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Creators	Garcia-Lara, Jorge, Weihs, Felix, Ma, Xing, Walker, Lucas, Chaudhuri, Roy R., Kasturiarachchi, Jagath, Crossley, Howard, Golestanian, Ramin and Foster, Simon J.

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Supramolecular structure in the membrane of *Staphylococcus aureus*

Jorge García-Lara^{a,1}, Felix Weihs^a, Xing Ma^{a,2}, Lucas Walker^a, Roy R. Chaudhuri^a, Jagath Kasturiarachchi^{a,3}, Howard Crossley^a, Ramin Golestanian^{b,4}, and Simon J. Foster^{a,4}

^aThe Krebs Institute, Department of Molecular Biology and Microbiology, University of Sheffield, Sheffield S10 2TN, United Kingdom; and ^bRudolf Peierls Centre for Theoretical Physics, University of Oxford, Oxford OX1 3NP, United Kingdom

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All life demands the temporal and spatial control of essential biological functions. In bacteria, the recent discovery of coordinating elements provides a framework to begin to explain cell growth and division. Here we present the discovery of a supramolecular structure in the membrane of the coccal bacterium *Staphylococcus aureus*, which leads to the formation of a large-scale pattern across the entire cell body; this has been unveiled by studying the distribution of essential proteins involved in lipid metabolism (PlsY and CdsA). The organization is found to require MreD, which determines morphology in rod-shaped cells. The distribution of protein complexes can be explained as a spontaneous pattern formation arising from the competition between the energy cost of bending that they impose on the membrane, their entropy of mixing, and the geometric constraints in the system. Our results provide evidence for the existence of a self-organized and nonpercolating molecular scaffold involving MreD as an organizer for optimal cell function and growth based on the intrinsic self-assembling properties of biological molecules.

membrane | model | staphylococcus | MreD | PlsY

The perpetuation of all cellular life requires the temporal and spatial management of essential biological functions, within the morphological framework characteristic of a specific organism. The underlying processes, which determine cell shape, are intimately intertwined with cell division and constitute pivotal issues for cell biology; their coordination in prokaryotes is mediated through counterparts of eukaryotic actin, tubulin, and intermediate filaments in addition to other specific components (1, 2). Several of these apparent cytoskeletal elements capitalize on their membrane binding properties, assembling along the longitudinal axis, between the poles of rod-shaped cells; they participate in many processes, including selection of the division site via the Min system and other components (3, 4); guidance and control of the cell wall biosynthetic machinery responsible for cell size, polarity, and shape through the actin-like protein MreB (5–11); and chromosome partitioning into daughter cells using another actin-like filament, ParM (12).

Despite this set of highly coordinated mechanisms, it has recently been shown that otherwise rod- and cocci-shaped bacteria can exist as largely spherical wall-less forms known as L-forms, with the capacity to divide (13). Importantly, the division of L-forms of the rod-shaped bacterium *Bacillus subtilis* is freed from the requirement of the classical tubulin-like division component FtsZ (14). L-forms appear to divide by scission after blebbing, tabulation, or vesiculation dependent on an altered rate of membrane biosynthesis (15); this harks back perhaps to a more evolutionary primitive mechanism permitting cellular proliferation. Thus, are there underlying organizational mechanisms that exist, independent of apparent cytoskeletal elements? The fluid mosaic model proposing the free diffusion of membrane proteins through the lipids has been challenged by growing evidence of the subcellular heterogeneity within the membrane resulting from diverse clustering of lipids and proteins (16–18).

Membrane curvature can act as a cue for localization of components (16). Raft aggregation of transmembrane proteins and the presence of compartment boundaries are insufficient explanations for such patterning. The physiological principles and molecular processes governing pattern formation are largely unknown.

Staphylococcus aureus is a coccal bacterium that can grow and divide in three consecutive orthogonal planes with fidelity (19); however, it lacks key morphogenetic components, such as MinCDE and MreB (20). Hence, what are the spatial organizers in *S. aureus* (21)?

Here we present the discovery of a supramolecular structure in the membrane of *S. aureus* that has been unveiled by studying the distribution of essential proteins involved in lipid metabolism (PlsY and CdsA) and the cell division component MreD. Such novel distribution of proteins complexes can be explained mainly as a by-product of the energy cost of bending that potential complexes exert on the membrane, and the geometric constraints imposed by the latter. A model based on such basic

Significance

The fundamental processes of life are organized and based on common basic principles. Molecular organizers, often interacting with the membrane, capitalize on cellular polarity to precisely orientate essential processes. The study of organisms lacking apparent polarity or known cellular organizers (e.g., the bacterium *Staphylococcus aureus*) may enable the elucidation of the primal organizational drive in biology. How does a cell choose from infinite locations in its membrane? We have discovered a structure in the *S. aureus* membrane that organizes processes indispensable for life and can arise spontaneously from the geometric constraints of protein complexes on membranes. Building on this finding, the most basic cellular positioning system to optimize biological processes, known molecular coordinators could introduce further levels of complexity.

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¹Present address: School of Medicine and Dentistry, College of Clinical and Biomedical Sciences, University of Central Lancashire, Preston PR1 2HE, United Kingdom.

²Present address: Unilever Future Leader Programme, Unilever R&D, Sharnbrook, Bedford MK44 1LQ, United Kingdom.

³Present address: Centre for Inflammation Research, College of Medicine and Veterinary Medicine, University of Edinburgh, Edinburgh EH16 4TJ, United Kingdom.

⁴To whom correspondence may be addressed. Email: s.foster@sheffield.ac.uk or ramin.golestanian@physics.ox.ac.uk.

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properties offers a new fundamental organizing framework in nondifferentiating coccal bacteria such as *S. aureus*, and may be one of the early cues that dictates the position of a target protein in the ensemble of proteins (22).

Results and Discussion

PlsY Is an Essential Protein with a Patterned Distribution in the Membrane. PlsY is an acyltransferase required for the indispensable process of phospholipid biosynthesis, and highly conserved across bacteria (23, 24). We show that PlsY is essential for the growth of *S. aureus*, demonstrated by the creation of a conditional lethal strain (*SI Appendix*, Fig. S1 *A* and *B*). PlsY-depleted cells exhibit misplaced cell division septa, altered cell wall morphology, and defective cytokinesis, suggesting a problem in cell division (Fig. 1 *A–D* and *SI Appendix*, Fig. S1*F*). To examine the link between PlsY and cell cycle progression, we studied the distribution of divisome proteins EzrA and PBP2. EzrA forms a septal ring at the midplane of the cell and was found at unexpected division planes when cells are depleted for PlsY (Fig. 1 *E–G* and *Movie S1*) (25, 26). PBP2, a septal protein responsible for cell wall biosynthesis at the site of division (27), was also delocalized (*SI Appendix*, Fig. S2), suggesting PlsY involvement in septum placement and/or division progression.

A chromosomally encoded and functional PlsY–GFP fusion under the control of the *plsY* promoter demonstrated septal localization, but also distributed in a pattern of foci around the membrane throughout the cell cycle (Fig. 2 *A–C*, *SI Appendix*, Figs. S3–S5, and *Movie S2*). Immunoblot and FACS analysis demonstrated that PlsY is membrane associated (*SI Appendix*, Fig. S6), and *plsY* repression led to a decrease of PlsY (*SI Appendix*, Fig. S1 *C–E*).

The fluorescent translational fusion reporter data were supported by immunofluorescence microscopy observations of PlsY distribution, which also revealed a pattern (Fig. 2*D* and *SI Appendix*, Fig. S7*A*), and optical sectioning of fluorescent or immunolabeled PlsY that showed a 3D distribution of foci (*SI Appendix*, Fig. S3 and *Movie S3*). The pattern was present in both fixed and unfixed samples, demonstrating that the observed pattern is not an artifact of the sample preparation procedure (*SI Appendix*, Fig. S8*A*). CdsA, a CDP-diacylglycerol synthase

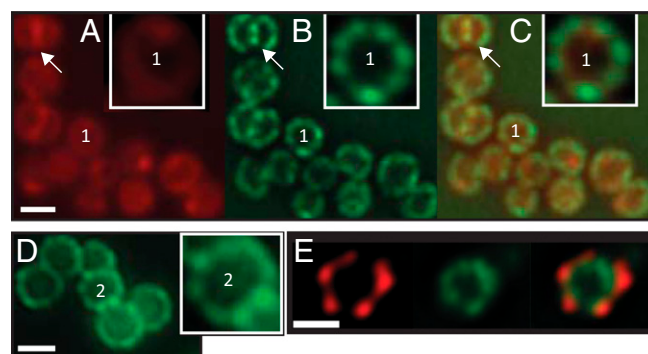


Fig. 2. (*A–C*) PlsY localizes to the cell division site (see arrow) but also distributes in a pattern in the membrane of *S. aureus* whether detected by GFP labeling [*B* and *C*, counterstained with Nile red (*A*) or immunolabeling with α -731 antibody (*D*). (*E*) CdsA (GFP labeled, green) has a similar distribution pattern to PlsY and colocalizes with immunolabeled PlsY (red). (Scale bars, 1 μ m.)

downstream of PlsY in the glycerophospholipid biosynthesis pathway, localized in a similar pattern and colocalized with PlsY (Fig. 2*E*, *SI Appendix*, Figs. S7*C* and S8*B*, and *Movies S4–S6*), whereas other membrane proteins did not (e.g., YmdA and SecY; *SI Appendix*, Fig. S7*B*). Consistent with those observations, PlsY and CdsA also showed a weak interaction in a bacterial two-hybrid analysis (BACTH; *SI Appendix*, Fig. S9). To confirm interaction between both components at the molecular level in their natural environment, we implemented a FRET-based system in *S. aureus* using fluorescent protein fusions. A donor bleaching strategy was applied and revealed a specific interaction between PlsY and CdsA (*SI Appendix*, Fig. S10). To our knowledge, these are the first pieces of evidence of the suspected colocalization and interaction between the enzymes in this pathway, and support the notion of metabolic channelling. Lipid clusters are also organized with PlsY in the membrane of unfixed cells (*SI Appendix*, Fig. S11). Such a distribution could correspond to the differential accumulation of specific fatty acid species or lipid conformations in the membrane (28).

MreD Is Needed for the Formation of a Supramolecular Structure in the Membrane. PlsY and/or CdsA seemed unlikely candidates as organizers of the patterned distribution, as did FtsZ, despite its link to the localization of phospholipid synthases (29). The MreBCD complex in rod-shaped bacteria localizes along the membrane, acting as a spatial organizer of various processes (7, 9, 30–33). In *Escherichia coli*, MreB (an actin-like molecule) is required to regulate phospholipid and membrane biosynthesis (31). However, MreB is missing in *S. aureus* and other coccal cells, whereas MreC and MreD are present (9).

S. aureus deprived of MreC grew identically to the parent, whereas lack of MreD or MreCD led to a growth defect with larger cells of abnormal morphology (Fig. 3 *A–C* and *SI Appendix*, Figs. S12 and S13), reminiscent of the PlsY-depleted phenotype. The lack of an apparent phenotype for the single *mreC* mutation was interesting given its importance in other organisms (34). Genome sequencing of two individual *mreC* mutants revealed one single nucleotide polymorphism in each strain leading to an amino acid substitution (*SI Appendix*, *Whole Genome Sequencing*). However, those substitutions seem unlikely candidates to justify the lack of an *mreC* phenotype. Loss of MreD can be complemented by ectopic expression of the *mreD* gene (*SI Appendix*, Fig. S13 *F* and *G*). MreD is required for the observed pattern of PlsY (cf. Fig. 2 *B* and *D* with Fig. 3*D* and *SI Appendix*, Fig. S14*A*) and the characteristic placement of EzrA (cf. Fig. 1*G* with Fig. 3*E*, *SI Appendix*, Fig. S14*B*, and *Movie S7*); this is consistent with the immunodetection of a MreD–GFP fusion that revealed a

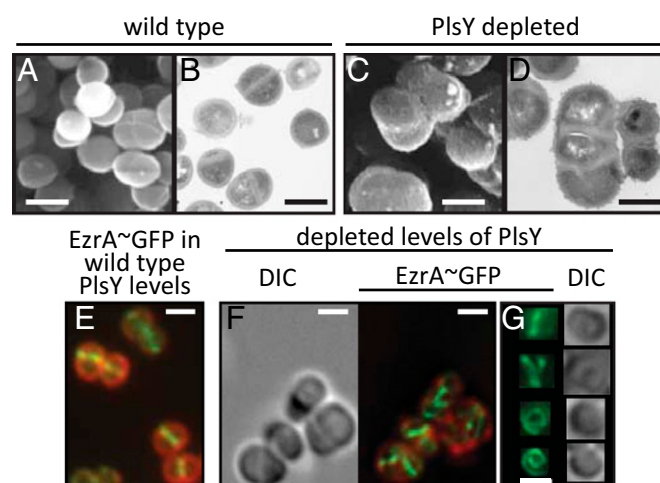


Fig. 1. Scanning (*A* and *C*) and transmission (*B* and *D*) electron micrographs of PlsY-depleted *S. aureus* JGL166 (*C* and *D*) reveal aberrant cellular morphologies compared with wild-type cells (*A* and *B*). (*E–G*) Fluorescence microscopy of the cell division-related EzrA protein tagged with GFP (green) reveals its septal location in wild-type cells (*E*); cell membrane is stained with Nile red (red). In contrast, EzrA–GFP is delocalized in the absence of PlsY (*F* and *G*). DIC, differential interference contrast microscopy. (Scale bars, 1 μ m.)

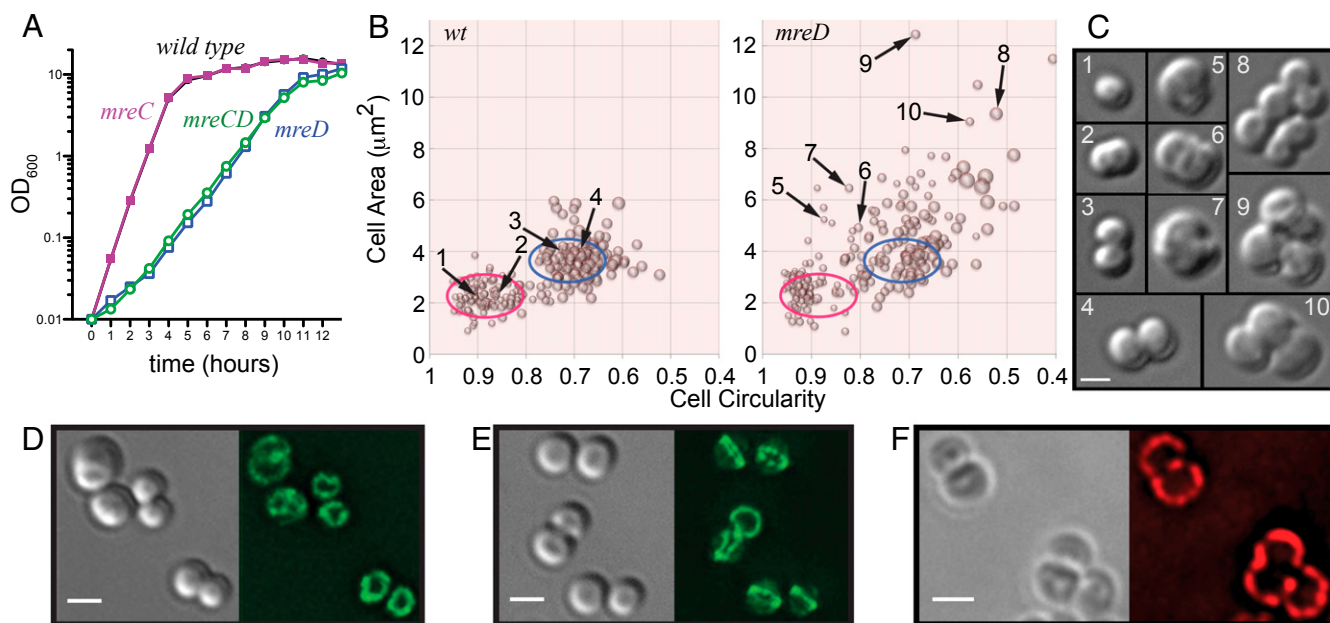


Fig. 3. Role of MreD in cell growth and morphology. (A) Growth of *S. aureus* SH1000 wild type (●, black), *mreC* deletion mutant (strain SJF4098; ■, pink), *mreD* deletion mutant (strain SJF2976; □, blue), and *mreCD* deletion mutant (strain SJF2625; ○, green). (B) Cell size distribution in SH1000 ($n = 235$) and *mreD* mutant (strain SJF2976; $n = 262$) populations of *S. aureus* at OD₆₀₀ = 0.5–0.8. Pink ellipse encloses single cells; blue ellipse encloses normally dividing cells. Cell sizes >4.5 μm² correspond to single, dividing, or clusters of cells of abnormally large size. Two shape descriptors have been used to illustrate cell morphology: cell circularity (x axis of the graphs) and cell roundness (illustrated by the diameter of the symbols). (C) Examples of the cell categories plotted in Fig. 3B. MreD depletion in *S. aureus* leads to the delocalization of GFP-tagged PlsY (D) and EzrA (E) even in cells with wild-type morphology. (F) Immunolocalization of MreD-GFP with anti-GFP antibodies reveals a distribution of MreD in the *S. aureus* membrane similar to PlsY. (Scale bars, 1 μm.)

localization pattern equivalent to PlsY and CdsA (Figs. 2B and D and 3F and SI Appendix, Figs. S3 and S15). BACTH analysis also showed protein–protein interactions between MreD and CdsA (SI Appendix, Fig. S9). Similarly, FRET provided supporting evidence of the interaction between MreD and PlsY (SI Appendix, Fig. S10).

Spontaneous Formation of a Patterned Distribution of Protein Complexes in the Membrane. Without cytoskeletal components, how might MreD, PlsY, and CdsA adopt a pattern in the membrane? Their distribution could arise from the interplay between the potential complex they contribute to and the geometric constraints imposed on the membrane, doing what cellular organizers do, but spontaneously, and leading the way. If the integral membrane protein complex inflicts a sufficiently large local curvature on the membrane, a random distribution would be frustrated due to the high bending energy cost that results from accommodating the complexes across the membrane. This mechanism of protein organization can account for pattern formation of FtsZ in liposomes (35, 36), and is also consistent with the sensing of 2D membrane geometry by the Min system (37).

The apparently spherical membrane of *S. aureus* and any surface deformation can be expressed in spherical coordinates as $\mathbf{R}(\theta, \varphi) = R[1 + u(\theta, \varphi)]\hat{\mathbf{r}}$, where R is the radius of the unperturbed sphere, $u(\theta, \varphi)$ describes the surface deformations, and $\hat{\mathbf{r}}$ is the radial unit vector. The distribution of the protein complexes across the membrane is described with the surface number density $\rho(\theta, \varphi) = \rho_0[1 + \psi(\theta, \varphi)]$. We assume that each protein complex covers a patch of area a_0 and interacts with the membrane geometry by imposing a spontaneous curvature H_p at its location. A free energy functional for the system can be written as the sum of three contributions $\mathcal{F} = \mathcal{F}_b + \mathcal{F}_d + \mathcal{F}_t$; this comprises a bending energy contribution, $\mathcal{F}_b = \kappa/2 \int dA [H^2 - 2H_p \rho a_0 H]$, due to membrane deformation, a contribution due to density

fluctuations of the diffusing protein complexes of the form $\mathcal{F}_d = 1/2\chi \int dA [\xi^2 (\nabla \psi)^2 + \psi^2]$ that originates from their entropy of mixing, and a harmonic potential $\mathcal{F}_t = kR^2/2 \int dA u^2$ that results from the tethering of the membrane to the vicinity of the cell wall and the turgor pressure. Here, κ is the bending rigidity, $H/2$ is the mean curvature, χ is the compressibility and ξ is the correlation length of the density fluctuations, and k is an effective spring constant (per unit area) representing the confinement potential. Using this free energy, we can define the governing dynamical equations for $u(\theta, \varphi)$ and $\psi(\theta, \varphi)$, which involve the corresponding mobility (transport) coefficients L_u and L_ψ (SI Appendix, Fig. S16 and Fig. 4A). Keeping the leading order in deformation and density fluctuations, and using expansion in spherical harmonics $Y_{lm}(\theta, \varphi)$ for both fields, a linear stability analysis can be performed to find the growth rates of the characteristic modes of the system. It is convenient to group the many parameters in the system into the dimensionless variables $K = kR^4/\kappa$, $M = L_\psi/(\kappa L_u \chi)$, $P = \xi^2/R^2$, and $W = \kappa \chi \rho_0^2 a_0^2 H_p^2$.

Fig. 4B shows the relevant growth rate λ_+ in units of $\kappa L_u/R^2$ as a function of the mode number ℓ , at selected values of K , M , and P for $W = 1$. It shows that λ_+ acquires a positive value (Fig. 4C) for the $\ell = 8$ mode, which means that this mode becomes unstable. The instability in the intermediate $\ell = 8$ mode develops as the parameter W is changed from 0.8 to 1 (Fig. 4D), leading to the development of slowly growing patterns that are linear combinations of the $Y_{8m}(\theta, \varphi)$ for different values of m , which could lead to eight foci in a cross-section of the sphere (Fig. 4E). This example demonstrates that patterns such as those observed (Figs. 2B–E, 3F and G, and 4C and D, SI Appendix, Figs. S7A and C, S8A and B, and S11B, and Movies S3–S6) can be generated as a result of such intermediate wavenumber instability.

Assuming that the protein complexes are in the dilute regime—namely, $\rho a_0 \ll 1$, we can estimate the compressibility as $\chi \approx 1/(k_B T \rho_0)$ (38), and write the coupling parameter as

In eukaryotes, membrane partitioning is a well-described process with features at different length scales (41). In prokaryotes, it would also be envisaged that multiple mechanisms are at play, resulting in the coordination of processes from the molecular to the cellular level. The ability to colocalize members of the same macromolecular biosynthesis pathway provides a dynamic mechanism to permit efficient metabolism. Phospholipid biosynthesis involves complex intermediates and potential substrate channelling between enzymes, which would require their colocalization. The development of FRET in *S. aureus* has provided the experimental framework to test juxtaposition of proteins in their native membrane setting, which has shown the interaction between PlsY and CdsA. It will be of great interest to determine the localization pattern of other phospholipid biosynthesis enzymes in the pathway and to extend this to further proteins involved in other metabolic processes and beyond. It may be that such protein assemblies generate localized physiobiochemical environments propagating effects to other membrane components in the same or different processes. The observation of apparent differential partitioning of PlsY and a lipid dye in the membrane supports such a hypothesis (*SI Appendix*, Fig. S11).

It has previously been shown that MreB, in rod-shaped cells, acts as a guidance mechanism for multiple biosynthetic components (11). However, the roles of MreC and MreD have remained more obscure. Here we find an important role for MreD in growth in an organism lacking MreB. Deletion of *mreD* results in pleiotropic effects, and we suggest that in rods, MreB may spatially constrain MreD, but it is MreD that is itself a key player in the actual coalescence of molecules. In *S. aureus* we also see evidence of cellular morphology effects on MreD-associated processes during the cell cycle. Thus, other spatiotemporal cues will be brought into play to allow coordination during growth and division. We have hypothesized that cell wall peptidoglycan architectural features can provide a template for division site recognition (42). It is likely a combination of intrinsic properties of proteins, such as MreD, that may provide localized perturbation to the membrane, coupled with larger scale morphological features derived from other cellular components and morphological dynamics generated from apparent cytoskeletal elements (such as FtsZ) that determine

the overall mapping of membrane components. Such a dynamic accumulation of cues optimizes cellular reactions within a given environment.

In a drive to understand the basis of life, a key goal is the ability to assemble a protocell capable of self-assembly (43). Unraveling the mechanisms underpinning such assembly is crucial. Our observations provide a novel and basic organizational drive of proteins in membranes, and catalyze further research to determine their behavior in vitro and in vivo to understand patterning in a lipid environment and in the interplay within and between cellular processes. The mathematical model presented here gives a theoretical framework to be tested and to begin to predict the behavior of components underpinning life.

Materials and Methods

This section briefly recapitulates the data analysis methods. A detailed description of methods can be found in *SI Appendix*. IPTG-inducible expression constructs of *plsY* in single copy in the staphylococcal chromosome, as well as *mreC* and *mreD* deletion strains, were made through allelic replacement by double crossover recombination. The process involved the use of episomal vectors with temperature-sensitive replicons, initially constructed in *E. coli* (TOP10), followed by passaging of the resulting constructs through a restriction-deficient strain (*S. aureus* RN4220) and subsequent transduction-mediated integration into the final test strain (*S. aureus* SH1000). The *mreD* complementation plasmid was based on *S. aureus* replication-stable episomes. A similar procedure was followed to generate GFP-transcriptional fusions but mediated through single crossover recombination that resulted in an intact copy of the gene of interest under the control of the regulatable promoter Pspac, and another copy of the gene fused to GFP under the control of the native promoter of the gene of interest. The rest of the techniques, including protein and antibody production, immunolabeling, epifluorescence and electron microscopy preparation, flow cytometry analysis, bacterial two-hybrid analysis, FRET, and cell fractionation, were performed according to standard techniques optimized to our project. Imaging was undertaken with a DeltaVision RT Deconvolution microscope (Applied Precision) linked to an Olympus IX70 microscopy system (Olympus U-RFL-T and IX-HLSH100 lamps, and Olympus UPlanApo 100×/1.35 oil iris lens), with SoftWoRx 3.5.1 software. Cell enumeration and perimeter, major axis, and minor axis measurements were performed using ImageJ 1.46o.

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