

# Structural and genetic analyses reveal the protein SepF as a new membrane anchor for the Z ring

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A key step in bacterial cell division is the polymerization of the tubulin homolog FtsZ at midcell. FtsZ polymers are anchored to the cell membrane by FtsA and are required for the assembly of all other cell division proteins. In Gram-positive and cyanobacteria, FtsZ filaments are aligned by the protein SepF, which in vitro polymerizes into large rings that bundle FtsZ filaments. Here we describe the crystal structure of the only globular domain of SepF, located within the C-terminal region. Two-hybrid data revealed that this domain comprises the FtsZ binding site, and EM analyses showed that it is sufficient for ring formation, which is explained by the filaments in the crystals of SepF. Site-directed mutagenesis, gel filtration, and analytical ultracentrifugation indicated that dimers form the basic units of SepF filaments. High-resolution structured illumination microscopy suggested that SepF is membrane associated, and it turned out that purified SepF not only binds to lipid membranes, but also recruits FtsZ. Further genetic and biochemical analyses showed that an amphipathic helix at the N terminus functions as the membrane-binding domain, making SepF a unique membrane anchor for the FtsZ ring. This clarifies why *Bacillus subtilis* grows without FtsA or the putative membrane anchor EzrA and why bacteria lacking FtsA contain SepF homologs. Both FtsA and SepF use an amphipathic helix for membrane binding. These helices prefer positively curved membranes due to relaxed lipid density; therefore this type of membrane anchor may assist in keeping the Z ring positioned at the strongly curved leading edge of the developing septum.

One of the first steps in bacterial cell division is the polymerization of the conserved protein FtsZ at midcell. FtsZ shares structural homology with eukaryotic tubulin and uses GTP to polymerize into filaments close to the cell membrane. These filaments then assemble into a ring-like structure, the Z ring, which recruits other proteins needed for the division septum (1). Several cell division proteins support the formation of a stable Z ring, such as ZapA that forms cross-links between FtsZ filaments (2, 3). In Gram-positive bacteria and cyanobacteria, the protein SepF also stimulates bundling of FtsZ polymers (4–6). Electron microscopic (EM) studies have shown that SepF assembles into large and regular protein rings with diameters of about 50 nm. In vitro, these rings are able to bundle FtsZ protofilaments into long tubular structures (4). Furthermore, SepF is essential for the synthesis of regular and smooth division septa (7–9).

The Z ring is associated with the cell membrane and the best-characterized membrane anchor is the conserved protein FtsA. This protein binds to FtsZ directly and contains a C-terminal amphipathic helix that binds to lipid bilayers in a membrane-potential-dependent manner (10–12). In *Escherichia coli*, FtsA is essential, but in *Bacillus subtilis*, *ftsA* can be deleted, although this affects Z-ring formation and cells become elongated. Because *B. subtilis* can grow without FtsA, there must be another protein that links the Z ring to the cell membrane. The essential *E. coli* cell division protein ZipA binds to FtsZ and contains an

N-terminal transmembrane domain (13). The necessity for ZipA can be bypassed by a gain-of-function mutation in FtsA (14). Gram-positive bacteria contain EzrA, which shows a similar topology to that of ZipA, with an N-terminal transmembrane helix and a large C-terminal domain that binds to the FtsZ C terminus (15). It therefore seemed likely that EzrA functions as an alternative membrane anchor for the Z ring in *B. subtilis* when FtsA is absent.

Comparison of the SepF amino acid sequence against protein databases did not reveal conserved motifs that provide clues to potential molecular mechanisms. To gain insight into the residues that are important for FtsZ interaction, we used a yeast two-hybrid screen. This revealed that the conserved C-terminal part of SepF comprises the FtsZ binding site. We were able to obtain diffracting crystals of this domain that revealed a tight dimer structure. Site-directed mutagenesis indicated that the protein polymerizes as units of dimers. Although the FtsZ interaction domain constitutes only 60% of the full-length protein, it appeared to be also sufficient for the formation of the large protein rings. This raised the question of what the function is of the N-terminal domain of SepF. Interestingly, high-resolution fluorescent-light microscopy suggested that SepF associates with the cell membrane. Subsequent analyses indicated that the membrane-binding domain resides in the N-terminal domain of SepF. Biochemical experiments using purified SepF showed that the protein binds specifically to lipid membranes. These data

## Significance

**A key step in bacterial cell division is the polymerization of FtsZ at midcell into a ring-like structure. This so-called Z ring forms a scaffold for the other cell division proteins. FtsA anchors the Z ring to the cell membrane; however, many bacterial species do not have FtsA. Here, we show that the conserved protein SepF, which forms large protein rings, also functions as a membrane anchor for the Z ring. We determined the molecular structure of the FtsZ-binding and ring-forming domain of SepF and show that the membrane-binding domain is located at the very beginning of the protein. These results explain why FtsA and SepF can fulfill similar functions in bacterial cell division.**

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Data deposition: The atomic coordinates and structure factors have been deposited in the Protein Data Bank, [www.pdb.org](http://www.pdb.org) (PDB ID code 3ZIE).

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suggested that SepF might function as a membrane anchor for the Z ring. Interestingly, we were able to delete both *ftsA* and the putative FtsZ-membrane anchor *ezrA* in *B. subtilis* and this did not affect viability. However, it turned SepF into an essential protein. Indeed, using purified SepF we were able to show that the protein recruits FtsZ to artificial lipid membranes. We conclude that SepF is a unique membrane anchor for FtsZ and the Z ring.

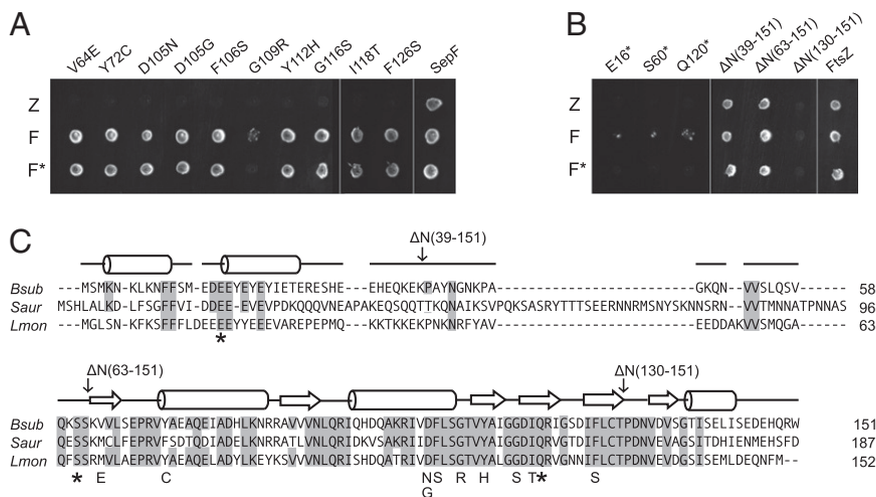
## Results

**Localization of FtsZ Interaction Mutants.** SepF binds directly to the C-terminal domain of FtsZ (6, 16). However, the protein has no apparent homology to proteins of known function and the amino acid sequence does not indicate a clear docking site for FtsZ. In earlier studies it was shown that FtsZ and SepF show a strong and specific interaction in yeast two-hybrid assays (8, 17). This assay was used to identify amino acid residues in SepF that are important for FtsZ binding. Mutations in SepF were introduced by means of error-prone PCR. SepF also shows strong self-interaction, and the mutants were tested for this as well (Fig. 1A and B). Thirteen different mutants were isolated, of which 10 were amino acid substitutions and 3 were nonsense mutations, the latter resulting in C-terminal truncations at amino acids 16, 60, and 120, respectively. To confirm that the SepF mutants show a defect in FtsZ interaction in *B. subtilis* cells, the mutants were expressed in an *ftsA* deletion strain. Previous work has shown that *sepF* becomes essential when *ftsA* is deleted in *B. subtilis* (17). Indeed, all mutants were unable to support growth in a *B. subtilis* *ftsA* mutant background (Fig. S1). Fig. 1C shows an amino acid sequence alignment of SepF homologs from three different bacterial species. An extensive ClustalW alignment can be found in Fig. S2. The 10 missense mutations are all located in the conserved C-terminal part of the protein. To further analyze this, three N-terminal truncation mutants were constructed, lacking the first 39 aa, 63 aa, or 130 aa, respectively (Fig. 1B). Interestingly, it was possible to delete the complete 63-aa-long N terminus without affecting FtsZ interaction and self-interaction in our assay. This suggests that the C-terminal part of the protein

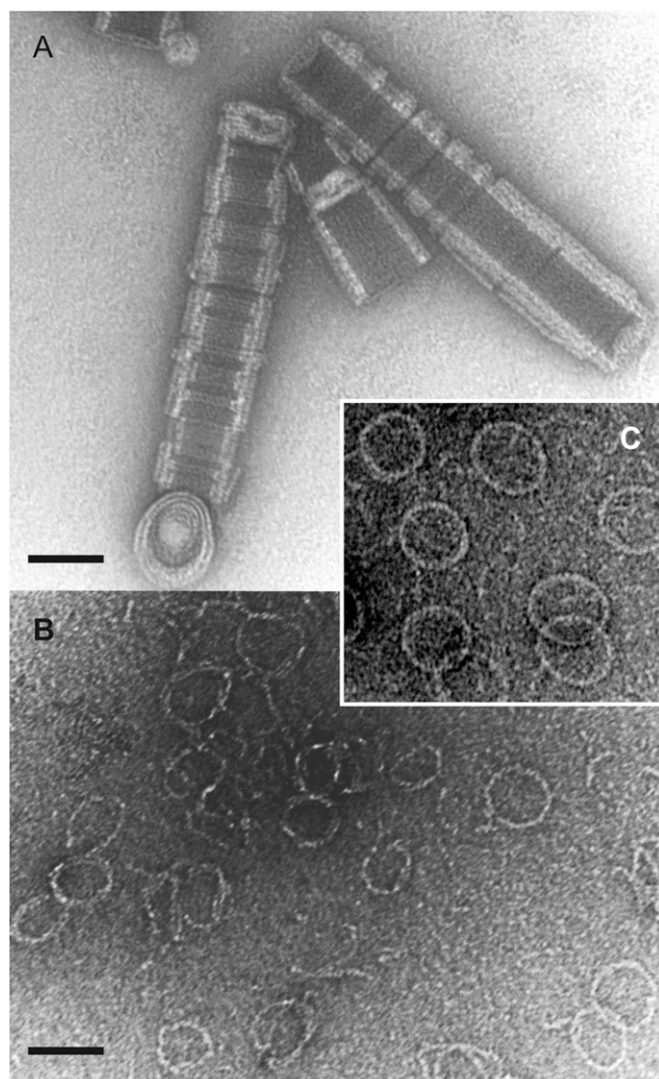
constitutes a separate domain carrying the FtsZ-interaction and self-interaction functionalities.

**Ring-Forming Domain.** We were unable to obtain diffraction-quality crystals of full-length SepF from *B. subtilis* or from other bacteria. One of the reasons could be that the protein contains unstructured domains that hamper the crystallization process. Mild proteolytic treatment is able to remove such domains, and indeed a controlled incubation with  $\alpha$ -chymotrypsin yielded a truncated protein that formed crystals, which diffracted to 2 Å resolution. Mass spectrometry and N-terminal sequencing revealed that the crystallized protein consists of residues 57–151, thereby neatly encompassing the C-terminal FtsZ-binding region identified in the yeast two-hybrid screen. EM analysis of this truncated SepF protein revealed tubular structures of ~50 nm in diameter, which were made up of stacks of SepF rings (Fig. 2A). Very high pH conditions (~pH 10) resolved the structures into individual rings (Fig. 2B). Thus, the C-terminal domain of SepF is responsible for the polymerization of SepF into large rings.

**Crystal Structure.** To elucidate the atomic structure, two additional structures of SepF-like proteins were solved by multi-wavelength anomalous dispersion (MAD) phasing, one from *Archaeoglobus fulgidus* and one from *Pyrococcus furiosus*, and used for molecular replacement of the *Bacillus* protein. The resulting crystal structure of the SepF C-terminal domain is shown in Fig. 3. The structure comprises residues Ser-61 to Ser-140. Because the protein spans residues 57–151, it means that the first 4 aa and the last 11 aa are disordered in the crystal. The crystal structure shows a tight dimer, with monomers that contain a compact  $\alpha/\beta$ -sandwiched arrangement of two  $\alpha$ -helices stacked against a five-stranded  $\beta$ -sheet (Fig. 3A and B). The dimer interface is formed by the  $\beta$ -sheets. Interestingly, the C-terminal  $\beta$ -strand from each monomer joins the  $\beta$ -sheet of the other monomer, such that it is parallel to its N-terminal  $\beta$ -strand. The tight packing of the dimer suggests that this is the functional unit in SepF polymers. In fact, many of the conserved residues are located at the interface between the monomers on the opposing  $\beta$ -sheets (Fig. 3C). Moreover, the *A. fulgidus* and



**Fig. 1.** Yeast two-hybrid screen for SepF-FtsZ interaction mutants. (A) SepF mutants were created using error-prone PCR, and FtsZ-interaction deficient mutants were selected using the yeast two-hybrid assay. Diploid colonies were obtained by mating PJ69-4 $\alpha$  strains harboring pGBT9 derivatives containing *sepF* mutants, and wild-type *sepF*, with PJ69-4A strains that harbored pGAD24 derivatives containing *ftsZ* (Z), wild-type *sepF* (F), or the corresponding *sepF* mutants (F\*). (B) Yeast two-hybrid interactions of the nonsense mutants and the different N-terminal deletion constructs. Remaining amino acid regions are indicated between parentheses. A and B are discontinuous images of the same plate. (C) Amino acid sequence alignment of SepF proteins, from *B. subtilis* (*Bsub*), *Staphylococcus aureus* (*Saur*), and *Listeria monocytogenes* (*Lmon*), using ClustalW. Secondary structure prediction (using PSIPRED) for *B. subtilis* SepF is shown above the sequences, whereby  $\alpha$ -helices and  $\beta$ -sheets are indicated by rods and arrows, respectively. Conserved amino acids are shaded. Yeast two-hybrid mutations are indicated.



**Fig. 2.** Negatively stained electron micrographs of purified chymotrypsin-treated SepF used for crystallization. (A) Under neutral pH conditions (50 mM Tris-HCl, pH 7.0) this protein forms large tubes made up of stacked 50-nm diameter rings. (B) Under basic pH conditions (50 mM glycine, pH 10.0) individual rings become visible. (C) Normal full-length SepF. (Scale bars, 50  $\mu\text{m}$ .)

*P. furiosus* SepF-like proteins show the same dimer packing in their crystals (Fig. S3; structure statistics are listed in Table S1).

**Polymerization of Dimers.** In Fig. 3D the yeast two-hybrid mutations from Fig. 1 are highlighted in the crystal structure. Several mutations that affect FtsZ interaction are located on the surface of the  $\alpha$ -helices. This is surprising because there is no clear pocket that could function as a docking site for the C-terminal domain of FtsZ on the dimer structure. However, the crystal packing reveals a side-by-side chain arrangement of very tightly packed dimers (Fig. 4), and in this arrangement an elongated pocket exists that could accommodate the FtsZ C terminus, which most likely forms an  $\alpha$ -helix (18, 19). In the SepF polymer, the  $\alpha$ -helices on the outside of dimers form the interfaces between the dimers. It seems highly likely that the SepF rings are polymers of SepF dimers. To determine whether the  $\alpha$ -helices are indeed involved in multimerization, we identified residues that are important for such interactions. At the interface between adjacent dimers there are two glycine residues (Gly109)

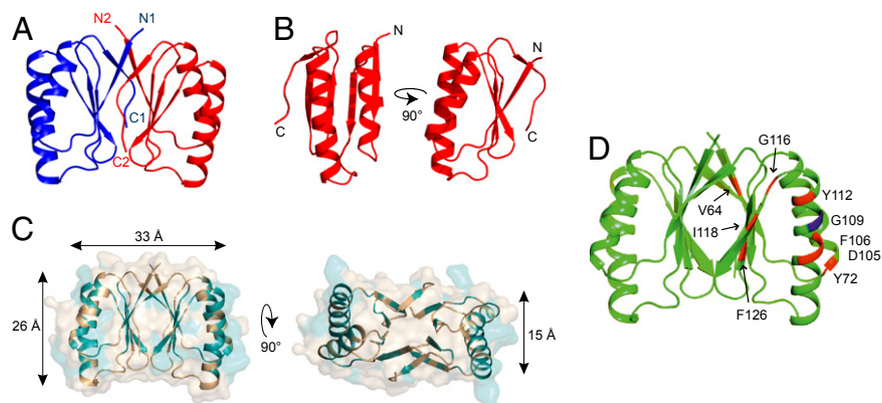
contacting each other, one contributed by each monomer (Fig. 4). The lack of side chains on this glycine leads to an unusually close packing of the two longest helices in the four-helix bundle. In fact, Gly109 is highly conserved (Fig. S2B), and the yeast two-hybrid experiments showed that an arginine substitution destroys self-interaction (Fig. 1A). Importantly, the crystal lattices of the *A. fulgidus* and *P. furiosus* proteins do not show the polymeric arrangement, and these proteins do not contain this conserved glycine residue. It should be mentioned that these proteins have currently no known function and were picked solely because of their sequence similarity to SepF. To confirm the role of Gly109 in multimerization, we replaced it with a lysine and crystallized the mutant protein. The G109K substitution has a dramatic effect, because in the new crystals the protein filaments are absent, but dimers are still present (Fig. 5A). EM analysis showed that this mutant completely abolishes ring formation. This was not a consequence of structural differences between the wild type and the G109K mutant proteins, because superposition of the two monomers shows close to no variation in the conformation of the C- $\alpha$  backbone between the two structures (Fig. S4B).

The G109K mutation should not affect dimerization. To test this, gel filtration and analytical ultracentrifugation were performed using full-length SepF with and without the G109K substitution. The gel filtration data indicate that, unlike wild-type SepF that elutes in the void volume, the G109K mutant is considerably smaller in size (Fig. 5B). Using three different protein concentrations, analytical ultracentrifugation assays show that the calculated molecular weight of the G109K mutant ranges between 30 kDa and 40 kDa, which is close to the molecular weight of the dimer (34 kDa, Fig. 5C). These experiments support the conclusion that SepF forms polymers of dimers and that the outside  $\alpha$ -helices form the polymerization interfaces.

One aspect that remains uncertain is the precise location of the FtsZ binding site. Differential scanning calorimetry using truncated SepF protein and FtsZ gave no conclusive results, and mixing the truncation with FtsZ did not result in tubular structures other than what is shown in Fig. 2A. Possibly, the close stacking of rings in these tubular structures interferes with FtsZ interaction. Yeast two-hybrid data show that the G109R mutation also abolishes FtsZ binding (Fig. 1A). The same was observed with the G109K mutation, using an affinity chromatography assay (Fig. S4C), suggesting that the interacting  $\alpha$ -helices are part of the FtsZ binding site.

A Dali structural similarity search (20) revealed that the C-terminal domain of SepF shows some resemblance to the *E. coli* protein TusA [YhhP, Protein Data Bank (PDB) ID 1dcj] (21). Because TusA is involved in thiomodification of tRNAs, it is unlikely that this similarity has any functional relevance (22).

**An *ftsA ezrA* Double Mutant Is Viable.** Because the C-terminal domain of SepF is involved in FtsZ interaction and ring formation, the question arises what the function is of the 60-aa-long N-terminal region. A surprising genetic result provided a possible clue. Superficially, the overall architecture of the cell division protein EzrA shows some similarity to that of the *E. coli* cell division protein ZipA, with an N-terminal transmembrane helix preceding a much larger cytoplasmic FtsZ-binding domain. Therefore, EzrA might be able to anchor FtsZ to the membrane, which would explain why *B. subtilis* cells are able to grow when the main membrane anchor of the Z ring, *ftsA*, is deleted. An *ezrA*-null mutant forms slightly longer cells (~20%) compared with wild-type cells (23), but an *ftsA* null mutant has problems dividing and forms long filamentous cells (24). Interestingly, when we transformed an *ftsA* deletion into a  $\Delta\text{ezrA}$  background, the resulting double mutant grew well and did not show a strong cell division defect (Fig. 6). We concluded that there must be another protein that links FtsZ to the cell membrane. Certain Gram-positive bacteria, such as Mycobacteria, lack FtsA but they



**Fig. 3.** Crystal structure of *B. subtilis* SepF residues 61–140. (A) The crystal structure reveals a dimer whereby the C-terminal  $\beta$ -strand of one monomer (C1, C2) joins the  $\beta$ -sheet of the other monomer and is parallel to its N-terminal  $\beta$ -strand (N1, N2). Structure statistics are listed in Table S1. (B) Monomer of SepF C-terminal domain. (C) Conserved residues are highlighted in teal on the SepF dimer structure. Most conserved residues are located at the interface between monomers ( $\beta$ -sheets) and the interface between dimers ( $\alpha$ -helices). Dimensions indicated by arrows are based on the protein backbone. (D) FtsZ-interaction mutants from the yeast two-hybrid screen are indicated on the crystal structure. G109 residue is indicated in blue.

do contain a SepF homolog. In addition, deleting both *ftsA* and *sepF* is lethal in *B. subtilis*, and so is the *ezrA sepF* double knockout (8, 17). These data might suggest that SepF can complement the function of FtsA. However, this is possible only if SepF binds to the cell membrane. Previously, it was shown that a SepF-GFP fusion binds to the Z ring and that the SepF-GFP signal becomes dispersed when FtsZ is depleted (8, 17). In fact, repetition of this experiment, but this time using high-resolution structured illumination microscopy (SIM), showed that the SepF-GFP signal clearly overlaps with the fluorescent signal of the cell membrane when FtsZ is depleted (Fig. S5), suggesting that SepF has indeed affinity for the cell membrane.

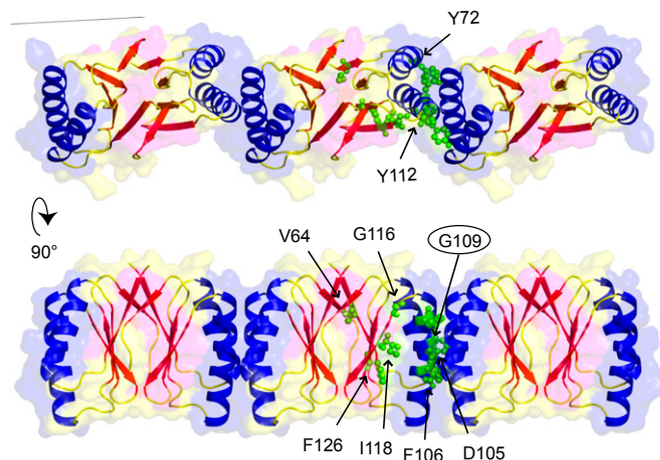
**Purified SepF Binds to Lipid Membranes.** SepF might interact with a membrane protein or SepF is a peripheral membrane protein and has an affinity for lipid bilayers itself. To test this, purified SepF was mixed with liposomes and after centrifugation the amount of SepF in the pellet and supernatant was determined. Excess BSA and 200 mM KCl were included to reduce non-specific interactions. Approximately four times more SepF is pelleted when liposomes are present, suggesting that SepF has some affinity for lipid membranes (Fig. S6A). However, even without liposomes, a substantial amount of SepF is spun down, presumably through polymerization. Therefore, a more suitable sucrose flotation experiment was performed. SepF and liposomes were loaded at the bottom of a sucrose gradient, and during centrifugation the liposomes migrated to a lower density, upward (lower sucrose concentration). Again BSA and 200 mM KCl were included throughout the gradient to increase specificity. As shown in Fig. 7A, a fraction of SepF clearly comigrates with liposomes, whereas purified GFP does not, lending further support to the idea that SepF can bind directly and specifically to the cell membrane. Another compelling indication was the observation that fluorescently labeled liposomes mixed with purified SepF started to aggregate and often fused into larger vesicles (Fig. S6C).

SepF forms remarkably large and uniform protein rings. To examine whether lipid membranes would influence these structures, SepF was mixed with small liposomes, negatively stained, and examined by EM. The presence of SepF resulted in large amorphous clumps of liposomes, as was observed with fluorescence light microscopy, but isolated liposomes were often strongly deformed, giving rise to elongated structures (Fig. 7B). Deformation of liposomes is regularly observed with peripheral membrane proteins, because insertion of the membrane-binding

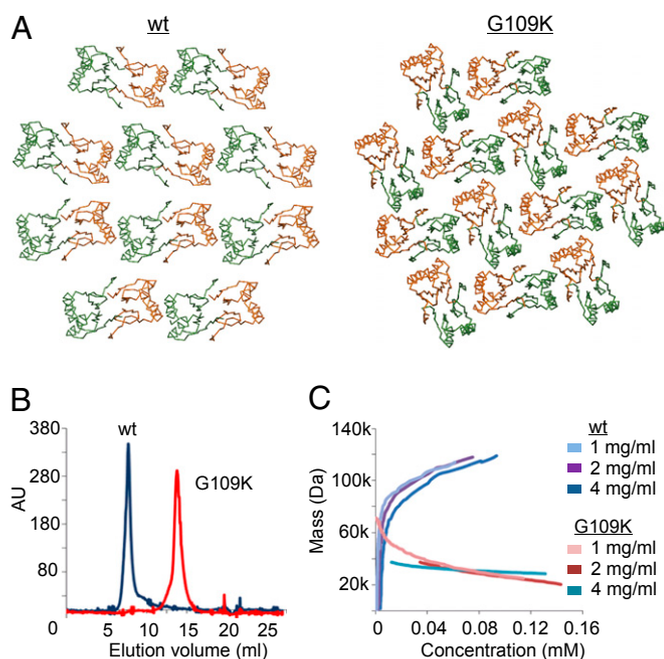
domain displaces lipid molecules that induce local curvature of the lipid bilayer (25). Interestingly, some SepF rings contained a liposome inside (Fig. 7B).

**Membrane-Binding Domain.** The SIM images suggested that binding to FtsZ is not required for the association of SepF with the membrane. To corroborate this, we tested the binding of two previously described mutations, A100V and F126S, which impair FtsZ binding but do not block protein ring formation (4). Indeed, both proteins comigrated with the liposome fraction in sucrose density gradients and induced liposome clustering (Fig. 7A and Fig. S6C).

To determine whether the membrane-binding domain is located in the main ring-forming part of SepF, we examined whether the SepF truncation used for crystallography showed any affinity for liposomes. As shown in Fig. 7A, the C-terminal domain (amino acids 61–139) does not comigrate with liposomes in sucrose gradients, and is not able to cluster liposomes (Fig. S6C). This implies that the membrane-binding domain resides in the 60-aa-long N-terminal region of SepF. In fact, prediction



**Fig. 4.** Polymer structure of SepF revealed by crystal packing. Crystal packing reveals a tight packing of dimers that assemble to polymers via a close interface between their outer helices. FtsZ-interaction mutants from the yeast two-hybrid screen are highlighted, as well as glycine residue G109 (encircled).



**Fig. 5.** G109K mutation diminishes polymerization. (A) Crystal lattice of wild-type SepF and crystal lattice of the G109K mutant showing disruption of the interdimer interface that leads to polymerization. (B) Gel-filtration profiles of full-length SepF wild type and the G109K mutant. (C) Analytical ultracentrifugation run at 21,000 rpm. Diagram of mass against concentration indicates that the G109K mutant has a molecular weight close to the theoretical mass of a SepF dimer (34 kDa). Concentrations used are indicated by different colors.

using Amphipaseek (26) suggested that amino acids 2–12 form a putative amphipathic helix (Fig. 8A), and this region is highly conserved in the N-terminal domain (Fig. S24). Such helical structures are known to insert into lipid membranes and are used by many peripheral membrane proteins as membrane-binding domains, including the cell division proteins FtsA, MinD, and MreB (11, 27–29). To examine whether the first 13 aa of SepF form an amphipathic helix upon interaction with lipid bilayers, this small peptide was synthesized and CD spectra with and without liposomes were determined. As shown in Fig. 8A, SepF1–13 is structurally unfolded in solution but exhibits a clear shift to an  $\alpha$ -helical fold in the presence of liposomes, a transition that is characteristic of membrane-binding amphipathic helices (30, 31).

To test whether this amphipathic helix is able to recruit proteins to the membrane *in vivo*, we constructed a C-terminal GFP fusion with the first 25 aa of SepF (N25-GFP) and analyzed the localization of the fusion protein in *B. subtilis* cells. As shown in Fig. 8B, the fluorescence signal is primarily diffuse although in some cells a weak membrane signal can be observed at division sites (arrowheads). Because SepF forms large polymers, it is likely that a weak membrane affinity of the monomers is compensated by multimerization. In fact, the G109K mutation that blocks polymerization also abolishes comigration with liposomes in sucrose gradients and clustering of liposomes (Fig. 7A and Fig. S6C). Indeed, introduction of this mutation into a SepF-GFP reporter fusion [SepF(G109K)-GFP] resulted in a diffuse fluorescent signal (Fig. 8B). Previously, Szeto et al. (32) have shown that the 12-aa membrane-binding amphipathic helix of *E. coli* MinD could target GFP to the cell membrane only when they fused this domain to the c-Jun leucine zipper. When the c-Jun leucine zipper dimerization domain was introduced between the first 25 aa of SepF and GFP (N25-JunLZ-GFP), membrane

binding became very clear (Fig. 8B). Interestingly, when only the first 13 aa of SepF were fused to GFP (N13-GFP), the fusion product also localized to the cell membrane, even without the presence of a dimerization domain (Fig. 8B). Mutating the hydrophobic leucine residue of the amphipathic helix to an aspartate [N13(L7D)-GFP] abolished membrane binding (Fig. 8B). These data strongly suggest that the amphipathic helix at amino acid positions 1–12 constitutes the membrane targeting domain of SepF. To further corroborate this conclusion, a SepF variant was purified, lacking the first 13 aa ( $\Delta$ N13). As expected, this mutant showed no lipid interaction (Fig. 7A). When the L7D mutation was introduced into full-length SepF, lipid binding was also abolished (Fig. 7A).

To support these findings, the importance of the amphipathic helix for the function of SepF in cell division was tested. It has been shown that SepF expressed *in trans* is able to support growth in the absence of both *ftsA* and wild-type *sepF* (17). However, we were unable to delete both *sepF* and *ftsA* when a SepF mutant was expressed that lacked the first 13 aa, indicating that the membrane binding of SepF is essential to support cell division. This result was not a consequence of an increased instability of the mutant protein (Fig. S6D). Interestingly, the replacement of the native amphipathic helix with the membrane-binding amphipathic helix of *B. subtilis* MinD restored growth in a strain lacking *ftsA* and wild-type *sepF* (Fig. 8C), demonstrating the key membrane-binding function of the N terminus of SepF. It should be mentioned that the *in vitro* binding experiments were performed using *E. coli* polar lipid extract. However, SepF also binds to liposomes formed from *B. subtilis* lipid extract (Fig. S6E).

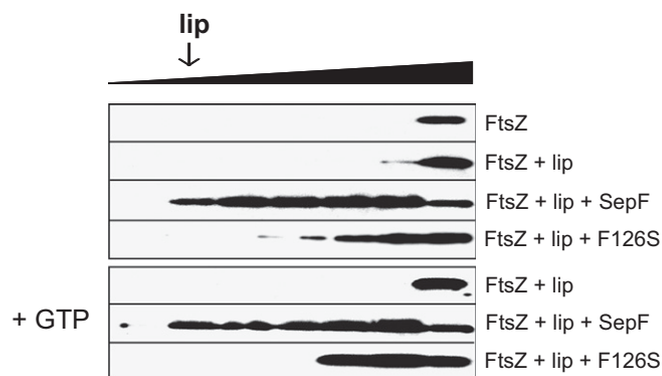
**SepF Recruits FtsZ to Lipid Membranes.** If SepF functions as a membrane anchor for the Z ring, then the protein should be able to recruit FtsZ to lipid membranes. To test this, FtsZ was included in the sucrose gradient centrifugation assays. As shown in Fig. 9, FtsZ remains in the bottom fraction of the gradient when mixed with liposomes. However, when SepF is included, FtsZ migrates with the liposomes. As a control, the experiment was repeated with mutant F126S that shows a reduced affinity for FtsZ but binds normally to liposomes. Indeed, this SepF variant is unable to recruit FtsZ to the liposome-containing gradient fraction (Fig. 9). FtsZ polymerizes into long protofilaments in the presence of GTP. However, previous work has shown that the binding of FtsZ to SepF does not depend on GTP (4, 6), and when the flotation experiments were performed in the presence of GTP, similar results were obtained (Fig. 9).

It has been shown that SepF rings assemble FtsZ protofilaments into very long 50-nm-wide tubular structures (4). In fact, these tubular structures were easily observed on EM grids when liposomes were added to a mixture of purified SepF and FtsZ, (Fig. S7). The main difference is that there are many more FtsZ-SepF tubules when liposomes are present. Charged surfaces,



**Fig. 6.** Viability of *ftsA ezrA* double mutant. Images show membrane-stained wild-type,  $\Delta$ ftsA,  $\Delta$ ezrA, and  $\Delta$ ftsA  $\Delta$ ezrA double-mutant *B. subtilis* cells. (Scale bars, 3  $\mu$ m.)





**Fig. 9.** SepF recruits FtsZ to lipid membranes. Recruitment of FtsZ to liposomes was analyzed using sucrose gradient centrifugation. FtsZ was detected using Western blotting. The F126S mutation reduces SepF-FtsZ binding (4). The presence of GTP (+GTP) has no effect on lipid interaction. Liposomes were made from *E. coli* polar lipid extract.

that the cell membrane thickness approximates to  $\sim 5$  nm, then the average width of septa is more in the range of 43 nm, which comes close to the inner diameter of SepF rings. This would mean that arcs of SepF polymers would fit neatly on top of the leading edge of the developing septum (Fig. 10). This model also better accommodates the fact that FtsZ protofilaments become curved upon GTP hydrolyses. In previous work it was shown that this bending of protofilaments separates the ends of FtsZ-SepF tubules, resulting in splayed tips containing highly curved FtsZ filaments (4).

**Localization by Curved Membrane Area.** Finally, both SepF and FtsA use an amphipathic helix to bind to the cell membrane. These protein structures bind stronger to convex (positively curved) membrane surfaces because curvature creates space in between lipid molecules (25). By far the most prominent convex membrane area in a bacterial cell appears at the leading edge of a developing division septum (Fig. 10). Thus, it seems likely that both FtsA and SepF have an intrinsic binding preference for this area in the cell. This characteristic might help to keep the Z ring positioned at the edge of the growing septum during cell division.

## Materials and Methods

**Strains and General Methods.** Yeast, *E. coli*, and *B. subtilis* strains and primers used in this study are listed in Tables S2 and S3, respectively. Molecular cloning, PCR reactions, yeast transformation and mating experiments, and *E. coli* transformations were carried out using standard techniques. *B. subtilis* chromosomal DNA for transformation and PCR reactions was purified as described before (35). Transformation of competent *B. subtilis* cells was performed using an optimized two-step starvation procedure based on the method of Anagnostopoulos and Spizizen (36, 37). Selection of transformants was carried out on nutrient agar supplemented with 100  $\mu$ g/mL ampicillin for *E. coli* and, for *B. subtilis*, 4  $\mu$ g/mL erythromycin, 50  $\mu$ g/mL spectinomycin, 5  $\mu$ g/mL tetracycline, or 0.5  $\mu$ g/mL phleomycin.

**Construction of Plasmids for Protein Purification.** pMal-C2X (New England Biolabs) was used to clone and purify variants of SepF used in this study as N-terminal MBP-SepF fusions. Using *B. subtilis* 168 genomic DNA as template, the  $\Delta$ N13 SepF truncation (oligos inc22/inc6) was amplified, digested with SmaI/XbaI, and ligated into pMal-C2 linearized with XmnI/XbaI, resulting in plasmid pHJ5106 (MBP-SepF $\Delta$ N13). The L7D and G109K exchanges were introduced into pNC12 with quick-change mutagenesis, using oligos H5492/H5493 (L7D) and oligos inc40/inc41, resulting in plasmids pHJ5107 and pNC13.

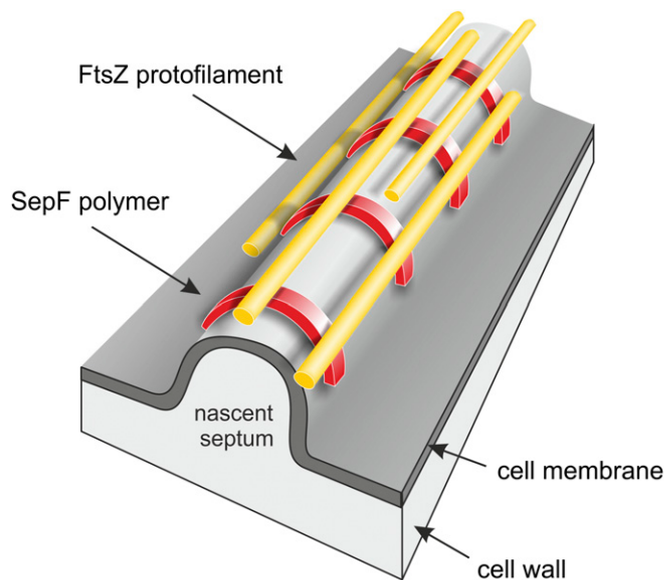
**Construction of Plasmids and Strains for Complementation Assays.** For the construction of the *B. subtilis* *ftsA* mutant containing different SepF mutants, *sepF* genes were amplified from the pGBT9-*sepF* derivatives with oligos *sepF*-mutIn-f2 and pGBT9-mutIn-r. The upstream region, containing the 5' end of the *sepF* coding sequence, and the downstream region of *sepF*

were amplified from *B. subtilis* 168 genomic DNA, using the oligo sets *sepF*-mutF-f/*sepF*-mutF-r2 and *ylmG*-rPCR-f/*sepF*-mutB-r, respectively. The chloramphenicol-resistant cassette was amplified from pDLT3 (38) with oligos rPCR-CmF2 and rPCR-CmR(-ter). The four DNA fragments were then fused together by recombinant PCR, using the oligos *sepF*-mutF-f and *sepF*-mutB-r. The *sepF* mutations were introduced into *B. subtilis* MD194, which contains a complete deletion of the *sepF* gene, by transformation with the PCR products. To test functionalities of the SepF mutants, the *sepF* mutations were then introduced into a *B. subtilis* MD197 mutant strain, which contains a complete *sepF* deletion, and a xylose-inducible *ftsA* (*amyE*::*P<sub>xyI</sub>*-*ftsA*,  $\Delta$ *ftsA*::*erm*,  $\Delta$ *sepF*::*spec*), by transformation with the genomic DNA extracted from the resultant *sepF* mutant strains.

The strains encoding SepF without amphipathic helix ( $\Delta$ AH, strain HS230) and the native AH replaced with AH of *B. subtilis* MinD (strain HS232) were constructed by PCR amplification of *P<sub>xyI</sub>* promoter (oligos H5495/H5497 for  $\Delta$ AH, oligos H5495/H5498 for AH<sub>MinD</sub>) and SepF fragments (oligos H5500/H5501 for  $\Delta$ AH-SepF, oligos H5500/H5503 for AH<sub>MinD</sub>-SepF) followed by insertion into *aprE spec* delivery vector, using the Stargate cloning system (IBA) and direct transformation into *B. subtilis* 168. The correct integration in the *aprE* locus and the sequence of the constructs were verified by PCR amplification followed by sequencing.

**Construction of Plasmids and Strains for Microscopy.** The strains encoding N-terminal fragments of SepF fused to GFP were constructed by amplification of *sepF*1-39 (N13) and *sepF*1-75 (N25), using oligos H5478/H5480 and H5478/H5481, respectively. The amplified products were fused with the plasmid pSG1154 linearized with oligos H509/H5479, using the InFusion HD cloning kit (Clontech), resulting in plasmids pHJ5108 (N13-GFP) and pHJ5109 (N15-GFP). The homologous sequences required for InFusion cloning were designed so that a flexible SGSGSGS linker is positioned between the SepF fragments and GFP. This linker also contains a unique BamHI restriction site, which was used to insert *junLZ* amplified with oligos H5484/H5485 from plasmid pT537 into pHJ5109, resulting in plasmid pHJ5110. The plasmid pHJ5111 encoding N13-GFP with L7D exchange was constructed with quick-change mutagenesis of plasmid pHJ5108, using primers H5492/H5493. The plasmid pNC14 encoding SepFG109K-GFP was constructed with quick-change mutagenesis of plasmid pFG1 (8), using primers inc40/inc41. The plasmids were transformed into *B. subtilis* 168, resulting in the corresponding strains HS206, HS207, HS208, HS223, and NC21.

**Yeast Two-Hybrid Assay.** For library screening, *Saccharomyces cerevisiae* AH109 was used as a host, and *S. cerevisiae* PJ69-4A and PJ69-4 $\alpha$  were used for mating experiments. To construct a *sepF* mutant library in pGBT9



**Fig. 10.** Schematic representation of SepF binding model. SepF polymers (red) bind as arcs onto the convex leading edge of the nascent division septum and bundle FtsZ protofilaments (yellow). See main text for explanation of the model.

(Clontech), the *sepF* gene was amplified from pENTR-*sepF* (17) with oligos pDONR-F and pDONR-R to attach *attL1* and *attL2* sequences at the 5' and 3' ends, respectively, necessary for cloning with the Gateway cloning technique (Life Technologies, previously named Clontech). Random mutations were introduced by using Ex Taq DNA polymerase (Takara) (17, 39). The PCR product was then recombined with pBDGW, using the LR Gateway cloning reaction, and the reaction mixture was transformed to *E. coli*. Plasmids from  $\sim 1.2 \times 10^5$  transformants were collected. *S. cerevisiae* AH109 harboring pGAD424-ftsZ was transformed with the plasmid library, and transformants were selected on synthetic defined medium without tryptophan and leucine in the presence of X- $\alpha$ -gal, to monitor interactions between FtsZ and SepF. The pGBT9 derivatives containing mutant *sepF* genes were selected using the *E. coli* tryptophan auxotroph KCC8 strain and the mutations were identified by sequencing.

To check the N-terminal deletions of SepF in the yeast-two hybrid assay, *sepF* encoding  $\Delta N(39-151)$ ,  $\Delta N(63-151)$ , and  $\Delta N(130-151)$  regions were amplified with the oligo sets *sepF*- $\Delta N1$ -*gwF/sepFgwR*, *sepF*- $\Delta N2$ -*gwF/sepFgwR*, and *sepF*- $\Delta N3$ -*gwF/sepFgwR*, respectively, and then recombined with pDONR201 by the BP Gateway reaction. The resultant plasmids were then recombined with pGBT9 or pGAD424, using the LR Gateway reaction.

Interactions of the SepF mutants with wild-type SepF and self-interactions were tested by mating experiments, using the following plasmids. The mutated *sepF* genes from selected pGBT9 derivatives (pGBT9-*sepFm01* to -*sepFm15*) were amplified with oligos Adapter-*attB1* and Adapter-*attB2*, and the products were integrated into pDONR201 (Life Technologies), using the BP Gateway cloning technology. The inserts were then transferred into pGAD424 by LR Gateway reaction.

**Cloning and Protein Purification for Crystallography.** The genes encoding *B. subtilis* SepF (GI: 238054369) and residues 37–122 of *A. fulgidus* SepF-like protein (GI: 11498388) and residues 49–134 of *P. furiosus* SepF-like protein (GI: 33359575) were amplified from genomic DNA by PCR, using forward oligos containing an NdeI restriction site and reverse oligos carrying a BamHI restriction site. The PCR product was digested with NdeI and BamHI before ligation with plasmid pHis17. A single point mutation, Ile-105 to Met, was introduced in the sequence of the *A. fulgidus* protein, using a QuikChange Mutagenesis Kit (Stratagene) for phasing.

All proteins were expressed in *E. coli* strain C41. Following introduction of plasmids into cells by electroporation, cells were first grown overnight, at 37 °C, on three nutrient agar plates, containing 100  $\mu$ g/mL ampicillin. All cells from the agar plates were then transferred to 12 L of 2 $\times$  tryptone-yeast extract medium containing 100  $\mu$ g/mL ampicillin and grown at 37 °C. For *B. subtilis* SepF and SepF-G109K, protein expression was induced at OD<sub>600</sub> by the addition of 1 mM IPTG, at 25 °C for 6 h. In the case of the *A. fulgidus* and *P. furiosus* proteins, synthesis was induced with 1 mM IPTG, at 37 °C for 3–4 h. For purification, cells expressing *B. subtilis* SepF or SepF-G109K were resuspended in 25 mM Tris-HCl, pH 8.5, 1 mM EDTA, and 1 mM Na<sub>3</sub> and subsequently lysed using a cell disruptor (Constant Systems). The cell lysate was clarified by ultracentrifugation at 25.171  $\times g$ , which was followed by precipitation of the protein from the lysate supernatant by gradual addition of ammonium sulfate, to a final concentration of 25% (saturation), while incubating at 4 °C. The precipitated protein was pelleted at 17.590  $\times g$ ; and the pellet was resuspended in 25 mM Tris-HCl, pH 8.5, 1 mM EDTA, 1 mM Na<sub>3</sub>, and 100 mM NaCl; concentrated; and loaded onto a HiPrep Sephacryl S200 (GE Healthcare) gel filtration column. To obtain the truncated forms of *B. subtilis* SepF or SepF-G109K, before the gel filtration step, the protein concentrations were measured using a NanoDrop ND-1000 Spectrophotometer (Thermo), and the sample was incubated with  $\alpha$ -chymotrypsin in a 1:1,000 concentration ratio between protease and SepF protein. Limited proteolysis, at 37 °C for 1 h, was immediately followed by gel filtration on a Sephadex S200 column. Both SepF and SepF-G109K were concentrated to 10 mg/mL for crystallization trials.

For expression of selenomethionine-derivatized *A. fulgidus* and *P. furiosus* proteins, chemically competent *E. coli* B834 (DE3) cells (Novagen) were transformed with the relevant plasmids; grown overnight at 37 °C on agar plates containing 100  $\mu$ g/mL ampicillin; and then transferred to sterile M9 minimal media containing 2 mM magnesium sulfate, 0.4% (wt/vol) glucose, 25  $\mu$ g/mL ferrous sulfate heptahydrate, 1  $\mu$ g/mL each of a mixture of vitamins (riboflavin, niacinamide, pyridoxine monohydrochloride, and thiamine), 40  $\mu$ g/mL amino acid Complete Supplement Mixture (CSM, -Met), 40  $\mu$ g/mL of seleno-L-methionine (Fisher 259960025), and 100  $\mu$ g/mL ampicillin. Protein synthesis was induced with 1 mM IPTG, at 22 °C for 10 h. Pelleted cells were resuspended in 50 mM Hepes (pH 7.5) and 5 mM DTT and lysed using a cell disruptor, and lysate was clarified by ultracentrifugation at 25.171  $\times g$ . The lysate supernatant was loaded onto a 10-mL cation-

exchange column (Pharmacia), and protein was eluted with a linear gradient of 0–1 M NaCl. Relevant fractions were pooled, concentrated, and loaded onto a Sephacryl S100 gel filtration column preequilibrated in 25 mM Tris-HCl, pH 7.5, 1 mM EDTA, 1 mM Na<sub>3</sub>, and 5 mM DTT. For crystallization trials, *P. furiosus* protein was concentrated to 15 mg/mL and *A. fulgidus* protein to 10 mg/mL.

**Protein Crystallization.** Crystallization experiments were carried out using the sitting-drop vapor diffusion technique, with incubation at 19 °C. *P. furiosus* protein crystals grew in 0.2 M ammonium sulfate, 0.1 M Hepes, pH 7.5, and 25% (wt/vol) PEG 3350 and were cryo-protected in mother liquor containing 20% (vol/vol) ethylene glycol. *A. fulgidus* protein crystals appeared in 0.2 M lithium sulfate, 0.1 M sodium acetate, pH 4.5, and 30% (wt/vol) PEG 8000 and were cryo-protected with 25% (vol/vol) glycerol in mother liquor. Truncated *B. subtilis* SepF was crystallized in 30% (wt/vol) PEG 8000 and 0.2 M ammonium sulfate and for data collection crystals were cryo-protected with 20% (vol/vol) glycerol in mother liquor. Crystals of truncated *B. subtilis* SepF-G109K grew in 15% (vol/vol) ethanol and 0.1 M Tris, pH 7.0, and were cryo-protected with 20% (vol/vol) ethylene glycol in mother liquor.

**Structure Determination.** The crystal structures of the *A. fulgidus* and *P. furiosus* proteins (P6<sub>5</sub>22 crystal form) were solved by multiple-wavelength anomalous diffraction, using selenomethionine-derivatized crystals. Selenium sites were located and phases obtained with SOLVE/RESOLVE (40). Clear electron density maps revealed the presence of three dimers in the asymmetric unit of the *A. fulgidus* crystal and two monomers packed as a dimer in that of *P. furiosus*. The structures of *B. subtilis* wild-type SepF and the P2 crystal form of the *P. furiosus* protein were solved by molecular replacement, with Phaser 2.1 (41), using the P6<sub>5</sub>22 form of the *P. furiosus* protein as template. The search revealed one dimer in the asymmetric unit of the *B. subtilis* SepF crystal and six monomers packed as three dimers for the *P. furiosus* protein (P2). *B. subtilis* SepF-G109K was also solved by molecular replacement, using the truncated *B. subtilis* wild-type SepF monomer as template, which revealed one monomer in the asymmetric unit. Model building was done with Coot (42), and refinement was carried out using Refmac (43) and Phenix (44).

**Analytical Ultracentrifugation and Gel Filtration Assays.** Gel filtration assays were carried out by loading 1.5 mg of protein onto a Superdex S-200 gel filtration column (GE Healthcare), preequilibrated in buffer containing 25 mM Tris-HCl, pH 7.5, 1 mM EDTA, 1 mM Na<sub>3</sub>, and 50 mM NaCl, and eluting at a flow rate of 0.5 mL/min.

Sedimentation equilibrium experiments were carried out in a Beckman Optima XL-I analytical ultracentrifuge with an An-60 Ti rotor, using interference and absorbance at 280 nm. Samples were spun at 20 °C, 21,000 rpm, in two-sector 12-mm path-length cells, until they reached equilibrium, as judged by the lack of changes in subsequent scans. The three different concentrations used were 1 mg/mL, 2 mg/mL, and 4 mg/mL. The data were analyzed using Ultraspin (<http://www.mrc-lmb.cam.ac.uk/dbv/ultraspin2/ultraspin1/>). Partial-specific volumes were calculated from the amino acid compositions of the proteins, using SEDNTERP, which was also used for estimation of the density and viscosity of buffers (45).

**Electron Microscopy.** Negative staining electron microscopy images of truncated SepF were collected on a Philips 208 TEM operating at 80 kV and at a magnification of 180,000 $\times$ . Protein was diluted to a concentration of 0.05 mg/mL in 50 mM Tris-HCl, pH 7.0, and was applied to carbon-coated grids that had been made hydrophilic by glow discharging for 10 s. Grids were stained with 2% uranyl acetate. To image SepF with liposomes, the preformed liposomes were diluted to 5 mg/mL in 10 mM Tris-HCl, pH 7.4, 50 mM KCl, and 2 mM MgCl<sub>2</sub> and sonicated for 15 min in a bath sonicator. SepF (0.3 mg/mL) was incubated with the sonicated liposome suspension (25  $\mu$ L final volume) for 10 min at room temperature. In the case that FtsZ was included (0.1 mg/mL) the buffer also contained 4 mM GTP. After incubation, 20  $\mu$ L of sample was applied to glow-discharged 200-mesh carbon-coated grids, which were negatively stained with 100  $\mu$ L uranyl-acetate (2% wt/vol). The grids were imaged using a Phillips CM100 electron microscope.

For the measurement of septum widths, *B. subtilis* wild-type (strain 168) cells were grown in LB at 37 °C until the midexponential phase and pelleted. Fixation of cells was done in 2% glutaraldehyde, and then cells were washed with sodium cacodylate buffer, followed by secondary fixation in 1% osmium tetroxide. Fixed cells were washed with distilled water. After fixation, cells were dehydrated with gradual addition of acetone and embedded in resin at 60 °C for 24 h. Resin blocks were sectioned to 70-nm samples with a Leica EM UC7, using a diamond knife. Ultrathin sections were collected on

copper mesh grids and stained with uranyl acetate and lead citrate, using an automatic stainer (Leica EM AC20). Grids were examined using a Philips CM100 Compustage (FEI) transmission electron microscope.

**FtsZ Interaction Assay.** The interaction of SepF with FtsZ was analyzed by a coelution experiment as described before (4). In brief, MBP-SepF and MBP-SepF<sub>G109K</sub> were expressed in *E. coli*, and cell extracts were mixed with amylose resin and washed with 50 mM Tris-HCl, pH 7.4, 200 mM KCl, 1 mM EDTA, and 0.5 mM DTT. Resin was mixed with purified *B. subtilis* FtsZ at 4 °C for 2 h, followed by washing and elution with 10 mM Maltose and 1% SDS. Elution fractions were analyzed by SDS/PAGE and Coomassie staining.

**Fluorescence Microscopy.** Overnight cultures were diluted 1:50 in fresh LB and grown to exponential phase at 30 °C. The cell membrane was stained with FM-95 (1 μg/mL). SepF-GFP was induced with 0.5% xylose. N-terminal fragments of SepF fused to GFP were induced on nutrient agar plates supplemented with 1% xylose. Cells were immobilized on microscope slides covered with 1.2% agarose in water. Epifluorescence images were acquired with Nikon Ti-E and Zeiss 200M fluorescence microscopes, using Metamorph 6 (Molecular Devices). Two-dimensional SIM images were obtained with a Nikon N-SIM microscope and NIS-Elements 4.1 (Nikon).

**Protein Purification for Lipid Interactions.** SepF mutants were purified as Maltose binding protein (MBP) fusions on amylose resin columns, as described before (4). For purification, overnight cultures of *E. coli* BL21 (DE3) with the relevant pMal-SepF plasmids were diluted 1:50 into fresh LB containing 100 μg/mL ampicillin. Cells were grown until an OD<sub>600</sub> of 0.5 was reached, and protein expression was induced with 0.5 mM IPTG for 3 h. Cells were resuspended in 20 mM Tris-HCl, pH 7.5, 200 mM KCl, and 5 mM EDTA with complete EDTA-free protease inhibitor tablets (Roche), followed by lysis with a French press using 20 kpsi pressure. MBP-SepF was purified using an amylose column and eluted with 20 mM Tris-HCl, pH 7.5, and 10 mM Maltose. MBP-SepF fusion was cleaved with Factor Xa (1 μg enzyme for 100 μg MBP-fusion protein), and SepF was separated from MBP with a HiTrap Q HP column (GE Healthcare), followed by concentration with Millipore Amicon Ultra protein concentration tubes (3-kDa cutoff). Small aliquots were frozen in liquid N<sub>2</sub> and stored at -80 °C. *B. subtilis* FtsZ was purified as described previously (46, 47).

**Lipid Interaction Assays.** Lipid binding experiments were carried out with large unilamellar vesicles prepared from *E. coli* polar lipid extract or synthetic phospholipids (Avanti), essentially as described in ref. 48. Dry lipid extract was solubilized in chloroform followed by evaporation of chloroform under an argon stream. Dry lipids were resolubilized in 50 mM Tris-HCl, pH 7.5, 2 mM β-mercaptoethanol, and 1.5% octylglycoside (20 mg/mL lipids) under an argon stream and dialyzed against 50 mM Tris-HCl, pH 7.5, and 2 mM β-mercaptoethanol. The preformed liposomes were aliquoted and stored at -80 °C.

*B. subtilis* lipids were extracted using the Bligh/Dyer extraction method (49). In brief, 1 L *B. subtilis* midlog culture grown in LB at 37 °C was harvested, washed with 170 mM NaCl, and resuspended in 10 mL ice-cold PBS. Cells were broken with a French press and cell debris was removed by centrifugation. Organic extraction (chloroform, methanol, water 1:2:0.8) was carried out for 2 h at room temperature under argon, followed by addition of chloroform and water (1:1:0.9) to induce phase separation. Chloroform phase was collected after centrifugation and evaporated under a dry argon stream. Lipids were solubilized by addition of 50 mM Tris-HCl, pH 7.5, 2 mM β-mercaptoethanol, and 1.5% octylglycoside. Insoluble material was removed by filtering. Liposomes were formed as described for *E. coli* lipid extracts.

For pelleting experiments, the preformed liposomes (20 mg/mL, polar lipid extract) were extruded through a 0.4-μm pore size membrane (Avanti Polar Lipids) in SepF-binding buffer (20 mM Tris-HCl, pH 8.0, 2 mM MgCl<sub>2</sub>, 200 mM KCl, 0.5 mM DTT, 1 mg/mL BSA). Purified SepF (0.25–0.5 mg/mL) and liposomes (5 mg/mL) were mixed in a total volume of 120 μL and incubated for 10 min at room temperature. The mixture was centrifuged in 0.8-mL ultraclear centrifuge tubes at 40,000 rpm for 10 min at 20 °C, in a Beckman TLA100 Rotor. Pellets and supernatants were analyzed by SDS/PAGE and stained with Coomassie Brilliant Blue G.

For sucrose gradient density (flotation) experiments, preformed liposomes (1 mg/mL, polar lipid extract) were diluted in SepF-binding buffer containing a final concentration of 45% sucrose. After extrusion through a 0.1-μm pore membrane, the liposomes (80 μL) were mixed with SepF (0.25–0.5 mg/mL), resulting in a final volume of 120 μL and 30% sucrose. BSA (1 mg/mL) was included and the mixture was incubated at room temperature for 10 min. A sucrose gradient was prepared using SepF-binding buffer containing 1 mg/mL BSA and 10% (100 μL), 15% (100 μL), 20% (200 μL), or 25% (200 μL) sucrose, loaded on top of the SepF-liposome mixture. Centrifugation was carried out for 2 h (25,000 rpm, 25 °C) in a Beckman Rotor MLS50, using 0.8-mL tubes (5 × 41 mm) and suitable adapters, followed by collection of 100-μL samples from top to bottom. Samples were analyzed by SDS/PAGE and Coomassie Brilliant Blue G staining. Interaction of FtsZ (3 μg/mL) with SepF (0.25–0.5 mg/mL) and liposomes (1 mg/mL, 0.1 μm diameter) was tested with and without GTP (2 mM) in 10 mM Tris-HCl, pH 7.4, 50 mM KCl, 2 mM MgCl<sub>2</sub>, and 1 mg/mL BSA. The reaction was performed at room temperature for 10 min. Gradient samples were analyzed by Western blotting, using FtsZ primary antibody and goat anti-rabbit HRP (Sigma).

For the microscopic liposome binding assay, liposomes (5 mg/mL, polar lipid extract) were sonicated for 15 min in a bath sonicator, followed by incubation with SepF (0.25–0.5 mg/mL) in binding buffer for 10 min at room temperature. Liposomes were stained with 0.1 mg/mL Bodipy FL C16 (Invitrogen). A Nikon Ti equipped with a spinning-disk confocal module and a 488-nm solid-state laser light source was used to visualize the fluorescently labeled liposomes. Images were acquired with Metamorph 6 and analyzed with ImageJ 1.46 (National Institutes of Health).

CD spectroscopy was carried out with a Jasco J-810 spectropolarimeter at 25 °C, using a synthetic peptide corresponding to SepF1-13 (Biomatik), and liposomes (phosphatidylglycerol, phosphatidylethanolamine, cardiolipin = 60:38:2 mol%) formed from synthetic pure phospholipids as described above (Avanti). Before the experiment, the preformed liposomes were sized to 30 nm through extrusion and mixed with the synthetic peptide, resulting in final concentrations of 2.5 mg/mL lipids and 250 μg/mL peptide (21:1 molar ratio) in 6 mM Tris-HCl, pH 7.5. Spectra are smoothed background subtracted averages of nine scans acquired using a 20-nm/min scan rate, a 4-s response time, and a 0.2-mm path length cuvette (Helma).

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