Chromosome Segregation and Peptidoglycan Remodeling Are Coordinated at a Highly Stabilized Septal Pore to Maintain Bacterial Spore Development

Graphical Abstract



Highlights

- Coordination of cell wall remodeling and chromosome segregation at a septal pore
- Balance between cell wall hydrolysis and synthesis is required for pore stability
- Pore stability is reinforced by the highly conserved SpollIAH-SpolIQ interaction
- Coordinating chromosome segregation and cytokinesis is fundamental for development

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In Brief

Coordinating chromosome segregation with cytokinesis is of primordial importance during development. In this issue, Mohamed et al. define how coordination between chromosome segregation and cell wall remodeling contributes to critical aspects of genetic and transcriptional compartmentalization during endospore formation in bacteria, one of the earliest forms of cellular development on Earth.







Article

Chromosome Segregation and Peptidoglycan Remodeling Are Coordinated at a Highly Stabilized Septal Pore to Maintain Bacterial Spore Development

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SUMMARY

Asymmetric division, a hallmark of endospore development, generates two cells, a larger mother cell and a smaller forespore. Approximately 75% of the forespore chromosome must be translocated across the division septum into the forespore by the DNA translocase SpoIIIE. Asymmetric division also triggers cell-specific transcription, which initiates septal peptidoglycan remodeling involving synthetic and hydrolytic enzymes. How these processes are coordinated has remained a mystery. Using *Bacillus subtilis*, we identified factors that revealed the link between chromosome translocation and peptidoglycan remodeling. In cells lacking these factors, the asymmetric septum retracts, resulting in forespore cytoplasmic leakage and loss of DNA translocation. Importantly, these phenotypes depend on septal peptidoglycan hydrolysis. Our data support a model in which SpoIIIE is anchored at the edge of a septal pore, stabilized by newly synthesized peptidoglycan and protein-protein interactions across the septum. Together, these factors ensure coordination between chromosome translocation and septal peptidoglycan remodeling to maintain spore development.

INTRODUCTION

The establishment and maintenance of genetic and transcriptional compartmentalization during cytokinesis is a key aspect of development in all organisms. In eukaryotic cells, multiple lines of evidence suggest that proteins localized at the division site coordinate chromosome segregation with cytokinesis (Fraschini, 2020). This coordination requires interplay between the reorganization of the actin and microtubule cytoskeletal elements, a variety of motor proteins and membrane trafficking (Fraschini, 2020; Seiler and Justa-Schuch, 2010). In eukaryotic cells with a cell wall (e.g., yeast), cytokinesis also requires chitin synthases and hydrolytic enzymes such as chitinases and glucanases (Lesage and Bussey, 2006). Importantly, the molecular mechanisms governing coordination between chromosome segregation and cytokinesis remain poorly understood, even in the simplest organisms, such as bacteria. Here, using the bacterium Bacillus subtilis as a model system, we investigate the mechanisms underlying the coordination between chromosome

translocation and the last steps in cytokinesis during sporulation, a primordial example of cellular development.

In dividing bacteria, genetic compartmentalization is achieved by segregating most of the chromosome into each daughter cell prior to cytokinesis. Here, a membrane-anchored DNA translocase of the broadly conserved FtsK/SpoIIIE family aids in the completion of chromosome segregation (Sherratt et al., 2010; Grainge, 2010; Barre, 2007). During endospore formation in Bacillus subtilis, a more complex situation occurs: asymmetric division generates two cellular compartments of different size and distinct developmental fates: a smaller cell called the forespore, which develops into the dormant spore, and a larger cell designated the mother cell (Stragier and Losick, 1996) (Figure 1A). Importantly, asymmetric division precedes chromosome segregation and traps approximately 25% of the chromosome in the forespore, while the remaining 75% is translocated into the developing forespore by SpoIIIE (Wu and Errington, 1998; Sullivan et al., 2009) (Figure 1A). Remarkably, this occurs concomitantly with remodeling of the division septum and the early steps



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Figure 1. Identification of SpollIM and Its Requirement, Together with PbpG, in Maintaining Compartmentalization (A) Diagram of chromosome translocation and the different stages of engulfment, showing membranes (black), PG (gray), chromosome (black squiggles), SpollIE (green), origin (oriC, blue), and terminus (ter, red). The forespore cytoplasm is shown in beige.



of spore envelope formation (Piggot and Hilbert, 2004). How these events are coordinated in a way that maintains transcriptional compartmentalization during development has remained unclear.

SpollIE is composed of three domains: an N-terminal transmembrane domain, a poorly conserved linker, and a motor domain (Burton and Dubnau, 2010). The N-terminal transmembrane domain is required for SpollIE stability and localization to the middle of the asymmetric septum (Wu and Errington, 1997; Sharp and Pogliano, 1999, 2002). The SpollIE motor domain contains three subdomains (α , β , and γ), being classified as a member of the RecA family of ATPases and sharing ~78% sequence similarity to FtsK (Barre, 2007). $\alpha\beta$ assembles into a hexameric ring containing the ATPase machinery and a central channel through which double-stranded DNA is threaded (Cattoni et al., 2014). The SpolIIE γ domain translocates DNA directionally by recognizing chromosomal DNA sequence motifs called SpolIIE recognition sequences (SRSs) (Ptacin et al., 2008; Cattoni et al., 2013; Besprozvannaya et al., 2013).

Two proposed models explain how SpollIE translocates DNA across the septum (Figures 1B and 1C). In the aqueous pore model, SpoIIIE is anchored at the edges of a toroidal aqueous pore surrounded by an unfused septal membrane through which the chromosome is translocated (Wu and Errington, 1997; Fiche et al., 2013). In this model, the SpoIIIE motor domains are predominantly localized on the mother cell side of the pore, and the transmembrane domain is predicted to stabilize and coordinate septal pore closure with completion of chromosome translocation (Fiche et al., 2013). In the membrane channel model, SpollIE exists on both sides of a fused septal membrane, and its transmembrane domain forms intercellular channels through which the DNA is translocated (Burton et al., 2007; Yen Shin et al., 2015; Fleming et al., 2010). A lack of clear genetic evidence in support of either model has kept the debate surrounding these models open.

Concurrent with chromosome translocation into the forespore, compartment-specific sigma factors are activated to control gene expression in the mother cell and forespore. Upon asymmetric division, σ^{F} is activated in the forespore (Figure 1A), which controls a signal transduction pathway that triggers the activation of σ^{E} in the mother cell (Piggot and Hilbert, 2004). σ^{E} directs the synthesis of cell wall peptidoglycan (henceforth abbreviated as PG) hydrolases that assemble into the DMP complex

(composed of SpoIID, SpoIIM, and SpoIIP), which thin the septal PG and mediate engulfment of the forespore (Morlot et al., 2010; Chastanet and Losick, 2007) (Figure 1D). In addition to PG degradation, engulfment is thought to involve PG synthesis by biosynthetic complexes, although these are yet to be defined (Ojkic et al., 2016). Upon completion of engulfment, membrane fission releases the forespore into the mother cell cytoplasm as a protoplast surrounded by two membranes and a thin layer of PG in between (Khanna et al., 2019; Tocheva et al., 2013; Doan et al., 2013). Finally, during engulfment a highly conserved SpoIIIAH-SpoIIQ zipper-like interaction holds the mother cell and forespore membranes together and functions like a ratchet, promoting efficient, forward movement of the engulfing membranes (Broder and Pogliano, 2006; Ojkic et al., 2014) (Figure 1D).

Here, we reveal that chromosome translocation and septal PG remodeling during engulfment are coordinated through an aqueous pore stabilized by the SpoIIIE protein itself, a protein called SpoIIIM (formerly YqfZ), septal PG synthesis by the fore-spore-specific PG synthase PbpG and by the SpoIIIAH-SpoIIQ interaction across the septal membrane (Figure 7A). In the absence of these factors, the asymmetric septum retracts due to septal PG hydrolysis by the DMP complex, resulting in loss of cytoplasmic and chromosomal compartmentalization and a block to spore development (Figure 7C). Collectively, our work defines the coordination between septal PG remodeling and chromosome segregation that ensures genetic and cytoplasmic compartmentalization during spore morphogenesis and demonstrates the primordial importance of coordinating chromosome segregation with cytokinesis during development.

RESULTS

A Screen for Synthetic Lethal Partners with *pbpG* Identifies *spollIM*

The forespore produces two functionally redundant class A penicillin-binding proteins (PBPs), PbpG and PbpF, that generate a thin layer of spore PG called the germ cell wall (Figure 1D) (McPherson et al., 2001). Sporulating cells lacking one of these proteins produce close to wild-type (WT) levels of spores; however, cells lacking both are severely impaired in sporulation (McPherson et al., 2001). WT and the *pbpF* mutant produce forespores with an oblong shape. However, sporulating cells lacking

⁽B) Diagram of aqueous pore model (zoomed-in from red box in Ai), showing SpoIIIE on the mother cell (mc) side and not on the forespore (fs) side, localized within unfused septal membranes that form a pore in the membrane.

⁽C) Diagram of channel model (zoomed-in from red box in Ai), showing SpoIIIE on both sides of the fused septal membranes.

⁽D) Diagram of the leading edge of the engulfing membranes (zoomed-in from blue box in Aii), showing the SpolIIAH-SpolIQ interaction across the engulfing membranes, PbpG (light teal) and PbpF (dark teal) in the forespore membrane, the DMP complex (purple), new PG (green dots and lines), and PG in the lateral wall of the sporangium (gray dots and lines).

⁽E) Tn-seq profiles in WT and $\Delta pbpG$ after 24 h of growth and sporulation in exhaustion medium. The height of each line reflects the number of sequencing reads at this position. Red box highlights the *spollIM* (*yqfZ*) locus, which is depleted for transposon insertions in the $\Delta pbpG$ library compared with WT.

⁽F) Average sporulation efficiency (% \pm SD, n = 3) of Δ spolIIM, Δ pbpG, Δ spolIIM Δ pbpG, and the respective spolIIM complementation strains, Δ spolIIM and Δ pbpG Δ spolIIM in Difco sporulation medium (DSM).

⁽G) Average frequency (\pm SD of 3 biological replicates) of miscompartmentalized cells during a sporulation time course in WT, Δ spolIIM, Δ pbpG, and Δ spolIIM Δ pbpG (n > 900 per time course, per strain, per replicate).

⁽H) Representative images of miscompartmentalization in WT, $\Delta spoIIIM$, $\Delta pbpG$, $\Delta spoIIIM$ $\Delta pbpG$ and $\Delta spoIIIE$ at T3.5 (related to Figure 1G). Scale bar, 2 μ M. (I) Diagram showing the engulfment block occurring in the $\Delta spoIID$ $\Delta spoIIP$ and its effect on compartmentalization by preventing PG hydrolysis.

⁽J) Representative images (from one biological replicate, out of 3 examined) of miscompartmentalization in Δ spolID Δ spolID Δ spolIP at T3.5, in WT, Δ spolIIM, Δ pbpG, Δ spolIIM Δ pbpG and Δ spolIIE backgrounds. Scale bar, 2 μ M.

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pbpG exhibit an abnormal, jellybean-like morphology (Figure S1A) (Rodrigues et al., 2016). Based on this morphological difference, we hypothesized that PbpG and PbpF may have specialized roles and function in separate genetic pathways. To investigate this hypothesis and identify factors that function in each pathway, we used transposon sequencing (Tn-seq) to screen for genes that become critical for sporulation in cells lacking either *pbpG* or *pbpF* (Meeske et al., 2016).

In validation of our genetic screen, *pbpF* was one of the top hits in the $\Delta pbpG$ mutant transposon library and *pbpG* was one of the top hits in the $\Delta pbpF$ transposon library (Figures S1D and S1E). One of the strongest hits that was specific to the $\Delta pbpG$ library was a gene called *yqfZ* that we have renamed *spolIIM* (Figures 1E and S1B). Previous work indicates that *spolIIM* is transcribed in the mother cell compartment under σ^{E} control and encodes a 99-amino-acid protein of unknown function (Eichenberger et al., 2003). SpolIIM is predicted to have an N-terminal transmembrane segment and a C-terminal LysM domain (UniProt Consortium, 2018) that are implicated in binding to PG (Buist et al., 2008).

To verify that SpoIIIM becomes important for sporulation in $\Delta pbpG$ cells, we determined the sporulation efficiency of $\Delta spoIIIM$ and $\Delta spoIIIM \Delta pbpG$ mutants (Figure 1F). Consistent with previous reports, the $\Delta spoIIIM$ mutant had a mild sporulation defect (76.6% spores) and the $\Delta pbpG$ mutant sporulated to near WT levels (87.3% spores). In validation of the Tn-seq screen, the sporulation efficiency of the $\Delta spoIIIM \Delta pbpG$ double mutant was 8.8% (Figure 1F). Expression of spoIIIM from an ectopic chromosomal locus restored sporulation efficiency of $\Delta spoIIIM$ and $\Delta spoIIIM \Delta pbpG$ mutants to WT and $\Delta pbpG$ levels, respectively (Figure 1F). Finally, consistent with the results of the Tn-seq screen and in support of the idea that SpoIIIM is specifically required in cells lacking PbpG, no synergy was observed in the $\Delta spoIIIM \Delta pbpF$ double mutant (Figure S1C).

PbpG and SpollIM Are Required for Forespore Morphology and Compartmentalization

To begin to unravel how SpollIM and PbpG contribute to sporulation, we used fluorescence microscopy to determine if cells lacking these proteins exhibit morphological defects during development. Sporulating cells were imaged every 30 min after the onset of asymmetric division (1 h and 30 min after the initiation of sporulation; T1.5). No obvious differences in forespore morphology were observed between the WT and spollIM mutant at early stages of development; however, as the cells neared the completion of engulfment (T3) the spollIM mutant exhibited forespore morphologies that we categorized into four types: (1) WTlooking, (2) dwarf (small spherical forespores), (3) mislocalized (dwarf forespores positioned closer to the mid-cell of the mother cell), and (4) abnormal (forespores that lacked a spherical appearance and contained irregular membranes) (Figures S2B and S2E). Consistent with the Tn-seq and sporulation efficiency data, the $\Delta spollIM \Delta pbpG$ double mutant produced even more dwarf, mislocalized, and abnormal forespores (Figure S2E). Transmission electron microscopy (TEM) of *AspollIM* and $\Delta spolliM \Delta pbpG$ confirmed the presence of dwarf forespores with wrinkled membranes (Figure S2F). Interestingly, closer inspection of $\Delta pbpG$ cells revealed that 2.5% of the forespores have a dwarf phenotype (Figures S2B and S2E), suggesting that the absence of SpoIIIM enhances this defect.

The severity of the morphological defects in the $\Delta spollIM$ and $\Delta spollIM \Delta pbpG$ mutants led us to wonder if their forespores retain cytoplasmic compartmentalization. We engineered strains to express a cytoplasmic, cyan fluorescent protein (CFP) in the forespore under the control of a $\sigma^{\text{F}}\text{-dependent}$ promoter (P_{spollQ}), and examined the distribution of the CFP signal by fluorescence microscopy during a sporulation time course (Figure 1H). As expected, in WT cells the CFP signal was confined to the forespore throughout the entire time course (Figure 1H). In the Δ spollIM mutant we observed some cells with CFP signal confined solely to the forespore, some with stronger CFP signal in the forespore and weaker CFP signal in the mother cell and others with strong CFP signal throughout the entire sporangium (Figure 1H). The frequency of $\Delta spollIM$ sporulating cells with any CFP signal (weak or strong) in the mother cell increased over time and reached 33% at T3.5 (Figure 1G). These observations suggest that *AspolIIM* forespores leak their cytoplasmic contents into the mother cell and lose compartmentalization. We observed an even higher proportion of miscompartmentalized sporulating cells in the $\Delta spollIM \Delta pbpG$ double mutant (90%) at T3.5) (Figure 1G). Expression of spollIM in trans restored compartmentalization in the double mutant (Figure S2A). Miscompartmentalization was also observed in the $\Delta pbpG$ mutant, albeit at a low frequency (3%) (Figures 1G and 1H). Highlighting the specific genetic relationship between spollIM and pbpG, the degree of miscompartmentalization in the *AspoIIIM ApbpF* double mutant was comparable to the ΔspollIM mutant (Figures S2C and S2D). These data suggest that SpollIM and PbpG are also required for forespore compartmentalization.

Interestingly, miscompartmentalization and abnormal forespore morphology have both been reported for sporulating cells lacking spollIE (Liu et al., 2006; Hilbert et al., 2004; Doan et al., 2013). A side-by-side comparison of the $\Delta spoIIIM \Delta pbpG$ double mutant and $\Delta spoll E$ revealed similar forespore morphologies and degrees of miscompartmentalization (Figures 1H and S2B). Furthermore, combining $\Delta spoIIIM$ and/or $\Delta pbpG$ with a hypomorphic allele of spolIIE (SpolIIE D584A) (Burton et al., 2007), which exhibits miscompartmentalization in a small fraction of cells, resulted in a dramatic increase in miscompartmentalization (Figures S3A and S3B). These observations led us to investigate if the $\Delta spoIIIM$ and $\Delta pbpG$ mutant have other phenotypes in common with ∆spollIE. Previous work demonstrated that *∆spollIE* miscompartmentalization can be suppressed when cells lack the engulfment hydrolases, SpoIID and SpoIIP (Hilbert et al., 2004). Sporulating cells lacking both SpolID and SpolIP have flat septa due to a block in septal PG hydrolysis (Figure 1I) (Chastanet and Losick, 2007). In a AspoIID AspoIIP mutant, AspoIIIE miscompartmentalization was almost completely abolished (Figure 1J). Interestingly, this was also the case in Δ spollIM and Δ spolIIM Δ pbpG mutants (Figure 1J), suggesting that SpollIE, SpollIM, and PbpG help maintain compartmentalization, by counteracting the effects of PG hydrolysis during engulfment.

SpollIM and PbpG Are Required for the Dispersal of SpollIE Complexes

Based on the loss of compartmentalization in the $\Delta spolllE$ mutant, it was previously hypothesized that the chromosome





Figure 2. SpollIE Foci Persist in the Absence of SpollIM and PbpG

(A) Diagram showing the dynamic localization of SpoIIIE during engulfment.

(B) Representative images of the localization of SpoIIIE-GFP in WT, ΔspoIIIM, ΔpbpG, and ΔspoIIIM ΔpbpG at T3.5. Scale bar, 2 μΜ.

(C) Representative image of the diffused localization of SpoIIIE in WT cells that have completed engulfment. Scale bar, 1 µM.

(D) Average frequency (\pm SD of 3 biological replicates) of cells with SpoIIIE-GFP foci in WT, Δ spoIIIM, Δ pbpG and Δ spoIIIM Δ pbpG during a sporulation time course (n > 1,000 per time course, per strain, per replicate).

(E) Average fluorescence (± STDEVP, *n* > 400, per time point, per strain) intensity of SpoIIIE-GFP foci in WT, Δ*spoIIIM*, Δ*pbpG* and Δ*spoIIIM* Δ*pbpG* during a sporulation time course.

spanning the septum, and/or the absence of SpoIIIE, creates an opening in the septum through which forespore cytoplasmic contents could leak (Hilbert et al., 2004; Liu et al., 2006). We considered the possibility that SpoIIIM and PbpG are required for SpoIIIE stability, and in their absence SpoIIIE becomes unstable, generating this opening. To investigate this possibility, we examined the localization of a functional SpoIIIE-GFP fluorescent fusion (Burton et al., 2007) during a sporulation time course in WT, Δ spoIIIM, Δ pbpG and Δ spoIIIM Δ pbpG mutant cells (Figure 2). Consistent with previous work, in WT cells, SpoIIIE-GFP localizes as a discrete focus at the center of the asymmetric septum (Figures 2A and 2B) (Burton et al., 2007; Wu and Errington, 1997; Sharp and Pogliano, 1999). As cells progress through engulfment, the focus is often located in the engulfing membranes, and as engulfment nears completion, the focus disperses, with SpoIIIE-GFP localizing as diffuse signal in the forespore membranes (Figures 2A–2C). SpoIIIE-GFP dispersal is thought to represent detachment of SpoIIIE from the DNA (Ben-Yehuda et al., 2003). Consistent with this idea and previous reports (Sharp and Pogliano, 1999), the number of WT sporulating cells with foci declined over time (at T3.5, 35% of the cells had a SpoIIIE-GFP focus) (Figures 2B and 2D). Importantly, SpoIIIE-GFP formed a discrete focus in $\Delta spoIIIM$ and $\Delta spoIIIM \Delta pbpG$ sporulating cells that resembled the focus observed in WT (Figure 2B). Furthermore, quantification of SpoIIIE-GFP foci revealed similar signal intensities in WT and mutant strains (Figure 2E). Thus, the miscompartmentalization in these mutants cannot be explained by instability of SpoIIIE. Interestingly, in the $\Delta spoIIIM$ and $\Delta spoIIIM \Delta pbpG$ mutants, the number of cells with SpoIIIE-GFP foci persisted for a longer

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period of time: at T3.5, 52% and 84% of the sporulating cells, respectively, contained a SpolIIE-GFP focus (Figures 2B and 2D). Thus, SpolIIM and PbpG are required for SpoIIIE dispersal into the forespore membranes. These results argue that forespore miscompartmentalization in cells lacking SpoIIIM and PbpG is not due to instability of SpoIIIE, suggesting that these factors play a more direct role in maintaining compartmentalization at a septal pore.

SpollIM and PbpG Are Required for Efficient Chromosome Translocation and Retention of the Chromosome in the Forespore

The data above suggest that SpollIE can still assemble into complexes in sporulating cells lacking SpollIM and PbpG. To investigate if SpoIIIE is capable of DNA translocation in these mutants, we used an approach based on the Lacl-lacO system (Figure 3A) and engineered strains to produce LacI-GFP under the control of a forespore reporter (PspollQ-lacl-gfp) inserted at a chromosomal locus (amyE), present in the forespore at the time of asymmetric division (Wu and Errington, 1998; Marguis et al., 2008). These strains were also engineered to contain 48 lacO repeats at the pelB locus, which resides near the terminus (Figure 3A). Using this approach, if the *pelB* locus is translocated into the forespore, a GFP focus will be observed in the forespore; if not, then no GFP focus will be observed (Figure 3A). Due to miscompartmentalization in *AspolIIM* and *AspolIIM* Δ*pbpG* mutants, we would also expect to observe some sporulating cells with a GFP focus bound to the mother cell chromosome, in addition to the one in the forespore (Figure 3A). Furthermore, if cells fail to translocate the chromosome, we would expect to observe two GFP foci in the mother cell (Figure 3A).

In WT sporulating cells, DNA translocation takes ~20 min (Burton et al., 2007). Accordingly, virtually all forespores contained a GFP focus at all time points examined, indicating that the chromosome terminus was efficiently translocated into the forespore (Figures 3B and 3C). In Δ spollIM at T2, virtually all forespores (97%) contained a GFP focus, with a small fraction of cells containing two foci in the mother cell (3%) (Figure 3C). This suggests that the majority of cells translocate the chromosome into the forespore in the absence of SpollIM. Interestingly, at T2.5 the number of cells with a GFP focus in the forespore decreased to 90%, while the number of cells with two GFP foci in the mother cells increased to 10%. A similar trend was observed at T3. These data suggest that in the absence of SpolIIM the translocated chromosome is actively or passively effluxed back into the mother cell.

A similar but more dramatic trend was observed in the $\Delta spollIM \Delta pbpG$ double mutant (Figures 3B and 3C). At T2, the majority of the $\Delta spolIIM \Delta pbpG$ double mutant cells had translocated their chromosome (76% contained a GFP focus in the forespore). Remarkably, at T2.5 and T3, we observed a sharp decline in the number of cells with a GFP focus in the forespore and an increase in the number of cells with two mother cell foci (38% and 50% at T2.5 and T3, respectively) (Figures 3B, 3C, and 3E). Thus, most $\Delta spolIIM \Delta pbpG$ cells are capable of translocating the chromosome into the forespore, but they fail to retain it. Finally, we note that although the vast majority of sporulating cells lacking PbpG successfully translocate a chromo-

some into the forespore, a small fraction of cells had two foci in the mother cell at T3 (2%) (Figures 3B, 3C, and 3E). Thus, even in the absence of pbpG, a small proportion of cells exhibit chromosome translocation defects.

Remarkably similar results were obtained when we examined chromosomes harboring the *lacO* array adjacent to the origin at -7° (*yycR*) (Figure 3D), at -120° (*yrvN*) or $+87^{\circ}$ (*yhdG*) (Figures S3C and S3D). These results suggest that the entire forespore chromosome is effluxed into the mother cell in the absence of SpoIIIM and PbpG. Furthermore, analysis of origin-proximal (*yycR*) and terminus-proximal (*pelB*) loci in the *ΔspoIIIM ΔpbpG* cells, using different fluorescently labeled DNA-binding proteins (LacI-YFP and TetR-CFP), revealed a pattern of YFP and CFP foci that is more consistent with passive efflux of the chromosome than active reverse translocation (Figure S3E) (see Discussion). Collectively, these data suggest that the septal pore generated in sporulating cells lacking SpoIIIM and PbpG results in forespore chromosome loss, in addition to loss of its cytoplasmic contents.

Impairing Engulfment Promotes Compartmentalization and Efficient Chromosome Translocation in Cells Lacking SpollIM and PbpG

Our data indicate that blocking engulfment PG hydrolysis almost fully restores compartmentalization to *AspolIIM* and *AspolIIM* $\Delta pbpG$ cells. If forespore chromosome loss in these mutants is due to passive movement through a septal pore, blocking PG hydrolysis should similarly prevent chromosome loss from the forespore. This is indeed what we found. Sporulating cells lacking SpolID and SpolIP almost fully restored chromosome translocation, and retention, to the $\Delta spolliM$ and $\Delta spolliM$ $\Delta pbpG$ mutants (Figure 4A). Furthermore, using a mutant ($\Delta spollB$) that impairs but does not abolish PG hydrolysis during engulfment (Perez et al., 2000) (Figure 4C), we found that both miscompartmentalization and forespore chromosome efflux were largely suppressed in the mutant backgrounds (Figures 4B and 4C). Collectively, these data suggest that SpollIM and PbpG function to maintain compartmentalization of forespore cytoplasmic contents, including the chromosome, and counterbalance the activity of the PG hydrolases that remodel the septum during engulfment.

Blocking Chromosome Translocation Partially Suppresses Miscompartmentalization in Cells Lacking SpollIM and PbpG

Based on recent data showing that chromosome translocation into the forespore generates turgor pressure on the septal PG (Lopez-Garrido et al., 2018), we tested if chromosome translocation into the forespore contributes to miscompartmentalization in cells lacking SpoIIIM and PbpG. If so, a SpoIIIE mutant (*spoIIIE36*) that produces a stable SpoIIIE focus and does not impact compartmentalization but fails to translocate the chromosome (Sharp and Pogliano, 1999; Besprozvannaya et al., 2014) should suppress miscompartmentalization in cells lacking SpoIIIM and PbpG. Indeed, *spoIIIE36* partially suppressed miscompartmentalization in the *ΔspoIIIM* and *ΔspoIIIM* pbpG mutants (Figures 4E and 4F). This result suggests that SpoIIIM and PbpG also function to counterbalance the effect chromosome translocation has on the septal pore.



Figure 3. Efficient Chromosome Translocation Requires SpollIM and PbpG

(A) Diagram explaining experimental rationale of the Lacl-*lacO* system for visualizing chromosome translocation. Lacl-GFP, expressed in the forespore (P_{spollQ}) from the *amyE* locus, binds to *lacO*48 sites inserted at the *yhdG*, *pelB*, *yrvN*, or *yycR* chromosomal locus. The origin (*oriC*) and terminus (*ter*) loci are shown in blue and red, respectively. Successful chromosome translocation is indicated by a Lacl-GFP focus in the forespore, with an additional focus in the mother cell compartment in the event of miscompartmentalization. Unsuccessful chromosome translocation is indicated by an absence of Lacl-GFP foci or by two Lacl-GFP foci in the mother cell compartment in the event of miscompartmentalization.

(B) Representative images of LacI-GFP foci at T3 in WT, Δ*spoIIIM*, Δ*pbpG* and Δ*spoIIIM* Δ*pbpG* strains containing *lacO*48 inserted at the *pelB* locus (174°). Yellow arrowheads point to cells with two mother cell foci. Scale bar, 2 μM.

(C and D) (C) Average frequency (\pm SD of 3 biological replicates) of cells with a LacI-GFP focus in the forespore (successful translocation) or with no or two LacI-GFP foci in the mother cell (unsuccessful translocation, efflux) during a sporulation time course, with *lacO*48 integrated at the *pelB* locus (174°) and (D) at the *yycR* locus (-7°) in WT, Δ spolIIM, Δ pbpG and Δ spolIIM Δ pbpG (n > 650 per time course, per strain, per replicate).

(E) Representative images of cells at T3 with two LacI-GFP foci in the mother cell of $\Delta spollIM$, $\Delta pbpG$ and $\Delta spolIIM$ $\Delta pbpG$ sporulating cells containing *lacO48* inserted at the *pe/B* locus (174°). Scale bar, 1 μ M.

SpolIQ Is Required for Septal Pore Stability during Engulfment in Cells Lacking PbpG, SpolIIM, or SpolIIE

Since slowing down engulfment by reducing the efficiency of PG hydrolysis suppressed miscompartmentalization in the $\Delta spoIIIM$ and $\Delta spoIIIM \Delta pbpG$ mutants, we wondered if this would also be the case in sporulating cells lacking the SpoIIIAH-SpoIIQ zipper-like interaction. As described in the introduction, this zipper-like interaction contributes to the efficiency of engulfment (Broder and Pogliano, 2006; Ojkic et al., 2014). To this end, we intro-

duced the $\Delta spolIQ$ mutation into the $\Delta spolIIM$ and $\Delta pbpG$ single mutants, and the $\Delta spolIIM \Delta pbpG$ double mutant backgrounds, and examined compartmentalization with our forespore reporter (P_{spolIQ} -cfp) during a sporulation time course (Figure 5).

Surprisingly, instead of suppressing the compartmentalization defects, the $\Delta spol/Q$ mutant enhanced them. Among the sporulating cells with CFP signal, a subset appeared to completely lack a forespore compartment (Figure 5B). Since σ^{F} is required for the production of CFP and σ^{F} only becomes

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active after asymmetric division, sporulating cells with this phenotype must have developed an asymmetric septum, which then retracted. Indeed, closer inspection of these cells revealed vestiges of septum formation (Figure 5D). We refer to this phenotype as septal retraction (Figure 5A). Quantification of septal retraction during a sporulation time course revealed that the phenotype increases over time (Figure 5E) and represented 49% and 33% of the sporulating cells at T3, in the $\Delta spolIQ \Delta spolIIM$ and $\Delta spolIQ \Delta pbpG$ double mutant, respectively. In the $\Delta spolIQ \Delta spolIIM \Delta pbpG$ triple mutant, almost all cells (98%) displayed this phenotype by T3 (Figure 5E).

Since the $\Delta spollIM \Delta pbpG$ double mutant phenotypes closely resemble those in the $\Delta spoll E$ mutant, we examined our forespore CFP reporter in the AspolIQ AspolIIE double mutant. Strikingly, the AspolIQ AspolIIE mutant had a similar frequency of septal retraction and vestiges of septa (Figures 5B, 5D, and 5E). TEM confirmed the presence of retracted septa in the $\Delta spollQ \Delta spolliM \Delta pbpG$ triple mutant and $\Delta spollQ \Delta spollE$ double mutant (Figure S4A). Importantly, septal retraction in Δ spollQ Δ spollIM and Δ spollQ Δ pbpG double mutants or the $\Delta spollQ \Delta spolliM \Delta pbpG$ triple mutant was not due to destabilization of SpoIIIE, as SpoIIIE-GFP still localized as foci in cells with retracted septa (Figure S4B). In most cells with retracted septa, regardless of the mutant, SpollIE-GFP was observed as a single focus (Figures S4B-S4D). Interestingly, in a small fraction of cells, two SpolIIE-GFP foci were observed (Figures S4B-S4D). Three-dimensional structured illumination microscopy (3D-SIM) confirmed these observations and suggest that SpollIE-GFP foci reside at the edges of an enlarged septal pore in the $\Delta spollQ \Delta spollM \Delta pbpG$ triple mutant (Figures S4E and S4F).

Finally, we tested whether PG hydrolysis was required for septal retraction. In support of the idea that septal retraction requires PG hydrolysis, retraction was completely abolished and compartmentalization restored in all strains examined, that lack SpoIID and SpoIIP (Figure 5C). Collectively, these data suggest that the septal membranes have not undergone fission at the time of chromosome translocation and that SpoIIIE, PbpG, SpoIIIM, and the zipper-like interaction between SpoIIIAH and SpoIIQ, all function to prevent the septal pore from enlarging during hydrolysis of the surrounding PG.

Evidence that SpollIM, PbpG, and SpollIE Coordinate Pore Closure with Completion of Chromosome

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Building on the observations we describe thus far and taking advantage of the septal retraction phenotype, we tested the hypothesis that SpoIIIM and PbpG coordinate septal pore closure with completion of chromosome translocation. We hypothesized that sporulating cells lacking SpoIIIM and/or PbpG are unable to coordinate septal pore closure with completion of chromosome translocation during engulfment PG hydrolysis, and if these cells also lack SpoIIQ, the membranes that line the pore retract. We reasoned that if we hold back engulfment PG hydrolysis in the absence of SpoIIIM and PbpG, then the septal pore would have sufficient time to close and septal retraction would occur less frequently, unless chromosome translocation is yet to be completed and the septal pore is still open. This is indeed what we found, using the experimental system described in Figure S5A.

We performed this assay on strains that were otherwise WT, or lacked *spolIQ* or both *spolIQ* and *pbpG*. We induced the PG hydrolases at two time points after the onset of sporulation, T2 and T3, to give the cells different amounts of time to complete septal pore closure. At these time points, a similar proportion of the sporulating cells have two septa and CFP signal in the two forespore compartments (>86% for all strains) (Figures S5CA and S5E). Thus, after induction of the PG hydrolases, an increase in sporangia with CFP signal that have a single septum or lack septa would suggest septal retraction and failure to complete septal pore closure (i.e., membrane fission).

When the PG hydrolases were induced at T2 (Figures S5B and S5C) and cells with CFP signal were examined one or 2 h after induction (T3 and T4), in an otherwise WT background or in the $\Delta spol/Q$ mutant, most cells still had two septa and there was only a moderate increase in the number of cells with a single septum or no septa (i.e., the septa had retracted). By contrast, in the $\Delta spol/Q \Delta pbpG$ double mutant cells, the proportion of cells with a single septum or no septa was higher (Figure S5G).

When the PG hydrolases were induced at T3 (Figures S5D and S5E) and cells with CFP signal were examined 1 or 2 h after induction (T4 and T5), although the overall pattern of septal retraction between the strains was similar to when induction occurred at T2 (more septal retraction in $\Delta spol/Q \Delta pbpG$ cells, than in otherwise WT and $\Delta spol/Q$ cells), there was a reduction in the

Figure 4. Blocking or Impairing Engulfment Restores Efficient Chromosome Translocation and Compartmentalization in the Absence of SpolIIM and PbpG

⁽A) Average frequency (\pm SD of 3 biological replicates) of cells with a LacI-GFP focus in the forespore or with two LacI-GFP foci in the mother cell during a sporulation time course, with *lacO48* integrated at the *pelB* locus (174°) in *\DeltaspolID* \DeltaspolIP alone or combined with \DeltaspolIIM, \DeltapbpG and \DeltapbpG \DeltaspolIIM (n > 600 per time course, per strain, per replicate).

⁽B) Average frequency (\pm SD of 3 biological replicates) of cells with a LacI-GFP focus in the forespore or with two LacI-GFP foci in the mother cell during a sporulation time course, with *lacO48* integrated at the *pelB* locus (174°) in Δ *spolIB* alone or combined with Δ *spolIIM*, Δ *pbpG* and Δ *spolIIM* Δ *pbpG* (n > 600 per time course, per strain, per replicate).

⁽C) Schematic representation of the slower engulfment and membrane bulging phenotype observed in the Δ spollB. Cyan represents forespore-specific CFP. (D) Representative images of sporulating cells at T3 showing compartmentalization of forespore gene expression in Δ spollB mutant alone or combined with Δ spollIM, Δ pbpG, Δ spollIM Δ pbpG and Δ spollIE. Scale bar, 2 μ M.

⁽E) Average frequency (\pm SD of 3 biological replicates) of miscompartmentalized cells during a sporulation time course in the *spolllE*36 mutant in otherwise WT (gray), $\Delta spollIM$ (blue), $\Delta pbpG$ (green), and $\Delta spollIM \Delta pbpG$ (red) strains (n > 750 per time course, per strain, per replicate).

⁽F) Representative images of miscompartmentalization in the *spolllE*36 mutant in otherwise WT, $\Delta spollIM$, $\Delta pbpG$ and $\Delta spollIM$ $\Delta pbpG$ strains at T3.5. Scale bar, 2 μ M.

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Figure 5. Septal Retraction and Its Dependency on PG Hydrolysis

(A) Schematic representation of septal retraction, illustrating that as septal retraction progresses, CFP fluorescence (cyan) leaks from the forespore to fill the entire cell.

(B) Representative images of septal retraction in WT, $\Delta spollIM$, $\Delta pbpG$, $\Delta spolIIM$ $\Delta pbpG$ and $\Delta spolIIE$ in a $\Delta spolIQ$ background at T3. Scale bar, 2 μ M. Sporulation efficiency (%, average \pm SD, n = 3) is shown below the respective strains.

(C) Representative images of septal retraction suppression in cells blocked for engulfment in WT, $\Delta spollIM$, $\Delta pbpG$, $\Delta spolIIM \Delta pbpG$ and $\Delta spolIIE$ strains at T3. Scale bar, 2 μ M.

(D) Representative zoomed-in examples of septal retraction in Δ spolIIM, Δ pbpG, Δ spolIIM Δ pbpG, and Δ spolIIE at T3. Yellow arrowheads point to retracting septa. Scale bar, 1 μ M. Schematic representations of cells are shown on the right.

(E) Average frequency (\pm SD of of 3 biological replicates) of cells exhibiting septal retraction in WT, Δ spollIM, Δ pbpG, Δ spollIM Δ pbpG and Δ spollE during a sporulation time course (n > 600 per time point, per replicate).

number of retracted septa for all strains (Figure S5G). Thus, given sufficient time, the septal pore will undergo closure in the absence of SpolIQ, PbpG, and SpolIIM.

Importantly, since $\Delta spollQ \ \Delta pbpG$ sporulating cells have a higher frequency of retracted septa but a similar frequency of fully translocated chromosomes as otherwise WT or $\Delta spollQ$ sporulating cells at both time points of PG hydrolase induction (Figures S5F and S5G), these data suggest that septal pore closure is delayed relative to completion of chromosome translocation in sporulating cells lacking SpolIIM and PbpG.

Finally, we performed these experiments in $\Delta spollQ \Delta spollE$ sporulating cells that do not translocate the forespore chromosome. Interestingly, the $\Delta spollQ \Delta spollE$ mutant behaved simi-

larly to the $\Delta spollQ$ $\Delta pbpG$ mutant (i.e., septal retraction decreased when the PG hydrolases are induced at T3; Figures S5B–S5E and S5G). Thus, given sufficient time, the septal pore will undergo closure even in the absence of SpollIE (and in the presence of untranslocated DNA) (Figures S5F and S5H). This result establishes that SpollIE is not absolutely required for septal membrane fission.

PG Synthesis by PbpG Is Required for Septal Pore Closure

Our data point to the possibility that PbpG contributes to PG synthesis around the septal pore, to maintain pore stability and drive pore closure. If this is the case, then the catalytic activity of PbpG





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should contribute to closure of the septal pore. To test this idea, we generated a mutant of PbpG (PbpG^{E94A S365A}, referred to as PbpG*) harboring alanine substitutions at predicted catalytic residues in its glycosyltransferase and transpeptidase domains (Figure S6A), in the context of a functional N-terminal, hexahistidinetagged version of PbpG (His6-PbpG) (Figure S6C). We note that PbpG* may still retain some residual catalytic activity, since it partially complemented the pbpG-null allele in a $\Delta pbpG \ \Delta pbpF$ (0.16%) (compared with 0.01% of the non-complemented $\Delta pbpG$ $\Delta pbpF$ double mutant) and $\Delta spollIM \Delta pbpG$ double mutant background (57.4%) (compare Figures S6C with 1F). We examined the capacity of PbpG* to support septal pore closure by examining the different phenotypes reported thus far in the presence and absence of SpolIIM: compartmentalization (Figures 6A and S6D), SpolIIE localization (Figure 6C), chromosome translocation (Figure 6B), and septal retraction (Figures 6D and S6D). For all phenotypes tested, the catalytic activity of PbpG was required, although PbpG* produced phenotypes that were less penetrant than those observed in the pbpG-null mutant, except in the case of SpoIIIE localization and septal pore retraction, where it phenocopied the pbpG null. Thus, PbpG itself, independent of its catalytic activity, is also required for the phenotypes described here. For reasons unknown, we were unable to detect His6-PbpG with His6 antibodies by immunoblot and thus confirm the levels of PbpG*. However, PbpG* in the context of a partially functional but detectable, C-terminal, hexahistidine-tagged version of PbpG (PbpG-His6) produced similar levels of protein as the nonmutated counterpart (Figure S6B). Thus, we conclude that PbpG catalytic mutations do not impact protein levels. Collectively, these results support the idea that PG synthesis by PbpG contributes to septal pore closure.

SpollIM and PbpG Interact with SpollIE

If SpoIIIM plays a role in septal pore closure, then it would likely localize to the septal membrane and exhibit a similar localization pattern as SpoIIIE. Consistent with the idea that SpoIIIM is anchored in the membrane with its LysM domain surface exposed, we found that a functional SpoIIIM-His6 fusion (Figure S7C) was membrane-associated and susceptible to trypsin cleavage in a protease accessibility assay (Figure S7A). To determine SpoIIIM's localization, we constructed a GFP-SpoIIIM fluorescent fusion and examined its localization over a sporulation



time course (Figures 6E and 6F). GFP-SpolIIM is fully functional as the sole source of SpoIIIM in otherwise WT cells and partially functional in the absence of PbpG (Figure S7C). In cells that had just begun engulfment, GFP-SpolIIM localized as a faint septal focus or septal band (Figure 6E). A similar localization pattern was observed at intermediate stages of engulfment (Figure 6E). As engulfment neared completion, fewer cells had GFP-SpoIIIM signal (Figures 6E and 6F). GFP-SpolIIM localization was reminiscent of the dynamic localization of GFP-SpolIIE, suggesting that SpollIM may depend on SpollIE for its localization. To test this, we examined GFP-SpoIIIM localization in *AspoIIIE* over a sporulation time course (Figures 6E and 6F). In AspoIIIE cells, we rarely observed GFP-SpolIIM foci; instead, most cells contained GFP-SpoIIIM septal bands (Figure 6E). Furthermore, the frequency of cells with GFP-SpolIIM signal in ΔspolIIE cells decreased earlier relative to WT cells (Figure 6F).

To validate the localization relationship between SpoIIIE and SpollIM we took advantage of the spollIE36 mutant, which, unlike SpolIIE, fails to disassemble as engulfment nears completion but instead persists as a stable focus (Sharp and Pogliano, 1999). If SpollIM localization depends on SpollIE, then in the spollIE36 mutant, GFP-SpolIIM localization might also persist. We examined GFP-SpoIIIM localization in the spoIII36 mutant over a sporulation time course (Figure 6F). Quantification of the number of cells with GFP-SpolIIM signal (focus and bands) in the spollIE36 mutant revealed that GFP-SpolIIM foci persisted for a longer period of time (Figures 6E and 6F). Similar observations, both in the $\Delta spollE$ and the spollE36 mutant background, were made in a merodiploid background using a stronger mother cell promoter to drive GFP-SpolIIM (Figure S7D). Furthermore, immunoblot analysis of the SpollIM-His6 fusion confirmed that SpoIIIM stability depends on SpoIIIE (Figure S7B). Collectively, these results suggest that SpollIM localization depends on SpolIIE, and these proteins form a complex in vivo.

To investigate if SpoIIIM and SpoIIIE interact directly, we used bacterial two-hybrid assays in *Escherichia coli*. In this assay, *E. coli* strain BTH101 expressed SpoIIIE and SpoIIIM fusions to domains T25 and T18 of *Bordetella pertussis* adenylate cyclase (Karimova et al., 1998). Positive interactions between SpoIIIE and SpoIIIM reunite T25 and T18 domains, resulting in *lacZ* expression, β -galactosidase production and blue

Figure 6. Peptidoglycan Synthesis by PbpG Is Required for Developmental Characteristics and Evidence that SpollIE, SpollIM, and PbpG Form a Complex

⁽A) Average frequency (\pm SD of 3 biological replicates) miscompartmentalized cells during a sporulation time course in WT, Δ spolIIM, PbpG* and Δ spolIIM PbpG* strains (n > 900 per time course, per strain, per replicate).

⁽B) Average frequency (\pm SD of of 3 biological replicates) of cells with a LacI-GFP focus in the forespore (successful translocation) or with no or two LacI-GFP foci in the mother cell (unsuccessful translocation, efflux) during a sporulation time course, with *lacO*48 integrated at the *yycR* locus (-7°), in WT, Δ spolIIM, PbpG*, and Δ spolIIM PbpG* (n > 600 per time course, per strain, per replicate).

⁽C) Average frequency (\pm SD of 3 biological replicates) of cells with SpoIIIE-GFP foci during a sporulation time course in WT (gray), Δ spoIIIM, PbpG^{*}, and Δ spoIIIM PbpG^{*} (n > 950 per time course, per strain, per replicate).

⁽D) Average frequency (± SD of of 3 biological replicates) of sporulating cells exhibiting septal retraction during a sporulation time course in WT, ΔspolIIM, PbpG*, and ΔspolIIM PbpG* (n > 700 per time course, per strain, per replicate).

⁽E) Representative images of GFP-SpoIIIM localization in WT, $\Delta spoIIIE$ and spoIIIE36 strains at T2 and T3. Yellow arrowheads indicate GFP-SpoIIIM fluorescence that appears to disperse, disappear, or remain as a stable focus as engulfment nears completion in WT and the *spoIIIE36* mutant. White arrowheads indicate GFP-SpoIIIM fluorescence that appears as a septal band in the $\Delta spoIIIE$ mutant. Scale bar, 2 μ M.

⁽F) Average frequency (± SD of 3 biological replicates) of cells with GFP-SpolIIM fluorescence in WT, ΔspolIIE and spolIIE36 during sporulation time course (n > 750 per time course, per strain, per replicate).

⁽G) Bacterial two-hybrid assay of T18 and T25 fusions to SpolIIE, SpolIIM, and PbpG. T18-SImA and T25-SImA interaction is included as a positive control (Cho et al., 2011).





(B) Interaction network of SpolIIE, highlighting its role in chromosome translocation and its role in PG synthesis via interactions with SpolIIM and PbpG. We hypothesize that a putative unidentified PG synthase (PBP) may also be involved in PG remodeling via interaction with SpolIIM.

(C) Schematic representation of the septal membrane, before and after initiation of engulfment and how different molecular components contribute to ensuring developmental compartmentalization and chromosome translocation upon initiation of engulfment. Note that in the $\Delta spol/Q \Delta spol/M \Delta pbpG$ triple mutant, SpolIIE is still present as a focus in retracted septa (Figure S4), but for simplicity this is not shown in the bottom panel.

colonies on LB agar containing X-gal. We observed positive interactions when T25 and T18 were fused to SpoIIIE and SpoIIIM (Figure 6G). Interestingly, although PbpG appears to localize uniformly in the engulfing membranes (Ojkic et al., 2016), we also detected a positive interaction between PbpG and SpoIIIE (Figure 6G). Collectively these results suggest that SpoIIIE, SpoIIIM, and PbpG function together in maintaining PG synthesis around the septal pore during chromosome translocation, thereby ensuring compartmentalization in the face of septal PG hydrolysis.

DISCUSSION

Critical to a better understanding of the mechanisms coordinating cytokinesis with chromosome segregation during development is a detailed appreciation of these mechanisms in simpler organisms, such as bacteria. Our data demonstrate that coordination between cytokinesis and chromosome segregation during bacterial spore development requires an intricate balance between enzymatic remodeling of cell wall material, stabilization of the division septum, and chromosome segregation. Our data indicate

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that SpollIE, SpollIM, and PbpG coordinate forespore chromosome translocation with septal PG remodeling at a highly stabilized septal pore (Figures 7A and 7C). Our observations support a model in which SpolIIE plays two critical roles during sporulation: (1) it translocates a copy of the chromosome into the forespore compartment and (2) it functions to maintain the septal pore through which the chromosome travels during engulfment. This second role is achieved by the recruitment of SpoIIIM and PbpG. Our data further suggest that PbpG synthesizes PG around the septal pore to counteract the activity of the PG hydrolases, while SpollIM, and SpollIE itself, function to reinforce the pore through protein-protein and protein-PG interactions. Finally, we reveal an important role for the highly conserved SpollIAH-SpolIQ zipper-like interaction in the stabilization of the septal membrane surrounding the pore. Together these factors maintain the septal pore and prevent its expansion due to the action of the septal PG hydrolases during engulfment (Figure 7C).

Septal PG degradation by the DMP complex plays a central role in driving engulfment (Khanna et al., 2020). However, as our data suggest, PG degradation must be counteracted to ensure the septal pore, where SpoIIIE translocates the forespore chromosome, does not expand. Our data indicate that SpoIIIE, along with PbpG and SpollIM, are critical for this process and suggest that they act, in part, by restoring PG surrounding the pore that is lost due to the action of the DMP complex (Figure 7B). This model is in keeping with data suggesting that remodeling of the septum during engulfment requires a balance between PG synthesis and hydrolysis (Ojkic et al., 2016) and with cryo-electron tomography data demonstrating changes in septal PG thickness before and after PG hydrolysis (Tocheva et al., 2013). Our data suggest that PbpG is responsible for localized septal PG synthesis and we hypothesize that SpoIIIE and SpollIM function to constrain the pore, possibly by recruiting or activating additional PG synthases in this region (Figure 7B). Consistent with this idea, FtsW, an essential PG glycosyltransferase of the SEDS family, forms a focus in the middle of the septum at the time of asymmetric division (Burton et al., 2007).

A surprising phenotype we observed was the movement of the forespore chromosome back into the mother cell, in cells lacking SpollIM and PbpG (Figures 3, S3C, and S3D). A similar phenotype was observed by Pogliano and co-workers in cells where SpoIIIE is targeted for degradation at the initiation of chromosome translocation (Lopez-Garrido et al., 2018). In this situation, it was proposed that the chromosome is either passively expelled into the mother cell, to minimize the repulsive and hydration forces associated with packing the chromosome into the small forespore, or actively reverse translocated by SpollIE. Thus, it is possible that the movement out of the forespore reported here represents a similar scenario. Interestingly, we found that in cells lacking SpollIM and PbpG, SpollIE is not degraded but resides in active, stable complexes (Figure 2). Thus, we have identified a condition where SpollIE is present and capable of translocating the chromosome into the forespore, but the chromosome fails to remain there. While it is tempting to hypothesize that PbpG and SpolIIM normally function to prevent reverse translocation, we favor a simpler model in which these proteins prevent enlargement of the pore and passive efflux of forespore cytoplasmic content, and its chromosome, into the mother cell during engulfment PG hydrolysis. In support of this model, chromosome efflux in the

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absence of PbpG and SpollIM is suppressed when engulfment PG hydrolysis is blocked or impaired (Figures 4A and 4B).

An unexpected phenotype we identified in this study was that in actively engulfing cells lacking the highly conserved SpollIAH-SpolIQ zipper-like interaction and SpolIIE, or SpolIIM, or PbpG (or both), the asymmetric septum forms (and σ^{F} is activated) but then retracts, abolishing development (Figures 5 and S4A). The SpollIAH-SpollQ zipper-like interaction has been shown to be particularly important in sporulating cells where the PG is degraded by lysozyme (Broder and Pogliano, 2006; Blaylock et al., 2004). Similarly, we hypothesize that it also functions to prevent the septal membranes from retracting during septal PG hydrolysis. As hypothesized above, in the absence of SpollIE, SpollIM, or PbpG there is reduced PG synthesis and destabilization of the septal pore. The reduced PG synthesis, combined with the thinning of septal PG at the onset of engulfment and the decreased membrane stability occurring in the absence of SpolIIAH-SpolIQ zipper, likely leads to progressive pore enlargement as the DMP complex makes its way outward from the middle of the septum. Consistent with this hypothesis, we show that septal retraction can be prevented by blocking engulfment PG hydrolysis (Figure 5C).

Overall, our results provide a more complete picture of how bacteria differentiate into spores, a morphological process that involves an asymmetric septum with a pore that must coordinate multiple developmental processes at once: the translocation of a chromosome across it, the maintenance of compartmentalization, and the dramatic remodeling of the forespore envelope. More broadly, we show how coordination of different processes is critical to the establishment and maintenance of genetic and transcriptional compartmentalization during cytokinesis, key aspects of development and differentiation in all organisms. Finally, our results highlight the importance of modern approaches in genetics for distinguishing between disparate biological models.

SUPPORTING CITATIONS

The following reference appears in the supplemental information: Rodrigues et al., 2013.

STAR*METHODS

Detailed methods are provided in the online version of this paper and include the following:

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AUTHOR CONTRIBUTIONS

A.M., H.C., J.L., E.B., B.G., M.A., S.C., and C.D.A.R. designed and performed the experiments, and analyzed the data; A.M., H.C., C.M., L.C., D.L., D.Z.R., and C.D.A.R. supervised and oversaw the experimentation and data analysis; C.D.A.R. wrote the draft; A.M., H.C., J.L., C.M., L.C., M.A., D.L., and D.Z.R. revised the draft, C.M., D.Z.R., and C.D.A.R. obtained the funding.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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STAR*METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
THE His Tag Antibody, mAb, Mouse	GenScript	Cat#A00186
anti-FtsZ	Hajduk et al., 2019	N/A
anti-SpollIAG	Doan et al., 2009	N/A
Bacterial Strains		
bBB042	Burton et al., 2007	N/A
bBB069	Besprozvannaya et al., 2014	N/A
bDR2413 (168)	Zeigler et al., 2008	N/A
bWX1200	Wang et al., 2014	N/A
See Table S1 for a complete list of strains used in this study.	N/A	N/A
Chemicals		
FM4-64	Thermo Fisher Scientific	Cat#T13320
SYTOX® Orange	Thermo Fisher Scientific	Cat#S11368
TMA-DPH	Thermo Fisher Scientific	Cat#T204
Oligonucleotides		
See Table S2 for a complete list of oligonucleotides used in this study.	Integrated DNA Technologies (IDT)	https://sg.idtdna.com/
Recombinant DNA		
P _{lac} ::T25-sImA (kan)	Cho et al., 2011	pHC535
P _{lac} ::T18-slmA (amp)	Cho et al., 2011	pHC538
See Table S3 for a complete list of plasmids constructed for this study.	N/A	N/A
Software and Algorithms		
Fiji software	Schindelin et al., 2012	https://imagej.net/Welcome
MicrobeJ	Ducret et al., 2016	https://www.microbej.com/

RESOURCE AVAILABILITY

Lead Contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Christopher Rodrigues (christopher.rodrigues@uts.edu.au).

Materials Availability

All materials generated in this study are available from the Lead Contact without restriction.

Data and Code Availability

The datasets generated during this study are available upon request from the Lead Contact.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Bacterial Strains and Growth Conditions

All *B. subtilis* strains used in this study were derived from strain 168 or PY79. *B. subtilis* cells were induced to sporulate via resuspension at 37°C according to the Sterlini-Mandelstam method or through exhaustion of nutrients in supplemented Difco Sporulation medium (DSM). Deletion mutant strains were either obtained from the *B. subtilis* Single Gene Deletion Library (Addgene) (Koo et al., 2017) or from pre-existing strains in the laboratory. All BKE mutants were back-crossed into *B. subtilis* 168 before assaying and prior

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to antibiotic cassette removal. Antibiotic cassette removal was performed using a temperature-sensitive plasmid that constitutively expresses Cre recombinase. Genetic constructs were obtained from previously existing *B. subtilis* strains found within the laboratory or through restriction enzyme-based cloning of PCR products into ectopic-integration, double-crossover plasmid vectors that were then transformed into *B. subtilis*. Tables of strains (Table S1), plasmids (Table S2) and oligonucleotide primers (Table S3) can be found in the Supplemental Information.

METHOD DETAILS

Sporulation Efficiency Assays

Sporulation efficiency was examined using the heat-kill assay of 24-30-hour grown cultures, where the total number of heat-resistant (80°C for 20 min) colony forming units (CFUs) was compared to wild-type heat-resistant CFUs.

Transposon Insertion Sequencing

Transposon insertion sequencing (Tn-seq) was performed on WT (bDR2413), $\Delta pbpG$ (bCR1557) and $\Delta pbpF$ libraries (bCR1558), as described previously (Meeske et al., 2016). About 750,000 transformant colonies were harvested, aliquoted, and frozen. The aliquot culture was thawed on ice, resuspended in DSM, and diluted into 50 mL DSM at an OD600 of 0.05. Samples were collected at T24. The T24 samples were incubated at 80°C for 20 minutes to kill sporulation-defective cells and then plated on LB agar to allow gemination and outgrowth. Approximately, 750,000 colonies were harvested and their genomic DNA extracted and digested with Mmel enzyme to cut the Mmel restriction site inserted at the inverted repeat sequences of the magellan6 transposon. Barcode adapters were ligated and the transposon-chromosome junctions were amplified by PCR using 16 cycles. PCR products were gel-purified and sequenced on the Illumina HiSeq platform using TruSeq reagents (Tufts University TUCF Genomics facility). Reads were mapped to the *B. subtilis* 168 genome (NCBI NC_000964.3), tallied at each TA site.

Fluorescence Microscopy

Live-cell fluorescence imaging was performed by placing cells on a 2% (w/v) agarose pad prepared in resuspension medium and set using a gene frame (Bio-Rad). When sporulating cells reached the desired time-point, 200 μ L of the culture was pelleted by centrifugation, and then resuspended in 10 μ L of resuspension medium containing the membrane dye TMA-DPH (1-(4-trimethylammoniumphenyl)-6-phenyl-1,3,5-hexatriene *p*-toluenesulfonate) (0.05 mM) or FM 4-64 (*N*-(3-Triethylammoniumpropyl)-4-(6-(4-(Diethylamino) Phenyl) Hexatrienyl) Pyridinium Dibromide) (0.67 μ g/ μ L). After gentle vortexing, 2 μ L of the cell suspension was spread on the agarose pad, and a coverslip was placed on top of the gene frame. Cells were imaged by standard epifluorescence microscopy using a Zeiss Axioplan 2 Microscope equipped with 100x objective N/A 1.4. A DAPI filter was used to excite the TMA-DPH membrane dye with an exposure time of 400 ms. CFP, GFP, and YFP filters were used with exposure times of 600 ms, 800 ms, and 1000 ms, respectively. 3D-structured illumination microscopy was performed using the DeltaVision OMX SR microscope equipped with Olympus PlanApo N 60x objective lens N/A 1.42. The 1.515 immersion oil was selected after calculating the refractive index using softWoRx software. mCherry/A568 and GFP/A488 filters were used with exposure times of 15-20 ms and 10-15% intensity (%T).

TEM Imaging

For the images in Figure S2F, sporulating cells were pelleted in 1.5-mL Eppendorf tubes at 10,000 rpm for 3 min and placed into primary fixative, consisting of 2.5% glutaraldehyde in 0.1 M sodium cacodylate buffer, for 2 h at room temperature. The cells were rinsed in fresh sodium cacodylate buffer three times for 15 min each. Secondary fixation was performed using 1% osmium tetroxide and 1.5% potassium ferricyanide in cacodylate buffer for 1 h at room temperature. The cells were then rinsed in three washes of Milli-Q water for 15 min each. The fixed cell pellets were dehydrated by incubating in increasing concentrations of ethanol for 15 min, consisting of 30, 50, 70, 90 and 100% ethanol. Dehydrated cell pellets were incubated in a mixture of LR White resin and ethanol at a ratio of 1:1 for 6 h at room temperature, followed by a 2:1 LR White/ethanol mixture overnight. Cell pellets were incubated in 100% LR White resin for 6 hours, followed by another 100% resin change overnight. The cell pellets were then placed into gelatin capsules in 100% resin and the resin polymerised for 24 h in an oven at 60°C. Resin embedded tissue was sectioned with a Diatome diamond knife using a Leica UCS ultramicrotome. Sections of thickness 70 – 90 nm were collected onto formvar-coated 100 mesh copper grids and stained sequentially with 1% uranyl acetate for 10 min and lead citrate for 5 min. The sections were imaged in a JEOL 1400+ transmission electron microscope at 80kV, and images captured with a digital camera at a resolution of 2K x 2K.

For the images in Figure S4A, Sporulating cells were harvested by centrifugation at 3,220 × g for 10 min. The cell pellet (1.4 μ L) was dispensed on the 200- μ m side of a type A 3-mm gold platelet (Leica Microsystems), covered with the flat side of a type B 3-mm aluminum platelet (Leica Microsystems), and was vitrified by high-pressure freezing using an HPM100 system (Leica Microsystems). Following freeze-substitution at -90° C for 80 h in acetone supplemented with 1% OsO4, the samples were warmed up slowly (1°C/h) to -60° C (AFS2; Leica Microsystems). After 8 to 12 h, the temperature was raised (1°C/h) to -30° C, and the samples were kept at this temperature for another 8 to 12 h before being rinsed 4 times in pure acetone. The samples were then infiltrated with gradually increasing concentrations of resin (Embed812, EMS) in acetone (1:2, 1:1, 2:1 [v/v]) for 3 h while raising the temperature until 20°C. Pure resin was added at room temperature. After polymerization at 60°C, 80-nm-thin sections were obtained using an ultramicrotome UC7 (Leica Microsystems) and were collected on formvar-carbon-coated 100-mesh copper grids. The thin sections were post-stained for 5 min with 5% aqueous uranyl acetate, rinsed, and incubated for 2 min with lead citrate. The samples were observed



using a CM12 (Philips) or Tecnai G2 Spirit BioTwin (FEI) microscope operating at 120 kV with an Orius SC1000B CCD camera (Gatan).

Immunoblot Analysis

Whole-cell lysates from sporulating cells were prepared as previously described (Rodrigues et al., 2016). Samples were heated for 15 min at 50°C prior to loading. Equivalent loading was based on OD₆₀₀ at the time of harvest. Samples were separated on a 12.5% polyacrylamide gel and transferred to a PVDF membrane. Membranes were blocked in 5% non-fat milk with 0.5% Tween-20 for 1 h. Blocked membranes were probed with anti-His (1:4000) (Genscript), anti-SpolIIAG (1:10000) (Doan et al., 2009) or anti-FtsZ (1:30000) (Hajduk et al., 2019) primary antibodies diluted into PBS with 5% non-fat milk (w/v) with 0.05% Tween- 20 at 4°C overnight. Primary antibodies were detected with horseradish-peroxidase conjugated anti-mouse or anti-rabbit antibodies (BioRad) and detected with Western Lightning ECL reagent as described by the manufacturer.

Protease Accessibility Assay

Protease accessibility assays were performed in sporulating cells lacking the SpoIIQ (Q) protein to ensure that the membrane proteins present in the inner and outer forespore membranes would not be artificially inaccessible because of protoplast engulfment (Broder and Pogliano, 2006). Twenty-five milliliters of sporulating cells (induced by resuspension) were harvested by centrifugation at hour 2.5 after the onset of sporulation, washed, and resuspended in 2 mL 1× SMM buffer (0.5 M sucrose, 20 mM MgCl₂, 20 mM maleic acid, pH 6.5). The cells were protoplasted by lysozyme (5 mg/mL final concentration) for 10 min with slow agitation. The protoplasts were harvested by centrifugation and resuspended in 1 mL of 1X SMM. Protoplasts (100 μ L) were incubated with trypsin (30 μ g/mL final concentration) (Worthington), trypsin and Triton X-100 (2% final concentration), or 1X SMM for 30 min at 30°C. Reactions were terminated by the addition of 100 μ L of 2X SDS-sample buffer and incubation for 5 min at 95 °C. Five microliters from each reaction were analyzed by immunoblot.

Plasmid Construction

pAT001a [*ycgO*::*spoIIIM* (*spec*)] was generated in a two-way ligation with *Eco*RI-*Bam*HI PCR product containing the *spoIIIM* gene (oligonucleotide primers oAT001 & oAT002 and 168 genomic DNA as template) and pKM083 cut with *Eco*RI-*Bam*HI. pKM083 (*yc-gO*::*spec*) is an ectopic integration vector for double crossover integration at the non-essential *ycgO* locus.

pAT003a [*ycgO*::*PspoIIIM*-opt_{RBS}-gfp-spoIIIM (spec)] was generated in a three-way ligation with a *Hind*III-*Xhol* PCR product containing opt_{RBS}-gfp (oligonucleotide primers oAT005 & oAT006 and pCR227 DNA as template), and an *Xhol-Bam*HI PCR product containing the *spoIIIM* gene (oligonucleotide primers oAT007 & oAT002) with pAT002a (*ycgO*::*PspoIIIM*) as template cut with *Hind*III and *Bam*HI. pCR227 [*yhdG*::*PgerM-gerM-gfp*] (cat) contains *gfp* (C. Rodrigues - laboratory stock).

pAT024a [*ycgO*::*PspoIID-opt_{RBS}-gfp-spoIIIM* (*spec*)] was generated in a two-way ligation of an *Eco*RI-*Hind*III digest product from pCR113 containing *P_{spoIID}*, into an *Hind*III-*Bam*HI digest product of pAT003a (*ycgO*::*PspoIIIM-opt_{RBS}-gfp-spoIIIM*). pCR113 [*yc-gO*::*PspoIID-malE* (*cat*)] contains the *PspoIID* promoter (C. Rodrigues - laboratory stock).

pAT032a [*ycgO*::*PspoIIIM-spoIIIM-His6* (*spec*)] was generated in a two-way ligation with *Eco*RI-*Bam*HI PCR product containing the *spoIIIM* gene with a C-terminal hexahistidine tag (oligonucleotide primers oAT001 & oCR659 and 168 genomic DNA as template) and pKM083 cut with *Eco*RI-*Bam*HI. pKM083 (*ycgO*::*spec*) is an ectopic integration vector for double crossover integration at the non-essential *ycgO* locus.

pAT057a [*pelB*::*spolIIE-GFP* (*cat*)] was generated in a three-way ligation with a *Eco*RI-*Xhol* PCR product containing *spoIIIE* (oligonucleotide primers oCR694 & oCR695 and bBB042 DNA as template), and an *Xhol-Bam*HI PCR product containing *gfp* (oligonucleotide primers oCR696 & oCR697) with pKM020 (*pelB*::*tet*) as template cut with *Eco*RI and *Bam*HI.

pAT090 [*ycgO*::*spoIIIE36* (*tet*)] was generated in a two-way ligation with an *Eco*RI- *Bam*HI PCR product containing *spoIIIE36* (oligo-nucleotide primers oCR694 & oCR717 and bBB069 DNA as template) with pKM086 (*ycgO*::*tet*) cut with *Eco*RI and *Bam*HI.

pCR20a [amyE::PspoIIQ-spoIVF_{RBS}-lacI-gfp (spec)] was generated in a three-way ligation with HindIII-Xhol digest product containing spoIVF_{RBS}-lacI (oCR416 & oCR417 and bXW1200 DNA as template), Xhol-BamHI digest product containing gfp (oCR418 and oCR353 and pCR154 as template) and pCR154 (amyE::PspoIIQ-opt_{RBS}-gfp (spec) (C. Rodrigues - laboratory stock) cut with HindIII-BamHI.

pJL013 [*pelB*::*spolIIE36 (tet)*] was generated in a two-way ligation with an *Eco*RI-*Bam*HI digest product containing *spolIIE36* from pAT090 (*ycgO*::*spolIIE36*) with pKM033 (*pelB*::*tet*) as template cut with *Eco*RI and *Bam*HI.

pHC3 [*ycgO*::*PspoIIQ-opt_{RBS}-pbpG-His6* (*tet*)] was generated in a two-way ligation with *Hind*III-*Bam*HI PCR product containing the *pbpG* gene [oligonucleotide primers oAT139 & oAT160 and 168 genomic DNA as template] and pCR008 cut with *Hind*III and *Bam*HI. pCR008 [*ycgO*::*PspoIIQ-opt_{RBS}-gfp-spoIIQ* (*tet*)] contains the *PspoIIQ* promoter (C. Rodrigues - laboratory stock).

pHC23 [*ycgO*::*PspolIQ-opt_{RBS}-pbpG-His6* (E94A, S365A) (*tet*)] was generated by sequential rounds of site-directed mutagenesis of pHC3 using oligonucleotide primers oAT96 & oAT97 to introduce the E94A mutation, followed by oAT147 & oAT148 to introduce the S365A mutation.

pHC28 [*P_{lac}*::*T18-spolIIE (amp)*] and pHC32 [*P_{lac}*::*T25-spolIIE (kan)*] were generated in a two-way ligation with Xbal-BamHI PCR product containing the *spolIIE* gene [oligonucleotide primers oCR716 & oCR717 and 168 genomic DNA as template] and pUT18C and pKNT25 (Karimova et al., 1998), respectively, cut with Xbal and BamHI.

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pHC29 [*P_{lac}::T18-spolIIM (amp)*] and pHC33 [*P_{lac}::T25-spolIIM (kan)*] were generated in a two-way ligation with *Xbal-Bam*HI PCR product containing the *spolIIM* gene [oligonucleotide primers oCR718 & oCR719 and 168 genomic DNA as template] and pUT18C and pKNT25 (Karimova et al., 1998), respectively, cut with *Xbal* and *Bam*HI.

pHC30 [*P_{lac}::T18-pbpG (amp)*] and **pHC34** [*P_{lac}::T25-pbpG (kan)*] were generated in a two-way ligation with *Xbal-Bam*HI PCR product containing the *pbpG* gene [oligonucleotide primers oCR720 & oCR721 and 168 genomic DNA as template] and pUT18C and pKNT25 (Karimova et al., 1998), respectively, cut with *Xbal* and *Bam*HI.

Other Constructs

[ycgO::PpbpG-his6-pbpG (erm)] and [ycgO::PpbpG-his6-pbpG^{E94A,S365A} (erm)]

*Hind*III-*PpbpG* was amplified by oAT137 and oHC056 using168 genomic DNA as template. The *his6-pbpG-XhoI* and *his6-pbpG*^{E94A, S365A}-*XhoI* DNA fragments were amplified by oHC055 and oCR704 using pHC3 and pHC23 as templates respectively. *Hind*III-*PpbpG* and *his6-pbpG-XhoI* and similarly *Hind*III-*PpbpG* and *his6-pbpG*^{E94A, S365A}-*XhoI* were ligated by Isothermal assembly using oHC055 and oHC056 and then ligated into pKM084 (*ycgO*::*erm*) cut with *Hind*III and *XhoI*. Finally, after ligation into pKM084, the two constructs were transformed directly into *B. subtilis*.

QUANTIFICATION AND STATISTICAL ANALYSIS

Sporulation Efficiency Analysis

All the sporulation efficiency data are average (± SD) of 3 biological replicates.

Tn-seq Analysis

Mapped genes in which reads were statistically underrepresented were identified using the Mann Whitney U test. Visual inspection of transposon insertion profiles was performed with the Sanger Artemis Genome Browser and Annotation tool.

Image Analysis

Microscopy images were processed by adjusting the brightness and contrast using the Fiji software (Schindelin et al., 2012). All quantifications were performed using the manual counting tool of the Fiji software, and the raw data were then exported into Excel for data collation and graph generation. At least two fields of view (90.13 x 67.53 μ M) were acquired for each biological replicate and contained hundreds of cells, each. The criteria described below, or in results and legends, was used to quantify each phenotype.

- Quantification of forespore miscompartmentalization (Figures 1G, 4E, 6A, S2D, and S3B and): any cell with forespore-expressed CFP signal in the mother cell was classified as miscompartmentalized. The percentage of miscompartmentalized cells (average frequency ± SD of 3 biological replicates) was calculated based on the total number of sporulating cells that had progressed into the asymmetric division stage of development and contained CFP signal. *n*>750 per time-course, per strain, per replicate.
- Quantification of chromosome translocation using the LacI-GFP / *lacO* system (Figures 3C, 3D, 4A, 4B, 6B, S3C, and S3D): we followed the criteria described in Figure 3A and the percentage of cells (average frequency ± SD of 3 biological replicates) was calculated based on the total number of sporulating cells that contained a GFP focus and had progressed into or beyond asymmetric division. *n*>600 per time-course, per strain, per replicate.
- Quantification of chromosome translocation using the dual reporter system (TetR-CFP/ *tetO* & LacI-YFP/ *lacO*): we followed the criteria described in Figure S3E and the percentage of cells (average frequency ± SD of 3 biological replicates) was calculated relative to the total number of sporulating cells that contained four foci, two CFP foci and two YFP foci. Not all cells contained four visible foci within the plane of view during development: the four foci were clearly visible in approximately 25%, 30% and 47% of the sporulating cells, at T2, T2.5 and T3 respectively. n>30 per time-point, per replicate.
- Quantification of septal retraction (Figures 5E and 6D): any cell with forespore-expressed CFP signal in the mother cell and a non-continuous asymmetric septum, or vestiges of an asymmetric septum (based on the membrane signal), was classified as having a retracted septum. The percentage of cells with retracted septa (average frequency ± SD of of 3 biological replicates) was calculated based on the total number of sporulating cells that had progressed into or beyond asymmetric division. n>600 per time-point, per strain, per replicate.
- Quantification of cells with a SpolIIE-GFP focus (Figures 2D and 6C): the percentage of cells with a SpolIIE-GFP focus (average frequency ± SD of of 3 biological replicates) was calculated based on the number of sporulating cells that contained a GFP focus, relative to the total number of sporulating cells that contained SpolIIE-GFP signal (foci or diffuse) in the membranes of the forespore. n>950 per time-course, per strain, per replicate.
- Quantification of GFP-SpolIIM (Figure 6F): the percentage of cells with a GFP-SpolIIM focus (average frequency ± SD of 3 biological replicates) was calculated based on the number of sporulating cells that contained GFP signal (focus or band), relative to the total number of sporulating cells that had progressed into or beyond asymmetric division. *n*>750 per time-course, per strain, per replicate.



The MicrobeJ plugin (Ducret et al., 2016) designed for the Fiji software was used to detect the signal intensity of SpoIIIE-GFP foci (Figure 2E), on background subtracted images (Process > Subtract Background) to avoid false positive detections of the fluorescent signal. To determine the signal intensity of SpoIIIE-GFP foci, the brightfield image was set to Channel 1 and the image with the GFP signal to Channel 2. These channels were then merged into one image. For Channel 1 of the image with "Bright" background, the "Bacteria" tab was set to "Fit Shape" and "Rod-Shaped" to detect the bacteria in the brightfield image. Five parameters: "Exclude on Edges", "Shape descriptors", "Segmentation", "Intensity", and "Feature" were checked on. To refine the generated bacteria outlines, the shape descriptors (area, length, and width) were set differently for each time-point corresponding to the contours of individual cell. To resolve unprocessed cells, the manual editing tool was also used. For Channel 2 of the image with "Dark" background, the "Maxima" tab was set to "Point" and "Basic". The tolerance was set to 15, and the intensity was set from 15-max to ensure detection of the GFP foci with a minimum signal intensity of 15. Three parameters: "Exclude on Edges", "Shape descriptors", and "Associations" were checked on. The raw data were displayed on a MicrobeJ results table, which was then exported into Excel for data collation and graph generation.