critical role in mediating protein-membrane interaction. We then exploited mutations and gene fusions in PlsX to assess the role of membrane association in acyl-PO₄ production and flux through the PA pathway. Mutations that severely affect the association of PlsX to the membrane do not eliminate its transacylase enzymatic activity. Strikingly, PA synthesis is severely hampered in these cells, probably because membrane association is required for the

proper delivery of PlsX's product, acyl-PO₄, to PlsY, its partner protein. Thus, PlsX plays a dual role in PA synthesis, acting both as a catalyst and as a chaperone protein that mediates substrate channeling into the pathway.

Results

Identification of the membrane-interacting domain of PlsX

To determine how PlsX associates with membranes, we analyzed its structure (9, 10) and amino acid sequence searching for typical membrane-interacting domains. PlsX lacks terminal amphipathic helices like those mediating the association of peripheral proteins such as MinD and FtsA with the membrane (11, 12), but we found a hydrophobic loop (containing 70% hydrophobic residues, H = 0.648 and GRAVY value of 1.56, over 10 amino acids), which is solvent-exposed and next to a stretch of cationic amino acids (lysine residues) (Figs. 1B and 2A). This loop, which connects helices α 9 and α 10 and is disordered in the crystal structure, could play a role in targeting PlsX to the membrane (Fig. 1C). To test this possibility, we replaced some of the hydrophobic residues in the loop to glutamates and some of the basic amino acids in the adjacent α 10 helix to alanine residues and investigated how these mutations affected PlsX's ability to interact with phospholipids *in vitro*. We expressed and purified each mutant with a C-terminal His₆ tag and assayed their association with liposomes made with *B. subtilis* lipids. As shown in Fig. 2B, PlsX-His₆ sedimented with liposomes but when the same experiment was repeated with mutants in the hydrophobic loop (PlsX^{L254E}, PlsX^{L258E-A259E}, and PlsX^{V262E}), little or no protein was detected in the pellet, indicating the importance of these residues for lipid binding. In contrast, PlsX mutants with substitutions in positive residues (PlsX^{K264A} and PlsX^{K271A}) and in tyrosine 276 (PlsX^{Y276E}) of the α 10 helix sedimented with liposomes (Fig. 2B).

We also investigated the interaction between PlsX and membranes by differential scanning calorimetry (DSC). Fig. 2C shows the heat capacity profiles of 1,2-dimyristoyl-*sn*-glycero-3-phospho-(1'-rac glycerol) (DMPG) large unilamellar vesicles (LUVs) mixed with PlsX at different lipid-to-protein (L/P) molar ratios. Binding of PlsX to DMPG LUVs decreased the enthalpy change (ΔH) associated with the lipid phase transition (Table 1), and made this transition broader and less cooperative without changing the transition melting temperature (T_m). DSC was also employed to examine the role of head group charge in the PlsX-membrane interaction. As illustrated in Fig. S1, WT PlsX binds to LUVs made with another anionic phospholipid (phosphatidylserine, 1,2-dimyristoyl-*sn*-glycero-3-phospho-L-serine (DMPS)), but does not bind to LUVs made with a zwitterionic lipid (phosphatidylcholine, 1,2-dimyristoyl-*sn*-glycero-3-phosphocholine (DMPC)). This result highlights the importance of electrostatic interactions for PlsX-membrane association.

We next probed our mutants with DSC and observed behaviors that were in complete agreement with the sedimentation experiments: both lysine residue mutants (PlsX^{K264A} and PlsX^{K271A}) perturbed phase transition like the WT protein (Fig. 2C and Table 1), whereas the double mutant PlsX^{L258E-A259E} showed a partial effect compatible with its residual sedimenta-

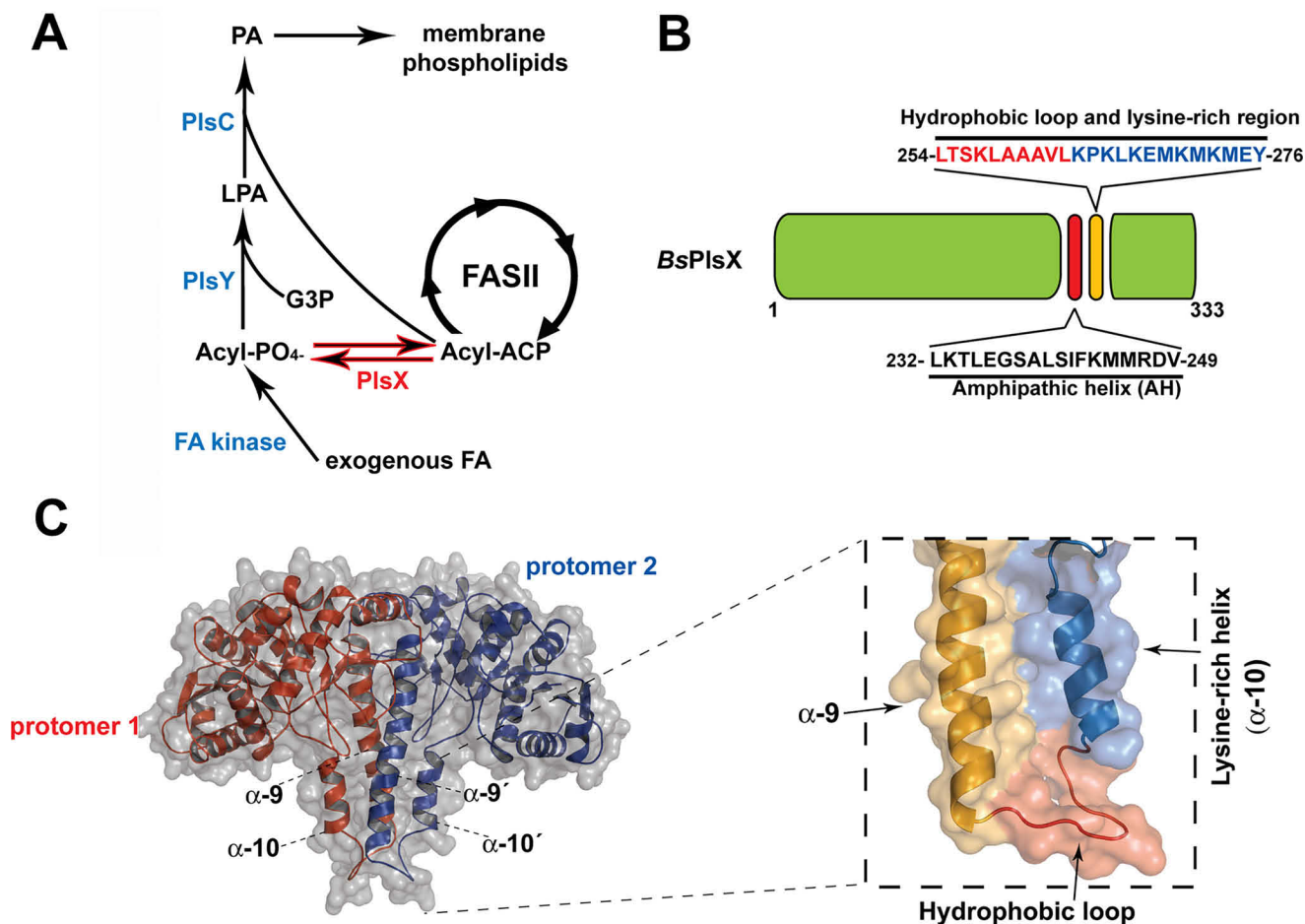


Figure 1. Phospholipid synthesis in bacteria and the structure of PlsX. A, phospholipid synthesis pathway in Gram-positive bacteria. B, diagram of the *B. subtilis* PlsX protein showing the position of the putative AH (232–249) and the hydrophobic exposed region (254–263) and lysine-rich region (264–276). C, right, structural model of *B. subtilis* PlsX dimer performed by Swiss Model homology-modeling server (31) (using PDB 1V11 as a template), indicating the localization of the hydrophobic loop and the lysine-rich helix α 10. Left, zoomed-in view of the α 9 and α 10 helices in a PlsX monomer.

tion with liposomes. Interestingly, the PlsX^{L254E} mutant had no effect on phase transition, confirming that this mutant has a severely compromised interaction with membranes (Fig. 2C).

To confirm that the hydrophobic loop identified *in vitro* was indeed necessary for membrane localization of PlsX *in vivo* we tagged the PlsX^{L254E} and PlsX^{K271A} mutant proteins with GFP and determined their subcellular localization by fluorescence microscopy. We used a strategy in which the GFP-tagged mutants were integrated at the nonessential *amyE* locus and expressed from an IPTG-responsive promoter (*P_{hyspank}*) in a strain where endogenous PlsX could be depleted by growth in the absence of xylose (DS28 background). Thus, the localization of mutant GFP-PlsX proteins was evaluated without the interference of endogenous PlsX, which could potentially hetero-oligomerize with the mutant proteins. As shown in Fig. 2D, the mutant proteins localized as expected from their ability to bind to lipids *in vitro*: GFP-PlsX^{K271A} was found in the membrane, like WT GFP-PlsX (8), whereas GFP-PlsX^{L254E} exhibited a cytosolic localization. Altogether, our data indicate that the hydrophobic loop between helices α 9 and α 10 plays a prominent role in the association and localization of PlsX to the membrane, both *in vitro* and *in vivo*.

Dimerization of PlsX is important for membrane interaction

The crystal structures of *B. subtilis* and *E. faecalis* PlsX showed that the protein is a dimer maintained mostly by interactions involving helices α 9 and α 10, which flank the hydrophobic loop responsible for membrane insertion (9, 10) (Fig. 1C). Helix α 9, in particular, is a long amphipathic helix (AH) whose hydrophobic side makes many of the contacts that stabilize the PlsX dimer. To test the role of dimerization for membrane targeting we constructed three mutants: (i) PlsX^{L235E} in which a conserved Leu in the middle of α 9 was mutated to a negatively charged Glu residue; (ii) PlsX^{AH(+2)} in which two residues (Ala-Glu) were inserted between Ile²⁴² and Phe²⁴³ to maintain the helicity but destroy the polar-apolar residue phasing of α 9; and (iii) PlsX^{AH(+3)} in which the insertion of three residues (Glu-Glu-Leu) between Ile²⁴² and Phe²⁴³ should maintain both the helicity and the amphipathic phasing of α 9 (Fig. 3A). The three mutants were expressed and purified and their oligomerization state was analyzed by gel filtration alongside the WT protein (Fig. 3B). As expected from the crystal structures and previously shown for *Streptococcus pneumoniae* PlsX (1), WT PlsX and PlsX^{L254E} eluted from the S-200 column with the expected size of a dimer (molecular mass of ~80 kDa, Fig.

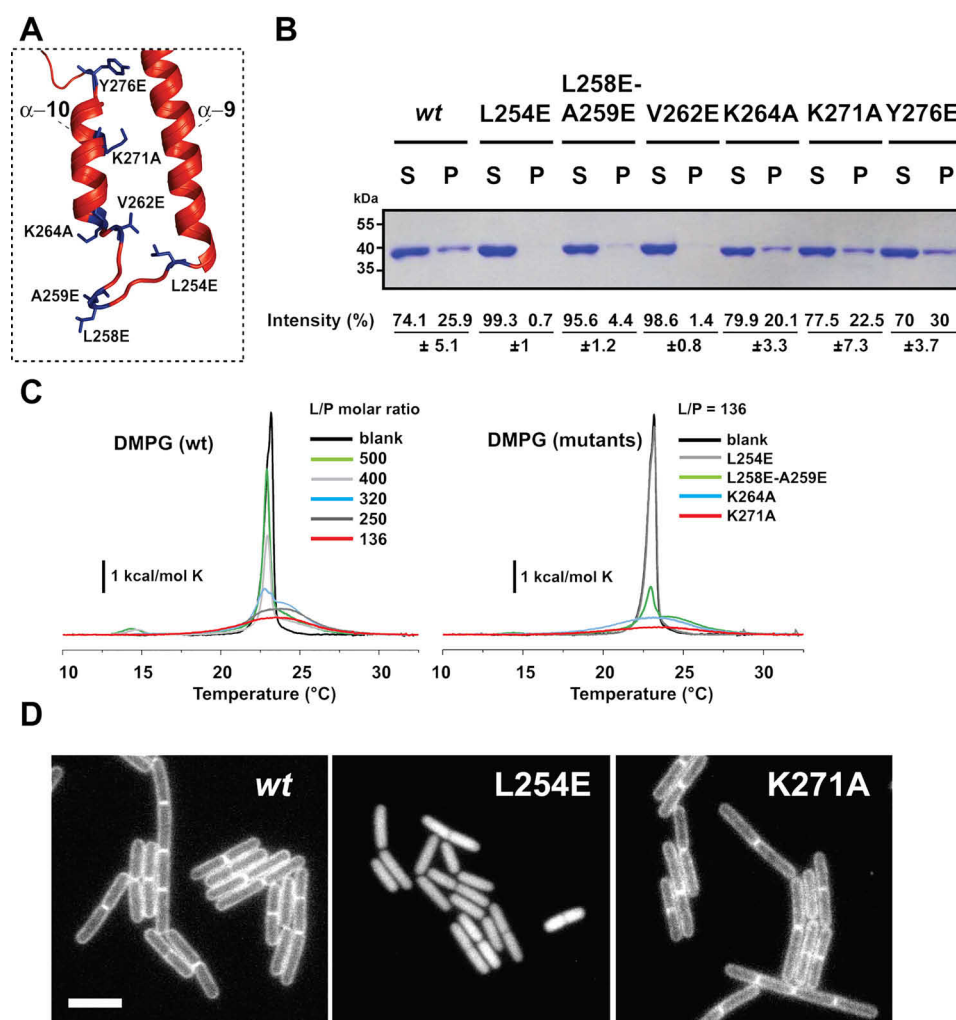


Figure 2. Mutations in the hydrophobic loop perturb PlsX membrane interaction. *A*, mutated amino acid residues of the hydrophobic loop and helix $\alpha 10$ of PlsX highlighted on the structure. *B*, cosedimentation assay showing the interaction of WT PlsX and different mutants with liposomes prepared with *B. subtilis* total lipid extracts. Data represent the mean $\pm$ S.D. from two independent experiments. *C*, effects of PlsX and selected mutants on the thermotropic behavior of lipid bilayer membranes probed by DSC. *Left*, representative DSC traces of DMPG LUVs in the absence and presence of different concentrations of PlsX. *Right*, excess heat capacity of DMPG in the absence (black) and presence of PlsX^{L254E}, PlsX^{L258E/A259E}, PlsX^{K264A}, and PlsX^{K271A} mutants at lipid-to-protein (L/P) molar ratio of 136:1. Lipid concentration was 800 μ M in all cases. *D*, fluorescence microscopy images showing the subcellular localization of GFP-PlsX wt (strain DS69; Pxyl-plsX amyE::Physpank-gfp-plsX) and PlsX^{L254E} (strain DS160; Pxyl-plsX amyE::Physpank-gfp-plsX^{L254E}) and PlsX^{K271A} (DS161; Pxyl-plsX amyE::Physpank-gfp-plsX^{K271A}); site-directed mutant. Images are representative of at least 100 cells analyzed for each condition in at least two different days. All images are at the same magnification. Scale bar = 5 μ m.

Table 1

Thermodynamic parameters associated to the DMPG phase transition

The phase transition temperature (T_m), the calorimetric enthalpy change (ΔH), and the linewidth at half height ($\Delta T_{1/2}$) were obtained from analysis of the thermograms with MicroCal Origin software. Lipid-to-protein molar ratios (L/P) are indicated. Data shown are representative of two independent experiments.

Sample	L/P	T_m °C	ΔH kcal/mol	$\Delta T_{1/2}$ °C
DMPG				
Blank		23.3 (2) ^a	5.0 (2)	0.7 (2)
+ PlsX wt	500	22.9 (1)	5.0 (1)	0.5 (1)
+ PlsX wt	400	22.8 (1)	4.5 (1)	0.6 (1)
+ PlsX wt	320	23.7 (2)	4.6 (2)	2.9 (1)
+ PlsX wt	250	23.6 (3)	4.2 (2)	4.4 (2)
+ PlsX wt	136	23.6 (3)	2.9 (3)	5.1 (2)
+ PlsX L254E	136	22.8 (1)	4.5 (1)	0.6 (1)
+ PlsX L258E-A259E	136	23.7 (2)	4.6 (2)	2.9 (1)
+ PlsX K264A	136	23.6 (3)	4.2 (2)	4.4 (2)
+ PlsX K271A	136	23.6 (3)	2.9 (3)	5.1 (2)

^a The numbers in parentheses correspond to the uncertainties in the last significant digit.

3B, Fig. S2). Significant changes in the elution pattern were observed for mutants PlsX^{AH(+2)} and PlsX^{L235E}, which displayed retention times corresponding to the monomer (~40 kDa), in addition to large aggregates (Fig. 3B), which probably arise because the mutant proteins can no longer dimerize. In contrast, mutant PlsX^{AH(+3)}, which bears a larger insertion than PlsX^{AH(+2)} but whose amphipathicity is maintained, behaved as a mixture of both monomeric and dimeric species (Fig. 3B).

To examine whether dimerization of PlsX is relevant for membrane association, the above mutants were fused to GFP and their cellular distribution was examined, employing the same strategy used to study the localization of the hydrophobic loop mutants (expression of mutant GFP-PlsX fusions from an IPTG-inducible promoter in the absence of endogenous PlsX). As shown in Fig. 3C, the two mutants that no longer dimerized, PlsX^{L235E} and PlsX^{AH(+2)}, had severely compromised membrane localization, as inferred by their cytoplasmic staining. In

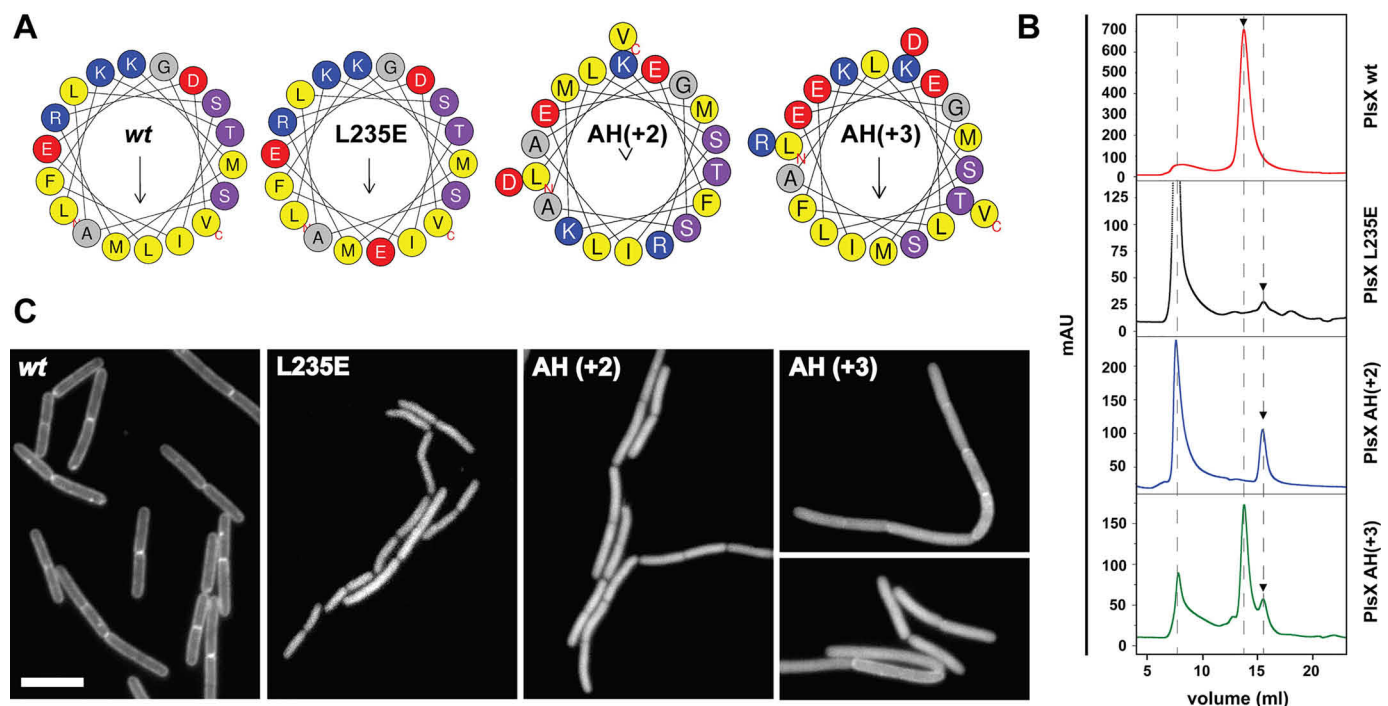


Figure 3. The amphipathic helix $\alpha 9$ is essential for the dimerization and proper localization of PlsX. A, helical wheel representation of the H9 amphipathic helix located at amino acids 232–249 of PlsX. Hydrophobic residues are marked in yellow. Mutation in one of the leucines (L235E), and insertion of two (AH(+2)) and three residues (AH(+3)) mutants are represented. Diagrams generated using the web server HELIQUEST (32). B, gel filtration chromatography assays. Purified PlsX wt (red) was a dimeric protein, whereas PlsX^{L235E} (black) and PlsX^{AH(+2)} (blue) presented monomeric and aggregated states. PlsX^{AH(+3)} (green) present both dimeric and monomeric states. Data shown are representative of two independent experiments. C, fluorescence microscopy showing subcellular localization of GFP-tagged PlsX wt (strain DS69; *P_{xyI}-plsX amyE::Physpantk-gfp-plsX*), PlsX^{AH(+2)} (strain DS34; *P_{xyI}-plsX amyE::Physpantk-gfp-plsX^{AH(+2)}*), PlsX^{AH(+3)} (strain DS48; *P_{xyI}-plsX amyE::Physpantk-gfp-plsX^{AH(+3)}*), and PlsX^{L235E} (strain DS58; *P_{xyI}-plsX amyE::Physpantk-gfp-plsX^{L235E}*) mutants. Images are representative of at least 100 cells analyzed for each condition in at least two different days. All images are at the same magnification. Scale bar = 5 μ m.

contrast, mutant PlsX^{AH(+3)}, which is partially a dimer *in vitro* (Fig. 3B), exhibited a mixed localization pattern, with both membrane and cytoplasmic staining (Fig. 3C). The importance of dimerization for membrane association was corroborated by *in vitro* experiments in which we observed that monomeric PlsX^{L235E} no longer co-sedimented with liposomes, whereas the partially dimeric PlsX^{AH(+3)} still sedimented, albeit to a lesser extent than the WT protein (Fig. S3).

Membrane association is necessary for PlsX function *in vivo*

The availability of mutants with severe defects in membrane association provided an opportunity to investigate if PlsX's membrane localization was necessary for its functioning *in vivo*. The function of PlsX mutants was monitored by their ability to support growth in the absence of WT PlsX. We created merodiploid strains with mutant PlsX alleles controlled by IPTG in our PlsX depletion strain background and assayed growth by spotting dilutions on plates containing IPTG but no xylose. As shown in Fig. 4A, there was a clear correlation between membrane association and the ability to support growth: mutants that were affected in membrane association because of alterations in the hydrophobic loop (PlsX^{L254E}) or because of impaired dimerization (PlsX^{L235E}, PlsX^{AH(+2)}) supported little or no growth in the absence of endogenous PlsX, mutant PlsX^{AH(+3)}, which exhibited less affected membrane localization, promoted intermediate levels of growth, and mutant PlsX^{K271A}, which associates with the membrane, was able to fully support growth (Fig. 4A).

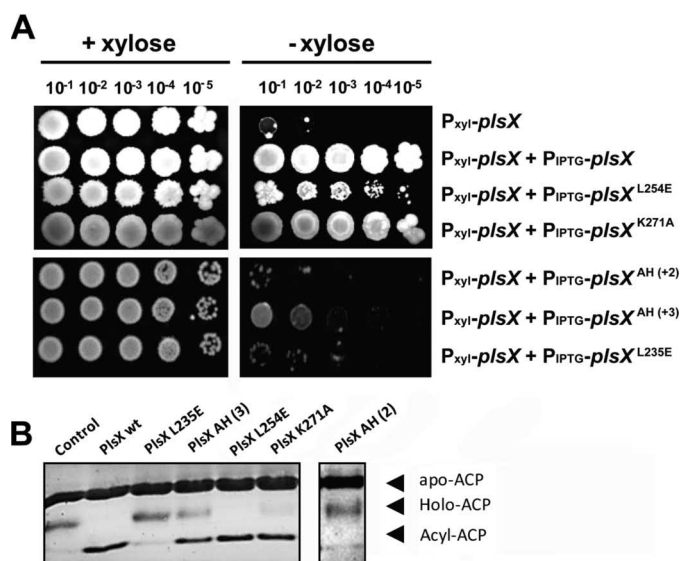


Figure 4. Proper membrane association is required for PlsX function *in vivo* but not for catalysis *in vitro*. A, growth complementation assays. Cells of *B. subtilis* DS28 (*P_{xyI}-plsX*), DS69 (*P_{xyI}-plsX amyE::Physpantk-gfp-plsX*), DS160 (*P_{xyI}-plsX amyE::Physpantk-gfp-plsXL254E*), DS34 (*P_{xyI}-plsX amyE::Physpantk-gfp-plsX^{AH(+2)}*), DS48 (*P_{xyI}-plsX amyE::Physpantk-gfp-plsX^{AH(+3)}*), and DS58 (*P_{xyI}-plsX amyE::Physpantk-gfp-plsX^{L235E}*) were grown in liquid LB to logarithmic phase, serially diluted and spotted (4 μ l) onto LB + 0.5 mM IPTG plates, with or without 0.4% xylose, and incubated at 37 °C for 24 h. B, *in vitro* acyltransferase activity assay of PlsX wt and mutants. The assay detects the synthesis of acyl-ACP from 16:0-PO₄ and holo-ACP by the appearance of a slow migrating band in conformationally sensitive gel electrophoresis (15% PAGE, 2 M urea). Data shown are representative of three independent experiments. Note that the PlsX AH(2) sample was run in a different gel and thus is depicted separate from the other samples.

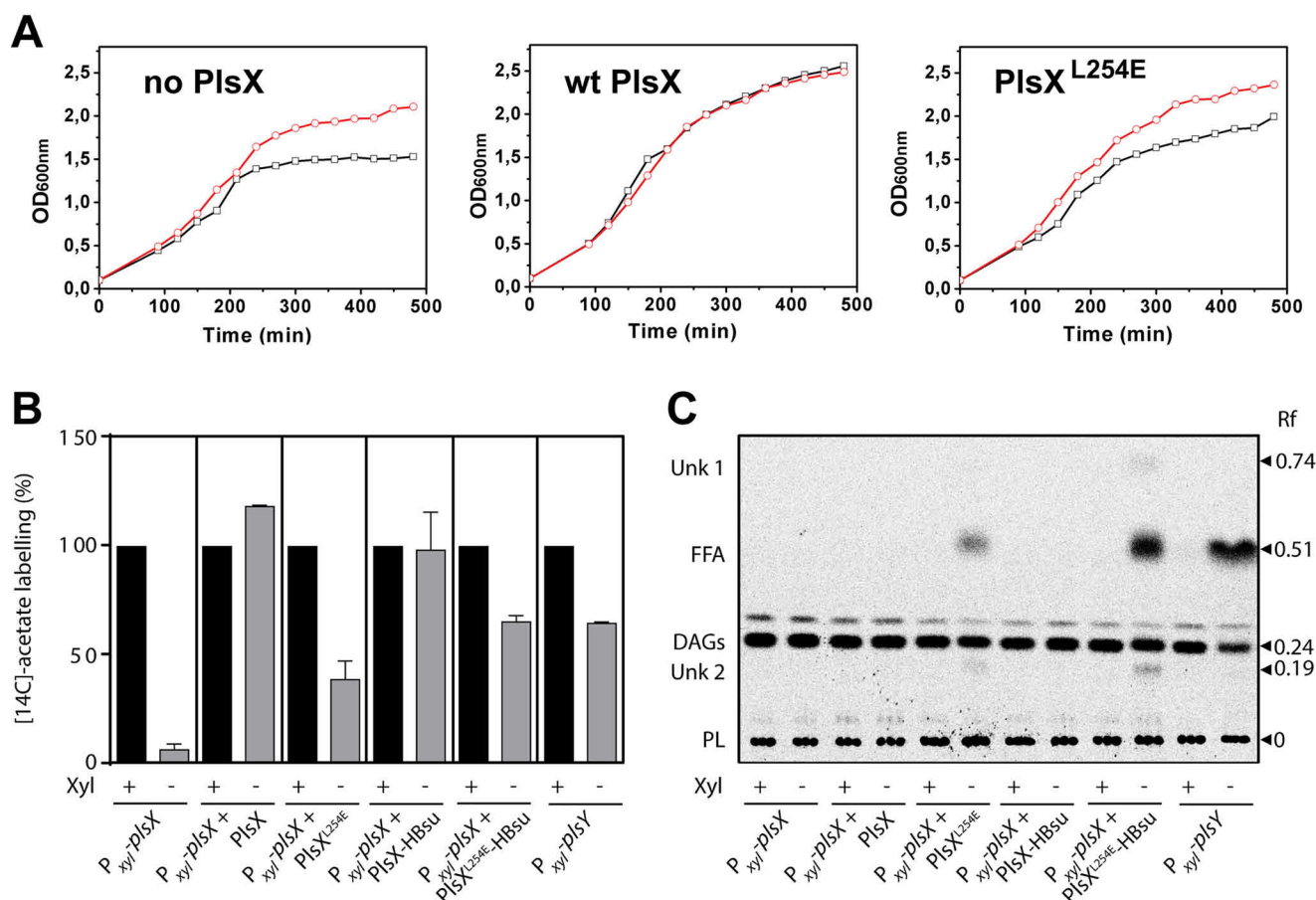


Figure 5. Removal of PlsX from the membrane blocks phospholipid synthesis. *A*, rescue of cells expressing PlsX^{L254E} by supplementation with *B. subtilis* fatty acids. Strains DS28 (P_{xyI}-plsX), DS69 (P_{xyI}-plsX amyE::Physpank-gfp-plsX^{L254E}), and DS160 (P_{xyI}-plsX amyE::Physpank-gfp-plsX^{L254E}), were grown in LB medium containing 0.5 mM IPTG and 1 mg/ml of BSA, and in the presence or absence of 0.4% xylose and/or fatty acid methyl esters (FAMES) (2 mg/ml). – FAMES, black lines/squares; + FAMES, red lines/circles. Data are representative of at least two independent experiments. *B*, lipid synthesis in cells expressing PlsX mutants. Strains were grown in LB medium in the presence of 1 mM IPTG to induce the expression of the different PlsX variants. The cultures were supplemented with xylose (0.4%) to induce the expression of WT PlsX (PlsXwt) or PlsY (in strain VD211 P_{xyI}-plsY). Strains DS28, DS160, DS171, and VD211 were labeled with 2 μ Ci/ml of [¹⁴C]acetate for 30 min during the transition from log to stationary phase. Strains DS69 and DS170 grown to midexponential phase were labeled with 2 μ Ci/ml of [¹⁴C]acetate for 30 min. Cells were harvested and total lipids were quantified in a scintillation counter. Bars represent the mean $\pm$ S.D. from three independent experiments. *C*, fatty acids accumulate in strains expressing PlsX that does not associate with the membrane. Total lipids of cells labeled with [¹⁴C]acetate as described in panel *B* were analyzed by TLC on preadsorbent Silica Gel G layers (Merck) developed with hexane-ethyl ether-acetic acid (60/40/1, v/v/v). Each lane was loaded with about 1,000 cpm of labeled lipids. The radioactive lipid species were identified by their co-migration with standards. FFA, free fatty acid; DAGs, diacylglycerols; PL, phospholipid; Unk, unknown. The example shown is typical of three independent experiments.

An important control to determine the role of PlsX localization *in vivo* was to rule out that the mutations affecting membrane association may have also affected catalysis by the enzyme. To study catalysis, we developed an *in vitro* acyltransferase assay measuring the reverse of the reaction normally executed by PlsX. We employed palmitoyl-PO₄ and holo-ACP as substrates and monitored the appearance of palmitoyl-ACP by conformation-sensitive gel electrophoresis. As shown in Fig. 4B, mutations that impaired dimerization (PlsX^{L235E}, PlsX^{AH(+2)}) abolished the production of palmitoyl-ACP by PlsX *in vitro*. This suggests that catalysis requires dimerization, a conclusion supported also by the fact that a mutation that rescues dimerization (PlsX^{AH(+3)}) is catalytically active, even though it introduces a larger insertion in helix α 9 than one that is inactive (PlsX^{AH(+2)}). Importantly, however, a mutation that compromises membrane association without affecting dimerization (PlsX^{L254E}) did not affect catalysis *in vitro* (Fig. 4B). Therefore, the PlsX^{L254E} mutation demonstrates that membrane binding and catalysis can be uncoupled in PlsX and

we further characterized this mutation to understand why membrane localization is necessary for PlsX function *in vivo*.

Removal of PlsX from the membrane blocks phospholipid synthesis and leads to free fatty acid accumulation

The growth impairment of cells expressing PlsX^{L254E} was an indication that the phospholipid synthesis pathway becomes blocked when PlsX cannot appropriately associate with the membrane. To test this possibility, we investigated if PlsX^{L254E} growth could be rescued by exogenous fatty acids, as was shown for a *plsX* null mutant (13). In Gram-positive bacteria such as *Bacillus* and *Staphylococcus*, exogenous fatty acids can bypass the need for PlsX because they are phosphorylated directly by fatty acid kinase (Fig. 1A). Indeed, we found that supplementation of a strain expressing PlsX^{L254E} with a mixture of branched-chain fatty acids rescued its growth as well as it rescued growth of a *plsX* null mutant (Fig. 5A), a strong indication that the growth defect of the PlsX^{L254E} mutant was due to inhibition of phospholipid synthesis.

We envisaged two possible scenarios to explain why phospholipid synthesis was compromised in cells expressing PlsX^{L254E}: (i) the mutant protein is not catalytically active *in vivo*, or (ii) it is catalytically active but membrane association of PlsX is necessary for efficient delivery of acyl-PO₄ to PlsY to proceed with phospholipid synthesis. We reasoned that the two possibilities could be distinguished by comparing the *de novo* lipid synthesis profile of a PlsX^{L254E} mutant with those of *plsX*[−] and *plsY*[−] mutants. If PlsX^{L254E} is inactive *in vivo* it should exhibit a *plsX*[−] phenotype, but if the block in the pathway is at the level of product delivery, PlsX^{L254E} should exhibit a *plsY*[−] phenotype. As reported previously by our group, a *plsX*[−] strain has a strong block in *de novo* lipid synthesis, presumably because the accumulation of acyl-ACP shuts down FASII (3). In contrast, *de novo* lipid synthesis is only partially inhibited in a *plsY*[−] strain, but this mutant accumulates free fatty acids, likely derived from the acyl-PO₄ produced by PlsX and which cannot be used further in the absence of PlsY (3). Therefore, we used [¹⁴C]acetate labeling to analyze lipid synthesis in a strain expressing GFP-PlsX^{L254E}, and compared it with a strain expressing a WT GFP-PlsX fusion, a conditional *plsX* mutant (*plsX*[−]) and a conditional *plsY* mutant (*plsY*[−]). As shown in Fig. 5B, [¹⁴C]acetate incorporation in a strain expressing GFP-PlsX^{L254E} was partially inhibited, reaching ~30% of the WT levels, a value much higher than the incorporation observed in the conditional *plsX* mutant (7.0%) but similar to that observed in a conditional *plsY* mutant (Fig. 5B) (3). The inhibition of the GFP-PlsX^{L254E} mutant was not due to a negative effect of the GFP fusion because lipid synthesis was normal in a strain expressing a WT GFP-PlsX fusion (Fig. 5B). In addition, TLC analysis of the newly synthesized lipids showed that the GFP-PlsX^{L254E} mutant accumulated free fatty acids to a level comparable with the conditional *plsY* mutant (Fig. 5C). Thus, collectively, these data are most consistent with the hypothesis that GFP-PlsX^{L254E} is catalytically active *in vivo* and that the main defect of this mutant protein is in channeling its product into the next step of the pathway.

Mistargeting as a way to test the role of membrane localization in PlsX function

Another way to test if membrane localization is necessary for PlsX function is to artificially target the protein to a different subcellular compartment without mutating the protein's amino acid sequence. We attempted to do that by fusing PlsX to HBsu, a histone-like protein that binds to the bacterial nucleoid with great affinity, such that normally all HBsu in the cell is nucleoid-associated (14). The high affinity of HBsu for the nucleoid and the fact that this structure is spatially segregated from the membrane suggested that fusing PlsX to HBsu might suffice to “peel” it from the membrane (Fig. 6A). However, when we imaged a strain expressing a triple GFP-PlsX-HBsu fusion we observed that the protein localized as foci or patches that overlapped with the nucleoid but that also reached the membrane, indicating that the fused protein manages to bind simultaneously to both structures (Fig. 6, B and C). In retrospect, this observation is not unexpected because it is known that DNA loops are frequently found in close proximity to the bacterial membrane (15, 16).

Although the strategy of fusing PlsX to HBsu did not work to remove the WT protein from the membrane, we reasoned it could still be employed to further evaluate the importance of membrane association for PlsX function. Even though the PlsX^{L254E} mutant displays a strongly reduced interaction with the membrane, it still supports residual growth when expressed in place of endogenous PlsX (Fig. 4A, compare spots of PlsX^{L254E} and the parental depletion strain). Furthermore, a detailed biophysical analysis of the PlsX^{L254E} mutant showed that it still retains the ability to bind superficially to membranes, probably via electrostatic interactions (33). Therefore, it is certainly possible that the cytoplasmic molecules of PlsX^{L254E} will exhibit transient or superficial contact with the membrane sufficient to allow some transfer of acyl-PO₄ to the bilayer and maintain a low level of phospholipid synthesis. Alternatively, the residual growth may result from acyl-PO₄ produced in the cytoplasm diffusing and spontaneously inserting itself in the membrane. We reasoned we could distinguish between these possibilities by fusing PlsX^{L254E} to HBsu to completely restrict access of the protein to the membrane. In fact, as shown in Fig. 6, B and C, fusing PlsX^{L254E} to HBsu led to a dramatic change in its localization, which went from cytoplasmic to being exclusively associated with the nucleoid. We then asked if this protein, which now had no access to the membrane, was capable of supporting growth in the absence of endogenous PlsX. As shown in Fig. 6D, GFP-PlsX^{L254E}-HBsu was slightly worse than GFP-PlsX^{L254E} at complementing the lack of PlsX. The similar phenotypes of GFP-PlsX^{L254E} and GFP-PlsX^{L254E}-HBsu were also evident in the [¹⁴C]acetate labeling experiments: both mutants exhibited comparable inhibition of lipid synthesis (Fig. 5B). However, the strain expressing GFP-PlsX^{L254E}-HBsu accumulated 2-fold more free fatty acids (Fig. 5C), a sign that phospholipid synthesis is indeed more impaired when PlsX is completely prevented from accessing the membrane. The worse performance of GFP-PlsX^{L254E}-HBsu cannot be explained as a general negative effect of HBsu, because fusion of WT PlsX to HBsu (GFP-PlsX-HBsu) had little if any impact on growth (Fig. 6D) or lipid synthesis (Fig. 5, B and C). We also carried out a control, which showed that all protein fusions were successfully expressed (Fig. S4). Therefore, these results reinforce our conclusion that PlsX membrane association is crucial for the efficient channeling of acyl-PO₄ into the phospholipid synthesis pathway.

Discussion

Peripheral proteins associate with the membrane in a wide variety of ways but two of the most common motifs are amphipathic helices and protruding hydrophobic loops (17). Here we showed that *B. subtilis* PlsX associates with the membrane via a hydrophobic loop comprising residues 254–263 that projects from the end of its amphipathic dimerization helices. Typical loops have exposed hydrophobic residues at their tips, flanked by cationic residues that interact favorably with the negative head groups of lipids (18, 19). In agreement with our prediction that the loop is the motif that targets PlsX to the membrane, mutations in hydrophobic residues (L254E, L258E/A259E, and V262E) perturb to varying degrees membrane interaction. In contrast, we failed to disrupt membrane binding

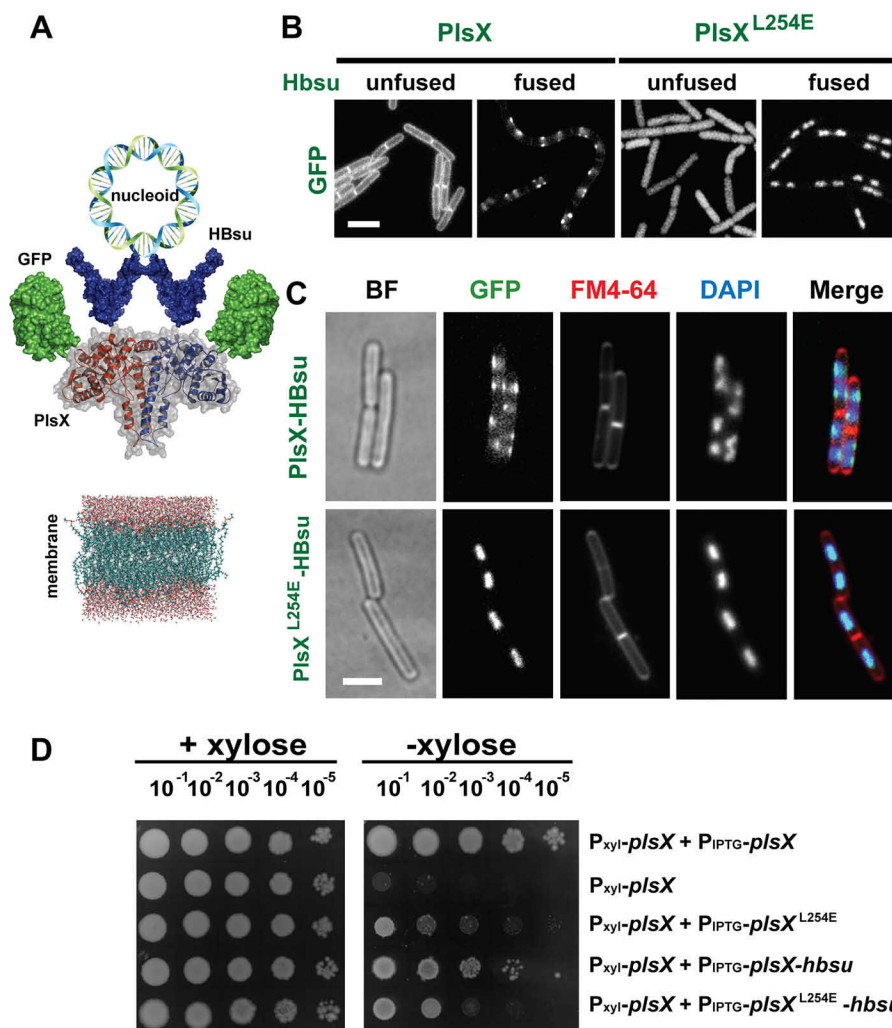


Figure 6. Subcellular localization of PlsX and PlsX^{L254E} fused to GFP and Hbsu. A, schematic representation of the triple fusion GFP-PlsX-Hbsu, designed as an attempt to remove PlsX from the cytoplasmic membrane, due to the interaction of Hbsu histone-like protein with the nucleoid. B, fluorescence microscopy showing localization of GFP-PlsX wt and GFP-PlsX^{L254E} fused and unfused to Hbsu in *B. subtilis* strains DS69 (P_{xyI}-plsX amyE::Physpank-gfp-plsX), DS160 (P_{xyI}-plsX amyE::Physpank-gfp-plsX^{L254E}), DS170 (P_{xyI}-plsX amyE::Physpank-gfp-plsX-hbsu), and DS171 (P_{xyI}-plsX amyE::Physpank-gfp-plsX^{L254E}-hbsu). C, fluorescence microscopy showing subcellular localization of GFP-PlsX-Hbsu fusions in *B. subtilis* DS170 (P_{xyI}-plsX amyE::Physpank-gfp-plsX-hbsu) and DS171 (P_{xyI}-plsX amyE::Physpank-gfp-plsX^{L254E}-hbsu) using FM4-64 and DAPI to stain membrane and nucleoid, respectively. Images are representative of at least 100 cells analyzed for each condition in at least two different days. All images are at the same magnification. Scale bar = 5 μ m. D, growth complementation assays of strains expressing PlsX-Hbsu fusions. Cells of DS28 (P_{xyI}-plsX), DS69 (P_{xyI}-plsX amyE::Physpank-gfp-plsX), DS160 (P_{xyI}-plsX amyE::Physpank-gfp-plsX^{L254E}), DS170 (P_{xyI}-plsX amyE::Physpank-gfp-plsX-hbsu), and DS171 (P_{xyI}-plsX amyE::Physpank-gfp-plsX^{L254E}-hbsu) were grown in liquid LB to logarithmic phase, serially diluted and spotted (4 μ l) onto LB + 0.5 mM IPTG plates, with or without 0.4% xylose, and incubated at 37 °C for 24 h. Data shown are representative of at least three independent experiments.

when single lysine residues (*i.e.* Lys-264 and Lys-271) of the adjacent α 10 helix were mutated. This suggests that the energetic contribution of each individual positive residue for membrane binding is modest, but we cannot rule out the possibility that the α 10 helix is not directly involved in membrane binding.

The existence of a hydrophobic loop flanked by positive residues in helix α 10 is a feature conserved in all PlsX homologues (Fig. S5). In proteobacteria, this region is slightly longer (15 residues, instead of 10) and contains more aromatic and positive residues, but still has the features expected of a membrane-binding hydrophobic loop. Hence, we expect that all PlsX proteins use similar motifs to associate with the membrane.

While this work was in preparation, Jiang and colleagues (20) described a new PlsX crystal structure in which the hydrophobic loop of one protomer is folded into a short amphipathic α helix, suggesting that the region we have so far described as a

hydrophobic loop may in fact change structure to become helical upon membrane interaction. This plausible model was supported by mutations that, similarly to ours, perturbed membrane interaction, but more detailed structural analysis in the presence of membranes will be necessary to ascertain the exact structure of PlsX. Irrespective of this structural detail, both works reinforce each other's conclusion that PlsX employs a stretch of hydrophobic and positively charged amino acids at the tip of its stalk as its membrane binding motif.

Another important observation of our work was that removal of PlsX from the membrane via mutation or the fusion to Hbsu resulted in inhibition of phospholipid synthesis. This observation may be explained if membrane association was necessary for PlsX catalysis, but we find this very unlikely for two reasons: (i) PlsX catalysis can be detected *in vitro* in the absence of lipids or detergents (see Refs. 1 and 20, and this

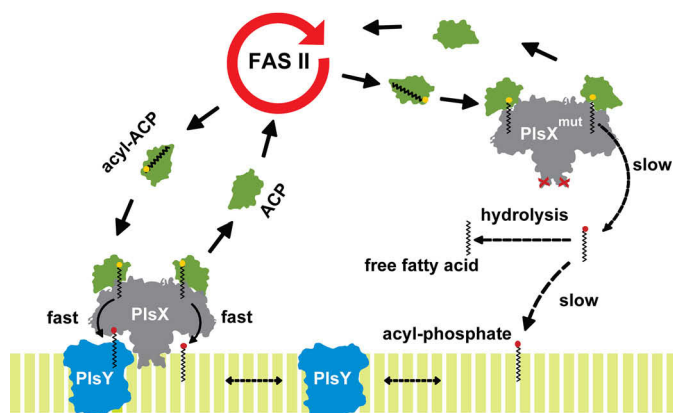


Figure 7. PlsX membrane association is crucial for acyl-PO₄ channeling to PlsY. Membrane association creates a path shielded from solvent for the transfer of acyl-PO₄ to the bilayer. Channeling may also involve transfer directly to PlsY, via protein-protein interaction. PlsX mutants that do not associate with the membrane still produce acyl-PO₄ but this cannot reach PlsY and is hydrolyzed to free fatty acids in the cytosol.

work), and (ii) we detect free fatty acid accumulation in cells expressing PlsX variants that do not localize to the membrane, confirming that the enzyme is catalytically active even when removed from the membrane. Why, then, would PlsX need to associate with the membrane to be fully functional? Being at the membrane interface makes sense for enzymes that act on substrates that are embedded in the bilayer, such as PlsC, which uses LPA, but this is not the case for PlsX whose substrates, acyl-ACP and P_i, are accessible in the cytoplasm. Instead, we propose that PlsX association with the membrane is needed to channel acyl-PO₄ efficiently into the next step of the pathway, which is carried out by PlsY, an integral membrane protein that likely access acyl-PO₄ only after it has partitioned into the membrane (5) (Fig. 7).

Acylphosphates are less water soluble than acyl-ACP (2) and this may represent a barrier to its diffusion from the cytosolic compartment and into the membrane. Although amphipathic molecules like acyl-PO₄, or even fatty acids, will spontaneously partition into the membrane from a water phase, the kinetics of the process may not be fast enough to support highly active pathways such as phospholipid synthesis (Fig. 7). In fact, the residual growth of strains containing PlsX variants that do not bind to the membrane (Figs. 4A and 6B) probably reflects the slow membrane partitioning of acyl-PO₄ that is being produced by PlsX molecules located in the cytoplasm. There are several examples of proteins that participate in cytoplasmic transport of fatty acids, whose diffusion is limited by their poor solubility (21). ACP itself can be thought of as an analogous acyl transporter, in the sense of avoiding solubility problems, because the acyl chain in an acyl-ACP molecule is tucked into a hydrophobic groove in the protein (22–24). Thus, it seems that acyl intermediates are not usually found freely surrounded by water inside cells, and PlsX would be the protein in charge of keeping acyl-PO₄ away from water as well. Finally, acylphosphates are very labile and the channeling (or chaperoning) role of PlsX may also be relevant to protect these molecules from hydrolysis, as pointed out by Li *et al.* (5). Consistent with this idea, we observe the accumulation of free fatty acids in strains in which

PlsX is cytoplasmic and no longer capable of efficiently channeling acyl-PO₄ into the membrane (Fig. 5B).

The detailed mechanism of acyl-PO₄ channeling by PlsX remains to be defined, but our data allows us to speculate that it depends on the proper and stable insertion of PlsX in the membrane. This can be concluded from the biophysical analysis of the PlsX^{L254E} mutant presented in the accompanying paper (33), which showed that this protein can interact with membranes only superficially. A similar conclusion is also supported by *in vivo* experiments, which showed that the phenotype of the PlsX^{L254E} mutant is not much worse than the phenotype of a strain in which PlsX can never access the membrane (PlsX^{L254E}-HBSu) (Fig. 6C). Together, these results mean that superficially accessing the membrane is not sufficient to ensure efficient transfer of acyl-PO₄. Stable insertion in the membrane may be necessary to create a path shielded from solvent for acyl-PO₄ to leave the active site and insert into the bilayer. This path may also involve the physical association between PlsX and PlsY, as these proteins were shown to interact in two-hybrid experiments (25). Proper interaction between PlsX and PlsY may only happen if PlsX is stably inserted in the membrane.

In conclusion, we have identified a loop rich in hydrophobic residues localized at the tip of the stem of PlsX that plays a critical role in mediating protein-membrane interaction and used this information to advance our understanding of bacterial phospholipid biosynthesis. On the basis of our combined results, we propose that the proper membrane interaction and localization of PlsX plays a key topological role in the channeling of acyl substrates from the cytoplasm and into the phospholipid synthesis pathway. These results highlight the importance of spatial organization for the proper functioning of certain biochemical pathways within cells. In addition, they contribute to the field of antibacterial drug discovery, considering the importance of bacterial lipid synthesis as a target for new antimicrobial compounds.

Experimental procedures

Bacterial strains and growth conditions

All bacterial strains used in this study were listed in Table S1. Drug concentrations for selection were: erythromycin, 1 μg ml⁻¹; lincomycin 25 μg ml⁻¹; kanamycin, 50 μg ml⁻¹; chloramphenicol, 5 μg ml⁻¹; spectinomycin, 100 μg ml⁻¹; ampicillin 100 μg ml⁻¹. In general, *B. subtilis* and *Escherichia coli* were grown in Luria-Bertani (LB) broth medium or on plates at 37 °C with aeration.

General molecular techniques

Molecular cloning techniques were performed as described (26). DNA fragments were obtained by PCR using Phusion DNA polymerase (New England Biolabs). Oligonucleotide primers were purchased from Exxtend SRL (Campinas, SP, Brazil) and Thermo Fisher Scientific (São Paulo, SP, Brazil) and are listed in Table S2. PCR products of the expected sizes were purified from gels using a GeneJET™ Gel Extraction Kit (Fermentas), ligated into the pTZ57R/T cloning vector (Fermentas) and transformed into *E. coli* DHα (Table S1). Plasmid DNA was prepared using a GeneJET Plasmid Miniprep kit (Thermo Sci-

entific). Sequencing was carried out using the Big Dye terminator cycle sequencing kit (Applied Biosystems) by the sequencing service of the Departamento de Bioquímica, IQ-USP (SP, Brazil). Transformation of *B. subtilis* was carried out by the method of Dubnau and Davidoff-Abelson (27). The amy phenotype was assayed for colonies grown for 24 h in LB-starch plates by flooding the plates with 1% I₂, KI solution.

Plasmid and strain construction

All plasmids used in this study are listed in Table S1. The *plsX* gene from *B. subtilis* was amplified and inserted into pET24b (T7 promoter) within NdeI and XhoI sites to generate pET24-*plsX*, which expressed BsPlsX with a C-terminal His₆ tag. Site-directed mutagenesis was performed using Phusion Site-directed Mutagenesis Kit (Thermo Fisher Scientific). GFP fusion constructs were designed for fluorescence microscopy. Construction of GFP-PlsX and GFP-PlsX-HBsu into pDR111 vector containing the double or triple fusion within NheI and SphI sites to generate pDR111-*gfp-plsX* and pDR111-*gfp-plsX-hbsu* vectors to integrate in *amyE* locus of *B. subtilis* DS28 strains.

Expression and purification of BsPlsX

Vectors pET24b-*plsX* were used to transform *E. coli* BL21 (DE3) pRIL-competent cells for protein expression. BsPlsX WT and variant mutants were induced with 0.8 mM IPTG for 4 h at 37 °C. Cells were collected by centrifugation (6000 rpm, 4 °C, 15 min), and cell pellets were lysed with a sonicator. Soluble proteins were applied to a Ni²⁺-agarose column and washed with buffer: 50 mM Tris-HCl (pH 7.5), 300 mM NaCl, and 50 mM imidazole. His-tagged proteins were eluted with the same buffer containing 300 mM imidazole. Purified proteins were applied to a P10 desalting column and eluted with buffer A without imidazole. The proteins were stored at −80 °C.

Gel filtration assay

The PlsX WT protein and variant mutants were purified by Ni²⁺ column and then were subjected to gel filtration chromatography using Superdex 200 Increase 10/300 GL resin in a HPLC column (10.0 × 300 mm) and employing the AKTA pure system (GE Healthcare). Proteins were eluted with buffer 50 mM Tris-HCl (pH 7.5), 300 mM NaCl at a flow rate of 0.5 ml/min and elution was monitored at 280 nm.

Fluorescence microscopy

For fluorescence microscopy, overnight precultures of *B. subtilis* were grown in LB medium containing antibiotics diluted in fresh medium without antibiotics to an optical density at 600 nm of 0.05 and grown at 37 °C. At appropriate time intervals, 5 μl of living cells was applied directly on glass slides on a thin film of 1.5% LB-agarose and examined using a ×100 oil immersion objective on a Nikon Eclipse Ti microscope, equipped with GFP BrightLineR Set (Semrock). Images were acquired on a system with the Nikon digital camera (Nikon, Tokyo, Japan). All images were captured using NIS software version 3.07 (Nikon) and fluorescence signals were analyzed using ImageJ software.

Synthesis of palmitoyl-PO₄

The 16:0-PO₄ was synthesized by a new mechanochemical method for the synthesis of acylphosphates.⁶ Palmitoyl chloride (370 μl, 1.2 mmol) was added to a mixture of tripotassium phosphate (K₃PO₄, 297 mg, 1.4 mmol) and 10 mol % tetra-*n*-butylammonium bromide (Bu₄NBr, 40 mg), followed by trituration. Trituration was continued until the reaction was completed as evident from the formation of a solid (about 5 min). The resulting mixture was filtered in a celite pad, and the retained solid was washed with chloroform (3 × 10 ml). The chloroform-soluble material was dried under vacuum to afford 400 mg of 16:0-PO₄ as a white powder (50% yield). ¹H NMR (CDCl₃, 300 MHz), δ 0.92 (t, 3H, J = 6.0 Hz), 1.31 (bs, 26 H), 1.65 (m, 2H), 2.46 (t, 2H, J = 7.0 Hz). ¹³C NMR (CDCl₃, 300MHz), δ 12.5, 21.8, 23.6, 28.6, 28.7, 28.8, 31.1, 34.0, 168.3. ³¹P NMR (CDCl₃) −7.2 ppm MS(ESI), *m/z* = 359 [M + Na]⁺; MS(ESI), *m/z* = 335 [M 2 H]².

PlsX activity assay

The acyltransferase activity was determined according to the procedures described by Lu *et al.* (1). The reaction for acyl chain transfer from 16:0-PO₄ to ACP contained 0.1 M HEPES (pH 7.4), 0.05% Tween 20, 2 mM MgCl₂, 40 μM 16:0-PO₄, and 40 μM ACP. Reaction was started by adding BsPlsX (5–30 μM) and was incubated at 37 °C for 1 h with agitation. Samples were analyzed by conformationally sensitive gel electrophoresis in 15% polyacrylamide gels containing 2 M urea (34, 35) and staining with Coomassie Brilliant Blue.

Western blotting

Samples were prepared as described Weart *et al.* (28). Briefly, harvested cells were resuspended in 50 mM Tris-HCl (pH 7.5), 300 mM NaCl, 1 mM phenylmethylsulfonyl fluoride, 2 mg ml^{−1} of lysozyme. Cells were incubated at 37 °C for 30 min and lysates were normalized by OD₆₀₀ at cell harvest, resuspended in sample buffer, and subjected to SDS-PAGE. GFP-PlsX was detected using the polyclonal rabbit anti-GFP antibody (29). FtsZ was detected using the polyclonal anti-FtsZ antibody (laboratory stock).

Liposome sedimentation assay

Initially, total phospholipids from *B. subtilis* PY79 were extracted with the Bligh and Dyer method (30). Liposomes were made by resuspension of the lipid film in 50 mM Tris-HCl (pH 7.5), 300 mM NaCl. For sedimentation assays, protein and liposomes were mixed at different concentrations in the assay buffer to a final volume of 150 μl. After incubation at room temperature for 30 min, samples were centrifuged at 65,000 rpm for 15 min in a Beckman TLA100 rotor. Supernatants were removed immediately and pellets were resuspended in an equal volume of buffer. Samples from supernatant and pellet were subjected to 12% SDS-PAGE and proteins were detected by Coomassie Blue staining phospholipid vesicles.

DSC experiments

The effects of PlsX on lipid bilayer membranes composed of DMPG, DMPC, and DMPS were analyzed by monitoring the

⁶ Marques Netto, unpublished data.

thermotropic phase behavior of lipid and lipid-protein samples using DSC. The thermograms were recorded in a VP-DSC MicroCal MicroCalorimeter (Microcal, Northampton, MA) using a heating rate of 23.4 °C/h. The analyses of the heat capacity curves were performed using Microcal Origin software (OriginLab Corporation, Northampton, MA). Lipid vesicles were prepared as follows: aliquots of lipid stock solutions in chloroform/methanol, 1:1 (v/v), were added into a glass tube, dried under a N₂ flow, and ultracentrifuged under vacuum for 5 h to remove traces of organic solvent. The resulting lipid film was hydrated in 50 mM Tris-HCl buffer (pH 7.5), 300 mM NaCl at a temperature above the main phase transition of the lipid for 2 h and submitted to six freeze-thaw cycles. LUV were obtained by extruding the preformed multilamellar dispersions using polycarbonate filters with a pore size of 100 nm (Nuclepore Corp., Cambridge, CA). An aliquot of the protein stock solution was added into the LUVs, gently vortexed for few seconds, and kept at either 37 °C (DMPC and DMPG) or 40 °C (DMPS) for 1 h prior to the DSC experiment. The phospholipids were purchased from Avanti Polar Lipids, Inc. (Alabaster, AL) and used without further purification.

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