Novel Secretion Apparatus Maintains Spore Integrity and Developmental Gene Expression in Bacillus subtilis

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Abstract

Sporulation in Bacillus subtilis involves two cells that follow separate but coordinately regulated developmental programs. Late in sporulation, the developing spore (the forespore) resides within a mother cell. The regulation of the forespore transcription factor σ⁵ is crucial at this stage. σ⁵ activity requires eight mother-cell proteins encoded in the spoIIA operon and the forespore protein SpoIIQ. Several of the SpoIIA proteins share similarity with components of specialized secretion systems. One of them resembles a secretion ATPase and we demonstrate that the ATPase motifs are required for σ⁵ activity. We further show that the SpoIIA proteins and SpoIIQ reside in a multimeric complex that spans the two membranes surrounding the forespore. Finally, we have discovered that these proteins are all required to maintain forespore integrity. In their absence, the forespore develops large invaginations and collapses. Importantly, maintenance of forespore integrity does not require σ⁵. These results support a model in which the SpoIIA-SpoIIQ proteins form a novel secretion apparatus that allows the mother cell to nurture the forespore, thereby maintaining forespore physiology and σ⁵ activity during spore maturation.

Introduction

How cells communicate with each other is a fundamental biological question relevant to the development of multi-cellular organisms, microbial pathogenesis and communities of microorganisms. In the bacterium Bacillus subtilis, cell-to-cell signaling pathways play a key role in coordinating gene expression during the process of spore formation. Upon the initiation of sporulation, the developing cell divides asymmetrically, generating two cells of unequal size and dissimilar fate: a small cell (the prospective spore), referred to as the forespore, and a large cell called the mother cell [1,2]. Initially, these two cells lie side by side, but shortly after polar division the mother cell membranes migrate around the forespore in a phagocytotic-like process, generating a cell within a cell. As a result of this engulfment process, the forespore is surrounded by two membranes, one derived from the mother cell, the other from the forespore. Shortly after the completion of engulfment, the mother cell packages the forespore in a protective coat while the forespore prepares for dormancy. Once the spore is fully mature, the mother cell lyses releasing it into the environment.

During this developmental process, the mother cell and forespore follow separate programs of gene expression directed by alternative sigma factors. However, cell-to-cell signaling pathways ensure that gene expression in one cell is coordinated with gene expression in the other [2–4]. Shortly after polar division, the first cell-type specific transcription factor (σ⁵) is activated in the forespore. σ⁵ is then responsible for the activation of σ⁶ in the mother cell. At a later stage, σ⁵ is required for the activation of σ⁵ in the forespore. Finally, σ⁵ sets in motion the activation of σ⁵ in the mother cell. The signal transduction pathways that govern the activation of the mother–cell transcription factors (σ⁵ and σ⁵) are well established [1,2,5]. By contrast, the mechanism underlying activation of σ⁵ in the forespore has remained enigmatic.

The late-acting forespore transcription factor σ⁵ appears to be regulated at multiple levels. First, the transcription of the gene encoding σ⁵ (sigG) is under the control of the earlier-acting forespore transcription factor σ⁵ [6,7]. However, unlike the other σ⁵-dependent genes, sigG transcription is delayed by approximately 30 min by an as yet unknown mechanism [8–10]. Once synthesized, σ⁵ activity requires eight mother cell proteins encoded in the spoIIA operon [11] and a forespore membrane protein SpoIIQ [12]. In addition, σ⁵ activity requires proper engulfment. Mutants that block the engulfment process are impaired in σ⁵-dependent gene expression [13–15]. Finally, once activated, σ⁵ recognizes its own promoter and its level increases rapidly in the forespore [6,7].
Author Summary

During development, cell–cell signaling pathways coordinate programs of gene expression in neighboring cells. Cell–cell signaling is typically achieved by the secretion of a signaling ligand followed by its binding to a membrane receptor on the surface of a neighboring cell (or cells). The ligand-bound receptor directly or indirectly triggers transcription factor activation. Here we present evidence for a non-canonical signaling pathway that links development of gene expression in the forespore and mother cell during spore formation in Bacillus subtilis. Our data support a model in which a novel secretion system is assembled in the double membrane that encases the spore within the mother cell. This apparatus is not involved in transducing a specific activating signal but rather allows the mother cell to nurture the spore and thereby maintain the spore program of developmental gene expression.

It is not known how the SpoIII proteins in the mother cell and SpoIIQ in the forespore participate in regulating $\sigma^G$. It has been suggested that they function to transduce a signal from the mother cell to trigger its activation [2,4,11,16–18]. It has also been proposed that these proteins are involved in monitoring the process of engulfment and sending an activating signal to the forespore upon its completion.

Fluorescence microscopy experiments have revealed that SpoIIQ and SpoIIIAG (the last gene in the spoIIIA operon) both localize to the membranes that surround the forespore [19,20]. Moreover, these two membrane proteins can interact across the double membrane [16,20]. The relevance of this interaction for $\sigma^G$ activation has been unclear. One clue to the role of these proteins in $\sigma^G$ activation is that several of the SpoIII proteins share weak similarity with components of specialized secretion systems [16,21,22]. In addition, recent experiments suggest that SpoIIQ forms large pores in the forespore membrane [22]. Based on these observations, it has been proposed that the SpoIII proteins and SpoIIQ form a channel between the mother cell and forespore [21,22]. This apparatus could transduce an activating signal to trigger $\sigma^G$ activation. Camp and Losick have recently found that the activity of transcription factors other than $\sigma^G$ is important for $\sigma^G$ activation [26–28]. The spoIIIAA gene was introduced into a B. subtilis strain harboring an in-frame deletion in the spoIIIAA gene and tested for their ability to activate $\sigma^G$ during sporulation. $\sigma^G$ activity was assessed in single cells by fluorescence microscopy with a $\sigma^G$-responsive promoter (P$_{\sigma^G}$) fused to cfp in a population-based assay using an $\sigma^G$-responsive promoter (P$_{\sigma^G}$) fused to lacZ. Mutations in all four of the conserved motifs blocked $\sigma^G$ activation to the same extent as the spoIIIAA null strain (Figure 1A and 1B, Figure S2). Moreover, the sporulation efficiency of the mutants was similar to the spoIIIAA null strain (Figure 1C and Figure S2). Importantly, almost all of the spoIIIAA mutant proteins were produced at levels similar to wild-type SpoIIIAA (Figure 1C and Figure S2). The NTases involved in type II and type IV secretion have been particularly refractory to biochemical reconstitution and our attempts to reconstitute SpoIIIAA ATPase activity in vitro were unsuccessful. Based on the mutational analysis, we tentatively conclude that SpoIIIAA is an ATPase and that ATPase activity is necessary for $\sigma^G$ activation and efficient sporulation.

Here we show that SpoIIIAA shares strong similarity to ATPases of the type II and IV secretion systems and that the conserved ATPase motifs are required for $\sigma^G$ activity and sporulation. Moreover, we demonstrate that at least six of the SpoIIIA proteins in the mother cell and SpoIIQ in the forespore are present in a multimeric membrane complex that spans the double membrane surrounding the forespore. Finally, we show that SpoIIQ, the SpoIIIA proteins and the ATPase motifs in SpoIIIAA are all required to maintain forespore integrity. In their absence, the forespore develops large invaginations and appears to collapse. Importantly, maintenance of forespore integrity does not require $\sigma^G$, indicating that forespore collapse is not a result of the failure to activate $\sigma^G$ (and the late program of forespore gene expression under its control). Instead, these results suggest that the block to $\sigma^G$ activity in the SpoIIIA and SpoIIQ mutants is a manifestation of the failure to maintain forespore integrity.

Consistent with this idea, we demonstrate that premature synthesis of $\sigma^G$ results in early $\sigma^G$ activity that does not require SpoIIIA or SpoIIQ whereas late $\sigma^G$ activity requires these proteins. Collectively, these data support and extend the feeding-tube model of Camp and Losick in which SpoIIIA-SpoIIQ form a novel secretion apparatus that allows the mother cell to nurture the forespore at late stages of development.

Results

The conserved ATPase motifs in SpoIIIAA are required for $\sigma^G$ activation

It has been reported previously that the first gene in the spoIIIA operon (spoIIIAA) is homologous to secretion superfamily ATPases [16,22,24]. Sequence identity searches indicate that SpoIIIAA most closely resembles proteins in the family of NTases involved in type II and type IV secretion (Figure S1). All members of this family contain four highly conserved motifs: the Walker A and B boxes and two additional motifs called the Aspartate and Histidine boxes. The SpoIIIAA ATPase contains four highly conserved motifs: the Walker A and B boxes and two additional motifs called the Aspartate and Histidine boxes. The Aspartate and Histidine residues in these motifs are present in close proximity to the nucleotide binding pocket and are thought to participate in ATP-binding or hydrolysis [25].

To investigate whether SpoIIIAA is an ATPase and whether this activity is important for $\sigma^G$ activation, we made amino acid substitutions in highly conserved residues in the conserved motifs (K149A in the Walker A box, D224A in the Walker B box, E180Q in the Aspartate box, and H250Y in the Histidine box). Similar mutations in other ATPases have been shown to abrogate ATP binding or hydrolysis [26–28]. The spoIIIAA mutants were introduced into a B. subtilis strain harboring an in-frame deletion in the spoIIIAA gene and tested for their ability to activate $\sigma^G$ during sporulation. $\sigma^G$ activity was assessed in single cells by fluorescence microscopy with a $\sigma^G$-responsive promoter (P$_{\sigma^G}$) fused to cfp in a population-based assay using an $\sigma^G$-responsive promoter (P$_{\sigma^G}$) fused to lacZ. Mutations in all four of the conserved motifs blocked $\sigma^G$ activation to the same extent as the spoIIIAA in-frame deletion (Figure 1A and 1B, Figure S2). Moreover, the sporulation efficiency of the mutants was similar to the spoIIIAA null strain (Figure 1C and Figure S2). Importantly, almost all of the spoIIIAA mutant proteins were produced at levels similar to wild-type SpoIIIAA (Figure 1C and Figure S2). The NTases involved in type II and type IV secretion have been particularly refractory to biochemical reconstitution and our attempts to reconstitute SpoIIIAA ATPase activity in vitro were unsuccessful. Based on the mutational analysis, we tentatively conclude that SpoIIIAA is an ATPase and that ATPase activity is necessary for $\sigma^G$ activation and efficient sporulation.

The SpoIIIA proteins form a multimeric complex in the membranes that surround the forespore

Seven of the SpoIIIA proteins (SpoIIIA through SpoIIIAH) are predicted to be integral membrane proteins and several share weak similarity to components of secretion complexes [21,22] (see Discussion). To investigate whether the SpoIIIA proteins form a membrane complex, we performed co-immunoprecipitation experiments. Membrane extracts were prepared from sporulating cells collected at hour 2.5 of sporulation (see Material and Methods). The membrane proteins were then solubilized using the non-ionic detergent dodecyl maltoside (DDM) and SpoIIAG was immunoprecipitated using anti-SpoIIAG antibody resin. Consistent with the idea that the SpoIIIA proteins form a complex, SpoIIAGE, SpoIIAF and a functional myc-tagged SpoIIAD fusion were co-immunoprecipitated with SpoIIAG (Figure 2A).
Analysis of the immunoprecipitate by Mass Spectrometry also identified SpoIIIAB, for which we do not have an antibody. Importantly, a mother cell membrane protein (SpoIID) required for engulfment that localizes to the membranes surrounding the forespore [29] was not present in the immunoprecipitate (Figure 2A). Moreover, none of the SpoIIIA proteins were found in the immunoprecipitate from a strain lacking SpoIIIAG (Figure 2A). Similar results were obtained when

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**Figure 1.** The ATPase motifs in SpoIIIAA are required for σ^G activity and sporulation efficiency. (A) σ^G activity was assessed in single cells by microscopy using a fluorescent reporter (P_{spoE}-cp) in a wild-type background (wt, BTD2779), a ΔspoIIAA mutant (ΔA, BTD2713) and a spoIIAA_{D224A} mutant (ΔA_{D224A}, BTD2775). Cells were visualized at hour 3 of sporulation. Forespore CFP fluorescence (false-colored green in the lower panel) and the fluorescent membrane dye TMA-DPH (false-colored red) are shown. Scale bar, 1 μm. (B) Expression of a σ^G-dependent sspB-lacZ translational fusion [53] was monitored during a time course of sporulation in a wild-type background (wt, BTD2919), a ΔspoIIAA mutant (ΔA, BTD2917), a spoIIAA_{D224A} mutant (ΔA_{D224A}, BTD2920) and a strain lacking σ^G (ΔsigG, BTD1331). Samples from sporulating cells were taken every hour and β-galactosidase activity (Miller Unit, m.u.) was determined. (C) Immunoblot analysis of whole cell lysates from sporulating cells shown in A using anti-SpoIIAA antibodies. Time (in hours) after the initiation of sporulation is indicated. Sporulation efficiencies of the same strains are shown below the immunoblot.

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SpoIIIA-SpoIIQ Complex Maintains Spore Integrity

SpoIIAG localizes to the membranes that surround the forespore

Previous studies of SpoIIAH and SpoIIQ indicate that these proteins localize to the membranes that surround the forespore [16,20]. Our co-immunoprecipitation data suggest that the other SpoIIA proteins reside in a complex with SpoIIAH and SpoIIQ, and should therefore also be present in these membranes. To investigate this, we analyzed the sub-cellular localization of SpoIIA using a CFP-SpoIIAH fusion. The gene fusion was placed under the control of the native spoIIA promoter and inserted at a non-essential locus in the genome. The fusion partially complemented a spoIIA mutant (2% sporulation efficiency, 1000-fold better than a spoIIA null strain). Analysis of CFP-SpoIIAH by fluorescence microscopy at hour 2 of sporulation revealed that the protein specifically localizes to the engulfing septal membranes (Figure 3). Moreover, the CFP-SpoIIAH fusion had a punctate staining pattern similar to what has been observed previously for SpoIIAH and SpoIIQ [16,19].

To determine whether this localization pattern requires SpoIIQ, SpoIIAH or the other SpoIIA proteins, we analyzed CFP-SpoIIAH in strains lacking these proteins. In the absence of SpoIIAH, SpoIIAH was mislocalized (Figure 3). CFP-SpoIIAH was present in all the cytoplasmic membranes of the mother cell; however, there was still some enrichment of the fusion protein in the membranes surrounding the forespore. Interestingly, proper localization of SpoIIAH did not require the other SpoIIA proteins. The absence of SpoIIAA or SpoIIAE had no impact on CFP-SpoIIAH localization (Figure 3 and data not shown). Moreover, the mislocalization of CFP-SpoIIAH in cells lacking the entire spoIIA operon was indistinguishable from the SpoIIAH mutant (data not shown). Finally, in the absence of the forespore

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spoIIB

spoIIIAB

spoIIIAH

spoIIIAF

spoIIIAE

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membrane protein SpoIIQ, CFP-SpoIIIAG had the most severe mislocalization phenotype with virtually no enrichment of the protein in the forespore membranes (Figure 3). Altogether, these data provide further support for the idea that SpoIIIAG and the rest of the SpoIIIA proteins reside in a multimeric membrane complex with SpoIIIAH and SpoIIQ. Moreover, they suggest that SpoIIIAH serves as an important link between SpoIIQ and the rest of the SpoIIIA complex.

The SpoIIIA and SpoIIQ proteins are required for integrity of the forespore

In the course of our analysis of the SpoIIIA mutants, we discovered that the SpoIIIA proteins play an important role in maintaining forespore integrity after engulfment is complete. During the process of engulfment, cells lacking the SpoIIIA proteins appear similar to wild-type (Figure 3) [31]. However, upon completion of engulfment, the SpoIIIA mutants exhibited morphological defects: the forespores appeared smaller and in many cases collapsed (Figure 4). To more carefully analyze this new phenotype, we used a forespore-specific CFP reporter to label the forespore cytoplasm and a YFP-SpoIVFA fusion that localizes to the mother-cell membranes that surround the forespore [20,32]. In wild-type cells at hour 3 of sporulation when engulfment was complete, the CFP reporter appeared homogenous in the forespore cytoplasm and the YFP-SpoIVFA fusion labeled the forespore membranes in a smooth and continuous fashion (Figure 4 and Figure S5). By contrast, in the SpoIIIA mutants the cytoplasmic CFP signal in the forespore appeared patchy and distorted (Figure 4A). Moreover, the YFP-SpoIVFA fusion localized in bright patches around the forespore (Figure 4B and Figure S5). The merged images revealed that the patchy YFP-SpoIVFA signal was coincident with regions of the forespore that lacked cytoplasmic CFP (Figure 4B and Figure S5). These results suggest that in the absence of the SpoIIIA proteins, the engulfed forespore membranes collapse, forming large invaginations. A similar collapsed forespore phenotype was observed in the absence of SpoIIQ and in a Walker B mutant of the SpoIIIAA protein (Figure 4A and data not shown). Importantly, the forespores in cells lacking σ^II did not display this phenotype and were indistinguishable from wild-type (Figure 4 and Figure S5). Thus, the failure to maintain proper forespore morphology is not due to the inability to activate σ^II. Instead, these results are consistent with a model in which the SpoIIIA-SpoIIQ complex as well as the ATPase activity of SpoIIIAA are necessary to maintain proper forespore integrity.

To more closely compare the collapsed forespores in the various mutants, we analyzed their morphology by electron microscopy. Cells at hour 3 of sporulation were fixed, embedded, sectioned, and stained using standard protocols (see Material and Methods). Analysis of sporulating cells by electron microscopy revealed several morphological defects in the mutants. Consistent with our observations using fluorescence microscopy, the forespores were significantly smaller than wild-type and many displayed large invaginations (Figure 5). Analysis of forespores in strains lacking...
SpoIIIA, SpoIIAB, SpoIIAE or SpoIIQ revealed that 30-52% (n = 50 per mutant) displayed membrane invaginations. Finally, the two membranes that surround the forespore had constrictions and bulges and in some cases appeared to have ruptured (carets in Figure 5). By contrast, the size and morphology of the forespores in the σ^5 mutant were indistinguishable from wild-type as observed previously [6]. Importantly, the phenotypes of the SpoIIIA mutant forespores (including the SpoIIAA ATPase mutant) and the SpoIIQ mutant forespores were comparable (Figure 5), suggesting a defect in a shared function.

Figure 4. Morphological defects in the absence of SpoIIIA or SpoIIQ. (A) Forespore morphology was monitored by fluorescence microscopy at hour 3 of sporulation in a wild-type background (wt, BCM703), a ΔspoIIAB mutant (ΔB, BCM704), a ΔspoIIIA mutant (ΔIII, BCM706), a ΔspoIIQ mutant (ΔIIQ, BCM716), and in a strain lacking σ^5 (ΔsigG, BCM708). All strains contained a forespore reporter (P_spoIIQ-cfp; false-colored blue in the lower panel) to visualize the forespore cytoplasm. The membranes (mb) from the same field were visualized with the fluorescent dye TMA-DPH (false-colored red in the lower panel). Carets highlight examples of “collapsed” forespores. (B) Larger images highlighting the cytoplasmic CFP signal and the localization YFP-SpoIVFA (IVFA; false-colored green) in wild-type (wt, BCM703) and a ΔspoIIAB mutant (ΔB, BCM704). Larger images of all strains can be found in Figure S5. Scale bar, 1 μm.

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Figure 5. Morphological defects in the absence of the SpoIIA or SpoIIQ proteins. Forespore morphology was assessed by electron microscopy at hour 3 of sporulation in wild-type (wt, PY79), a ΔspoIIA mutant (ΔIIA, BDR841), a ΔspoIIAB mutant (ΔB, BTD119), a ΔspoIIQ mutant (ΔIQ, BTD141), a ΔsigG mutant (ΔsigG, BDR104), a spoIIIAΔD224A ATPase mutant (AD224A, BTD2683), and a strain that contains low levels of SpoIIAE (low E, BTD3019). For each strain, two typical forespores are shown in the first two columns. Scale bar, 200 nm. The last column shows a 5× enlargement of the forespore membranes. The carets highlight bulged or ruptured membranes in the mutants.

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All together, these results suggest that the SpoIIIA-SpoIIQ complex (and the ATPase activity of SpoIIIAA) is required to maintain the integrity of the forespore. Importantly, since forespore integrity did not require $\sigma^G$, these results indicate that forespore collapse in the SpoIIIA and SpoIIQ mutants is not a consequence of the failure to activate $\sigma^G$ (and the late program of forespore gene expression under its control). Instead, these results suggest that the block to $\sigma^E$ activity is a manifestation of the failure to maintain forespore integrity.

**Low levels of SpoIIIA proteins can support efficient sporulation**

How might the SpoIIIA-SpoIIQ complex maintain forespore integrity? The localization patterns of SpoIIQ and the SpoIIIA proteins [16,19] suggest that the SpoIIIA-SpoIIQ complex could function in a structural capacity. These complexes are distributed throughout the forespore membranes in what appears to be helical arcs [16,19] and could play a cytoskeletal role in maintaining or reinforcing the architecture of the spore. On the other hand, the similarity of the SpoIIIA proteins to components of specialized secretion systems and the requirement of the ATPase domain of SpoIIIAA for proper forespore morphology argue in favor of an enzymatic function for the complex. For example, if the SpoIIIA-SpoIIQ complex functions as a secretion apparatus, its role in maintaining forespore integrity could be to nurture the forespore by secreting nutrients or osmolytes such as ions, metabolites or small proteins into the forespore as has been proposed by Camp and Losick [21,23].

We reasoned that if the SpoIIIA-SpoIIQ complex plays a structural role in forespore integrity, then reduction in the levels of proteins in the complex might result in a collapsed forespore phenotype and impaired sporulation. By contrast, if this complex plays an enzymatic function during spore maturation, a reduction in protein levels might not significantly affect spore morphogenesis. To investigate this, we generated strains that produced low levels of SpoIIIAA, or SpoIIIAE (see Material and Methods). As shown in Figure 6, the levels of SpoIIIAA and SpoIIIAE in these strains were barely detectable by immunoblot. Serial dilution of the extract from wild-type cells suggests that the levels of SpoIIIAA and SpoIIIAE are 5-10-fold reduced (data not shown). Importantly, despite these low levels, the sporulation efficiency of these strains was similar to wild-type (Figure 6). Moreover, examination of the forespores by fluorescence and electron microscopy indicates that low levels of SpoIIIAA and SpoIIIAE do not affect forespore integrity (Figure 5 and Figure S6). Similar results were obtained using a strain that produced low levels of SpoIIHAF (data not shown). These results suggest that the SpoIIIA-SpoIIQ complex has an enzymatic rather than a structural role in maintaining proper forespore integrity.

**SpoIIIA and SpoIIQ are not required for early $\sigma^G$ activity but are needed to maintain late $\sigma^E$ activity**

Collectively, our results are consistent with the feeding-tube model proposed by Camp and Losick [21,23] in which the putative SpoIIIA-SpoIIQ secretion apparatus secretes nutrients and/or osmolytes into the forespore, “feeding” the spore in its final stages of preparation for dormancy. In the context of this model, the role of the SpoIIIA proteins and SpoIIQ in $\sigma^G$ activation is not to secrete a specific signal into the forespore to trigger transcription factor activation. Instead, the “feeding” of the forespore is itself what allows $\sigma^G$-dependent gene expression. In this scenario, the lack of $\sigma^G$ activity in the SpoIIQ and SpoIIIA mutants is a consequence of the loss of metabolic potential in the forespore. This model is extremely compelling in its parsimony but is a significant departure from current thinking.

All current models for $\sigma^G$ regulation involve an unidentified inhibitor that holds the sigma factor inactive during the engulfment process. Relief of inhibition is then triggered upon the completion of engulfment and/or by a signal that is received from the mother cell. The data in support of such an inhibitor comes from experiments in which the $\sigma^G$ gene was fused to a strong $\sigma^E$-responsive promoter ($P_{sigG}$) that is active immediately after polar division. (The $\sigma^G$ gene is itself recognized by $\sigma^E$ but $\sigma^E$-dependent transcription of $\sigma^G$ occurs only after a 30-minute delay [8,10].) Cells harboring the $P_{sigG}$ fusion synthesize $\sigma^G$.

![Figure 6. Low levels of SpoIIIA or SpoIIIAE support efficient sporulation. Immunoblot analysis of whole cell lysates from sporulating cells. (A) Comparison of SpoIIIA (A) levels in wild-type (wt, PY79), a $\Delta$spoIIIAA mutant (A, BTD117), a strain expressing high levels of SpoIIIA (high, BTD2341), and a strain expressing low levels of SpoIIIA (low, BTD3023). (B) Comparison of SpoIIIAE levels (E) in wild-type (wt, PY79), a $\Delta$spoIIIAE mutant (E, BTD113), a strain expressing high levels of SpoIIIA (high, BTD2349), and a strain expressing low levels of SpoIIIA (low, BTD3019). The levels of $\sigma^E$ are similar in all lysates indicating that all strains entered sporulation with similar efficiency. Short and long (over-exposed) exposures of the anti-SpoIIIA (A) and anti-SpoIIIAE (E) immunoblots are shown. Time (in hours) after the initiation of sporulation is shown. Sporulation efficiencies of the same strains are shown below the immunoblots. doi:10.1371/journal.pgen.1000566.g006](image-url)
prematurely, prior to the completion of engulfment. However, despite premature synthesis, the timing of σG activity is indistinguishable from wild-type and is detectable only upon the completion of engulfment [2,9,21,33,34]. Thus, these data suggest that σG is held inactive until a signal from the mother cell and/or the completion of engulfment triggers relief of inhibition.

The compelling nature of the feeding-tube model prompted us to re-visit this important experiment that challenges its simplicity. Examination of σG protein levels in a sporulation time course using one of the original PspoIIQ-sigG fusions [34] revealed premature synthesis of σG as reported previously (Figure 7B). However, the levels of σG protein present prior to the completion

![Image of Figure 7](image)

**Figure 7.** σG is active when synthesized prior to the completion of engulfment. (A) σG activity was assessed by microscopy using a fluorescence reporter (PsspE-gfp; false-colored green in the lower panel) in a ΔsigG mutant (ΔsigG, BTD3004), a wild-type background (wt, BTD3002), a ΔsigG mutant containing one copy of PspoIIQ-sigG (PΔsigG, BTD3007), or three copies of PspoIIQ-sigG (3X PΔsigG, BCM791), and a ΔsigG, ΔspolIAE double mutant that contains three copies of PspoIIQ-sigG (ΔE, BCM816). Sporulating cells were monitored at hour 2 of sporulation. The membranes from the same field were visualized using the dye TMA-DPH (false-colored red) and merged with the GFP signal. The membrane dye inefficiently traverses the lipid bilayer and therefore reports on the engulfment status of the forespore [54]. Forespores that stain weakly with TMA-DPH (white carets) have been completely engulfed by the mother cell. Forespores that have not yet completed engulfment have strong signal due to the two membranes surrounding the spore. Forespores that have σG activity but have not completed engulfment are indicated (yellow carets). Scale bar, 1 μm. Similar results were obtained with a PsspB-gfp reporter (not shown). (B) Immunoblot analysis of whole cell lysates from sporulating cells. The levels of σG were analyzed in sporulating cells from the same strains described in A. σF levels were monitored to control for efficiency of sporulation. σA levels were monitored to control for loading.

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of engulfment (hour 1.5–2) were quite low. In fact, at hour 2, the level of σ^G in wild-type was slightly higher than the strain carrying the P_{spoIIQ-sigG} fusion (Figure 7B). Although the P_{spoIIQ-sigG} directed accumulation of σ^G may vary with the exact construct used or the medium employed to induce sporulation [33,34], these results suggest that σ^G levels in the P_{spoIIQ-sigG} fusion strain might have been too low for the transcriptional reporter to reveal early activity. Thus, these data raised the possibility that σ^G could be active prior to the completion of engulfment and not subject to inhibition.

To investigate this idea, we increased the levels of premature σ^G by introducing three copies of the P_{spoIIQ-sigG} fusion at three non-essential loci in the genome (see Material and Methods). Under these conditions, the levels of σ^G were approximately 2-fold higher at early time points (hour 1.5–1.75) of sporulation (Figure 7B). Strikingly, in this strain, we could easily detect premature σ^G activity. Specifically, a σ^G-responsive promoter (P_{gfp}) fused to gfp revealed fluorescence in the majority of cells that had not yet completed engulfment (Figure 7A). Importantly, early σ^G activity was independent of SpoIIQ; SpoIIIA and mutants blocked in engulfment (Figure 7A and Figure S7A, see yellow carets). Moreover, consistent with the idea that the SpoIIIA-SpoIIQ complex is required to maintain metabolic potential in the forespore, at late time points (T3.5), the intensity of GFP fluorescence in the SpoIIIA and SpoIIQ mutants was lower than in the matched wild-type control (Figure S7B). Thus, in the absence of SpoIIIA or SpoIIQ, we could detect early σ^G activity but at late stages of sporulation, σ^G-dependent gene expression was reduced, suggesting that it may have ceased. These results challenge the idea that there is an active mechanism holding σ^G inactive that is relieved upon the completion of engulfment (and/or by a signal received from the mother cell). Furthermore, they provide support for the idea that the role of SpoIIIA-SpoIIQ complex in σ^G activity is to maintain metabolic potential in the forespore.

Discussion

We have presented evidence that the SpoIIIA proteins in the mother cell and SpoIIQ in the forespore form a large multimeric complex that spans the two membranes that surround the forespore (Figure 8A). In addition, we have shown that the ATPase motifs in SpoIIIAA are required for SpoIIIAA function in vivo. Based on these findings and the recent work of Camp and Losick [21,22] and Moran and colleagues [35], we propose that the SpoIIA-SpoIIQ complex forms a novel secretion apparatus that links the mother cell and forespore. This putative secretion complex appears to have been cobbled together from components that are found in a variety of specialized secretion systems found in Gram-negative bacteria. SpoIIIAA resembles ATPases found in type II and type IV secretion systems. SpoIIAB shares weak similarity with the GspF and TadB/C proteins found in type II and type IV secretion systems, respectively [36,37]. SpoIIAH shares homology with the YscJ/FliF protein family found in type III secretion systems [21,22]. SpoIIIAG also shares weak similarity (albeit weaker than SpoIIAH) to the YscJ/FliF protein family [22]. SpoIIIAE shares weak similarity with FlhB (YscU) found in type III secretion systems [22]. Finally, SpoIIIAE is similar to ABC-type permeases involved in Type I secretion.

Further support for the idea that these proteins form a secretion apparatus comes from recent findings that the extracellular domains of SpoIIAH and SpoIIQ, that reside in the space between the mother cell and forespore membranes, are accessible to modification by a soluble biotin ligase produced in the forespore [22]. These results suggest that SpoIIQ forms a pore in the forespore membrane that allows the ligase access to the extracellular domains of SpoIIAH and SpoIIQ. One unresolved issue concerning this putative secretion complex is the role of SpoIIAH. SpoIIAH appears to lie at the heart of the complex linking the SpoIIA proteins in the mother cell to SpoIIQ in the forespore. Paradoxically, cells lacking SpoIIAH have a sporula-

Figure 8. The SpoIIA proteins in the mother cell and SpoIIQ in the forespore form a secretion complex. (A) Schematic diagram showing the SpoIIA-SpoIIQ secretion complex in the two membranes that surround the forespore. SpoIIAA (A), SpoIIAB (B), SpoIIAC (C), SpoIIAD (D), SpoIIAE (E), SpoIIAF (F), SpoIIAG (G), SpoIIAH (H), and SpoIIQ (Q) are shown. SpoIIAA (A) is shown as a hexamer by analogy to other traffic ATPases [25]. SpoIIQ (Q) is shown as a multimeric pore based on the experiments of Meisner et al [22]. The actual stoichiometry of proteins in the complex is unknown. (B) The feeding-tube model. In wild-type cells (wt), the SpoIIIA-SpoIIQ complex secretes an unknown metabolite/osmolyte (red circle) into the forespore that maintains forespore integrity and σ^G activity (indicated by a green forespore). In the absence of the SpoIIIA (ΔIIIA) or SpoIIQ proteins, the forespore loses metabolic potential; the forespore collapses and σ^G activity cannot be maintained.

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tion efficiency of 5% while all other mutants in the complex are 1,000-fold worse (Table S1). Apparently, in the absence of SpoIIAH, the other SpoIIA proteins can assemble a partially functional complex with SpoIQQ. One possibility is that SpoIIAH is principally involved in tethering the other SpoIIA proteins to SpoIQ and that in its absence they can still interact, albeit weakly, with the forespore membrane protein. Alternatively, the SpoIIAG protein, which, like SpoIIAH, shares weak homology with the Ysc/JEff protein family [35], might function in place of SpoIIAH. Either model is consistent with our findings that CFP-SpoIIIAG remains somewhat enriched in the membranes that surround the forespore in the absence of SpoIIAH and that this enrichment is lost in the SpoIQ mutant (Figure 3).

Our data also indicate that the SpoIIA proteins and SpoIQQ are required to maintain forespore integrity. In the absence of these proteins, the forespore appears normal at early stages of sporulation. However, at a time when engulfment is complete, the forespore develops large invaginations and appears to shrink and/or collapse. The collapse of the forespore membranes likely explains the instability of the forespore [38,39] and the loss of compartmentalized gene expression previously observed at late times in development in SpoIIA mutants [33,40]. We hypothesize that these phenotypes are due to lack of osmolytes and/or loss of metabolic potential. Indeed, it has been shown previously that mutations that impair glycerol metabolism in B. subtilis result in membrane collapse [41] and that carbon starvation leads to cell autolysis [42]. Moreover, defects in osmotolerance in Listeria monocytogenes affects cell morphology [43]. Finally, our data suggest that the role of SpoIIA in maintaining forespore integrity is enzymatic rather than structural. Collectively, these results support and extend the feeding-tube model proposed by Camp and Losick [21,23], in which the SpoIIA-SpoIQQ secretion apparatus secretes nutrients and/or osmolytes into the forespore, "feeding" the spore in the final stages of preparation for dormancy (Figure 8B).

In this model, the maintenance of metabolic potential in the forespore is what enables σG activity. Our data showing that there is not an active mechanism holding σG inactive that is relieved upon the completion of engulfment provides further support for this new model. It is noteworthy that Stragier and colleagues have recently proposed that σG activity is held inactive until engulfment is complete by a protein (Gin/CsB) synthesized in the forespore under the control of σG [34]. Although this model has been challenged [21,44], our data do not rule out the possibility that Gin could serve as a timing device to help prevent early σG activity as was originally proposed by Piggot [44]. For example, Gin could be a saturable inhibitor of σG. Once σG levels (synthesized under the control of σG) exceed this negative regulator and provided the forespore has metabolic potential, σG could flip its auto-regulatory switch. In support of this idea, the level of premature σG activity was ~2-fold higher in the strain harboring three copies of PspoIQQsigG when Gin was absent (Figure S8).

One outstanding question in the regulation of σG is how its activity is coupled to proper engulfment. Our data showing that premature synthesis of σG results in early σG activity even when engulfment is impaired suggests that a separate regulatory pathway that links σG activity and engulfment is unlikely. Instead, we favor the model that SpoIIA and SpoIQQ proteins fail to assemble into an active complex when engulfment is perturbed and this prevents σG activity. In this model, the assembly of an active complex serves as the "surveillance mechanism" that couples gene expression to proper engulfment [17]. Interestingly, we find that in engulfment mutants, a subset (11%) of the sporulating cells succeed in activating σG to levels comparable to wild-type (Figure S9). We interpret this class of cells as those that have successfully assembled an active SpoIIA-SpoIQQ secretion complex and therefore can maintain forespore physiology and σG activity despite the failure to engulf. Consistent with this idea, the subset of cells with σG activity was completely eliminated in a SpoIIAD224A ATPase mutant or in the absence of SpoIQQ or the SpoIIA proteins (Figure S9 and data not shown).

A secretion apparatus that allows the mother cell to feed the forespore could have arisen because the forespore becomes fully engulfed inside the mother cell and therefore isolated from the external environment. In addition, the forespore developmental program might involve the down-regulation (or shut-off) of genes required for certain metabolic functions [45]. Under both conditions, the forespore would become dependent on the mother cell for nutrients and/or osmolytes. The observation that in engulfment mutants the forespore remains in contact with the environment yet most forespores lack σG activity [8,13,14] (Figure S9) suggests that the forespore has lost the ability to provide for itself and has become dependent on the mother cell for its final stages of maturation.

In conclusion, our results and those of Camp and Losick are most consistent with a model in which the SpoIIA-SpoIQQ complex functions as a novel secretion complex allowing the mother cell to nurture the forespore at late stages in sporulation (Figure 8B). In this model, σG activity depends on this complex not for an activating signal but to maintain forespore physiology. Our data suggest that the principal regulation of σG activity is i) the delay in sigG transcription under the control of σF; ii) the saturable inhibitor Gin that prevents pre-mature σG activity; iii) the auto-activation of the sigG gene; and iv) the maintenance of metabolic potential by the SpoIIA-SpoIQQ complex. Moreover, our results provide a plausible explanation for the coupling of σG activity to proper engulfment. Finally, the feeding-tube model nicely accounts for why no signaling protein required for σG activation has ever been identified and why no mutant has been found that can bypass the requirement for SpoIQQ or the SpoIIA proteins in compartmentalized σG activity. The challenge for the future is to identify the factor or factors secreted by the SpoIIA-SpoIQQ complex that nurture the forespore in its final stages of development.

Materials and Methods

General methods

All B. subtilis strains were derived from the prototrophic strain PY79 [46]. Sporulation was induced by resuspension at 37°C according to the method of Sterlini-Mandelstam [47] or by exhaustation (in supplemented DS medium; [48]). Sporulation efficiency was determined in 36-h cultures as the total number of heat-resistant (80°C for 20 min) colony forming units (CFUs) compared with wild-type heat-resistant CFUs. B. subtilis strains harboring the PspoIIAsigG fusions were constructed by direct transformation of ligation products because sigG is toxic to E. coli. Whole cell lysates from sporulating cells (induced by resuspension) were prepared as described previously [49]. Samples were heated for 10 min at 50°C prior to loading. Equivalent loading was based on OD600 at the time of harvest. Tables and of strains, plasmids and oligonucleotide primers and descriptions of plasmid construction can be found online as supplementary data (Table S2, Table S3, Table S4; Text S1).

Protein purification and antibody production

His6-SpoIIA, His6-SpoIIAF and His6-SpoIIIAG fusion proteins were expressed in E. coli BL21 DE3 pLysS and purified on Ni²⁺-NTA agarose (Qiagen). GST-SpoIIIAE fusion protein
was expressed in *E. coli* NB42 and purified on glutathione-agarose (GE Life Sciences). A complete description of the purifications can be found in Text S1. Peak fractions were pooled and used to generate rabbit polyclonal antibodies (Covance). Crude serum was affinity-purified as described [50].

**Antibody resins preparation**

All antibody resins were prepared as described previously [30]. Briefly, affinity-purified antibodies (2 mg) were batched absorbed to 1 ml protein A-agarose (Amersham). The antibody resin was washed with phosphate buffered saline (PBS) and covalently cross-linked to the protein A-agarose by the addition of Disuccinimidyl Suberate (Pierce) to a final concentration of 3 mM. After 30 minutes the reaction was quenched by the addition of Tris pH 7.5 to a final concentration of 100 mM. The antibody resin was washed with 100 mM Glycine pH 2.5 to remove uncross-linked antibody and then neutralized with 1× PBS.

**Co-immunoprecipitation from detergent solubilized membrane fractions**

Preparation of crude membranes and detergent solubilization of membrane proteins was performed as described [30]. 50 ml cultures were harvested at hour 2.5 of sporulation (by resuspension) and washed twice with 1× SMM (0.5 M Sucrose, 20 mM MgCl$_2$, 20 mM Maleic acid pH 6.5) at room temperature. Cells were resuspended in 1/10 volume 1× SMM and treated with Lysozyme (0.5 mg/ml, 1 hour). Soluble and insoluble fractions were separated by centrifugation and flash frozen in N2(l). Thawed protoplasts were disrupted by osmotic lysis with 3 ml hypotonic buffer (Buffer H) (20 mM Hepes pH 8, 200 mM NaCl, 1 mM DTT, with protease inhibitors: 1 mM PMSF, 0.5 μg/ml leupeptin, 0.7 μg/ml pepstatin). MgCl$_2$ and CaCl$_2$ were added to 1 mM and lysates were treated with DNaseI (10 μg/ml) (Worthington) and RNaseA (20 μg/ml) (USB) for 30 min on ice. The membrane fraction was separated by centrifugation at 100,000×g for 1 hour at 4°C. The supernatant was carefully removed and the membrane pellet was dispersed in 200 μl Buffer G (Buffer H with 10% Glycerol). Crude membranes were aliquoted and flash frozen in N2(l).

Crude membranes were diluted 5-fold with Buffer S [Buffer H with 20% Glycerol and 100 μg/ml Lysozyme] and membrane proteins were solubilized by the addition of the nonionic detergent DDM (n-dodecyl-β-d-maltopyranoside, Sigma) to a final concentration of 0.5%. The mixture was rotated at 4°C for 1 hour. Soluble and insoluble fractions were separated by centrifugation at 100,000×g for 1 hour at 4°C. The soluble fraction (the load) was mixed with 40 μl antibody resin and rotated for 4 h at 4°C. The resin was pelleted at 5 K rpm and the supernatant (the flow through) was removed. The resin was washed 4 times with 1 ml Buffer S containing 0.01% DDM. Immunoprecipitated proteins were eluted by the addition of 90 μl of Sodium Dodecyl Sulfate (SDS) sample buffer and heated for 15 minutes at 50°C. The eluted material (the IP) was transferred to a fresh tube and 2-Mercaptoethanol was added to a final concentration of 10%. The load, flow through and IP were analyzed by immunoblot.

**Mass spectrometry analysis**

Immunoprecipitates were trypsinized and the peptides were separated on a nanoscale C18 reverse-phase high-pressure liquid chromatography capillary column, and were subjected to electrospray ionization followed by MS using an LCQ DECA ion-trap mass spectrometer.

**Immunoblot analysis**

Proteins were separated by SDS-PAGE on 15% polyacrylamide gels, electroblotted onto Immobilon-P membranes (Millipore) and blocked in 5% nonfat milk in phosphate-buffered saline (PBS) -0.5% Tween-20. The blocked membranes were probed with affinity-purified anti-SpoIIAA (1:10,000), anti-SpoIIAE (1:5,000), anti-SpoIIAF (1:10,000), anti-SpoIIAG (1:10,000), anti-spoIIAH (1:10,000) [30], anti-SpoIID (1:10,000) [49], anti-SpoIIQ (1:10,000) [49], anti-GFP (1:10,000) [32], anti-myc (1:1000) (Covance), anti-σ$^A$ (1:20,000) (a gift from M. Ho and R. Losick), anti-σ$^F$ (1:5,000) [51], and anti-σ$^G$ (1:10,000) [32]. Two primary antibodies were diluted into 3% BSA in 1× PBS-0.05% Tween-20. Primary antibodies were detected using horseradish peroxidase-conjugated goat, anti-rabbit G (BioRad) and the Western Lightning reagent kit as described by the manufacturer (PerkinElmer).

**Fluorescence microscopy**

Fluorescence microscopy was performed with an Olympus BX61 microscope as previously described [32]. Fluorescent signals were visualized with a phase contrast objective UplanFl 100×. Samples were captured with a monochrome CoolSnapHQ digital camera (Photometrics) using Metamorph software version 6.1 (Universal Imaging). Exposure times were typically 500 ms for GFP and CFP. The membrane dye TMA-DPH (Molecular Probes) was used at a final concentration of 0.01 mM and exposure times were typically 200 ms. Images were analysed, adjusted and cropped using Metamorph software. The cells were centrifuged (8 K rpm for 30 sec) prior to visualization. This step had no impact on the reported phenotypes.

**Electron microscopy**

Samples were collected at hour 3 of sporulation (by resuspension) and prepared for electron microscopy. Cells were resuspended in PBS and fixed (0.5% paraformaldehyde and 2.5% glutaraldehyde) for 4 h at room temperature followed by dehydration with acetone and embedding in Epon resin. Finally, ultrathin sections were stained with lead citrate. For evaluation of forespore morphology, only longitudinal sections were considered. Greater than 50 sporangia were examined per strain.

**Supporting Information**

**Figure S1** Sequence alignment of the C-terminal domain of SpoIIAA (IIAA) from *B. subtilis* with the ATPase domains of several secretion ATPases. The conserved motifs (red boxes) found in all secretion NTPases are highlighted. Mutated residues are indicated (blue asterisk). The alignment was made using ClustalW (http://www.ch.embnet.org/software/ClustalW.html) and BOX-SHADE (http://www.ch.embnet.org/software/BOX_form.html). Found at: doi:10.1371/journal.pgen.1000566.s001 (0.96 MB TIF)

**Figure S2** The ATPase motifs in SpoIIAA are required for σ$^G$ activity and sporulation efficiency. (A) σ$^G$ activity was assessed in single cells by microscopy using a fluorescent reporter (PspoE-β-gal) in a wild-type background (wt, BTD1609), a δspoIIHLH mutant (δA, BTD2713), a δspoIIHLH mutant containing a wild-type copy of spoIIHLH inserted at a non-essential locus (A(wt), BTD2719), a spoIIHLH$^{K149A}$ Walker B box point mutant (A(K149A), BTD2775), a spoIIHLH$^{D224A}$ Walker A box point mutant (A(D224A), BTD2906), a spoIIHLH$^{K140A}$ Walker A box point mutant (A(K140A), BTD2906), a spoIIHLH$^{K149A}$ Walker A box point mutant (A(K149A), BTD2906), a spoIIHLH$^{K140A}$ Walker A box point mutant (A(K140A), BTD2906), and a spoIIHLH$^{D224A}$ Walker A box point mutant (A(D224A), BTD2906). Cells were visualized at hour 3 of sporulation. Forespore CFP fluorescence (false-colored green in the lower panel) and the
fluorescent membrane dye TMA-DPH (false-colored red) are shown. Scale bar, 1 μm. Sporulation efficiencies of the same strains are shown below the fluorescent images. (B) Immunoblot analysis of whole cell lysates from sporulating cells shown in A. Time (in hours) after the initiation of sporulation is indicated.

Figure S3 The SpoIIIAE, SpoIIIAF and SpoIIIAJ proteins reside in a membrane complex. Immunoprecipitations were performed on detergent-solubilized membrane fractions derived from B. subtilis sporulating cells at hour 2.5 of sporulation. (A) Immunoprecipitates using anti-SpoIIIAE antibody resin from a spoIIIAE+ strain (wt, BDR94) and a δspoIIIAE mutant (δE, BDT2535) are shown. (B) Immunoprecipitates using anti-SpoIIIAF antibody resin from a spoIIIAF+ strain (wt, BDR94) and a δspoIIIAF mutant (δF, BDT2537) are shown. The detergent-solubilized membrane fraction prior to immunoprecipitation (Load), the supernatants after immunoprecipitation (Sup), and the immunoprecipitates (IP) were subjected to immunoblot analysis using anti-SpoIIIAE (E), anti-SpoIIIAF (F), and anti-SpoIIIAJ (G) antibodies. All four strains contained a δspoIVB mutation to prevent cleavage of proteins that have domains that reside in the intermembrane space (K. Marquis, N. Campo, TD, and DZR, unpublished observations).

Figure S4 SpoIIIAH resides in a membrane complex with SpoIIAH and SpoIIQ. Immunoprecipitations were performed with cleared lysates from sporulating B. subtilis. Cells were collected at hour 2.5 and treated with lysozyme. Proteoplasts were lysed in a hypotonic buffer (50 mM Tris pH 7.5, 1 mM EDTA). Lysates were centrifuged (at 10,000 ×g for 5 minutes) to remove cellular debris. Cleared lysates were subjected to immunoprecipitation with anti-FLAG M2-agarose (σ). (A) Immunoprecipitations with cells expressing FLAG-tagged SpoIIAH. Lysates were incubated in the presence or absence of 1% Triton X-100. Cleared lysates (Load), the supernatants after immunoprecipitation (Sup), and the immunoprecipitates (IP) were subjected to immunoblot analysis using anti-FLAG to detect SpoIIAH (H), anti-SpoIIIAJ (G), anti-SpoIIQ (IQ), and anti-SpoIID (IID) antibodies. (B) Immunoprecipitation with cells expressing FLAG-tagged FtsH. Cleared lysates (Load), the supernatants after immunoprecipitation (Sup), and the immunoprecipitates (IP) were subjected to immunoblot analysis using anti-FLAG to detect FtsH (Fsh) and anti-SpoIIIAJ (G) antibodies.

Figure S5 Morphological defects in the absence of SpoIIIA proteins. Forespore morphology was monitored by fluorescence microscopy at hour 3 of sporulation in a wild-type background (wt, BCM703), a δspoIIIBAB mutant (δB, BCM704), a δspoIIIC mutant (δE, BTD3062), and a δspoIIID mutant containing low levels of SpoIIAE (low E, BTD3063), and a δspoIIIAE mutant containing high levels of SpoIIAE (high E, BTD3064). All strains contained a forespore reporter (PspoIIQ-cfp; false-colored blue in the lower panel) to visualize the forespore cytoplasm and a YFP-SpoIIFA fusion (IYFA; false-colored green in the lower panel) that labels the mother cell membranes that surround the forespore. The membranes (mb) from the same field were visualized with the fluorescent dye TMA-DPH (false-colored red in the lower panel). Scale bar, 1 μm.

Figure S7 σG is active when synthesized prior to the completion of engulfment. (A) Larger fields of cells showing premature synthesis of σG result in early σG activity. σG activity was assessed by microscopy using a fluorescent reporter (PspIIQ-cfp) in a δsigG mutant (δsigG, BTD3004), a wild-type background (wt, BTD3002), a δsigG mutant containing one copy of PspIIQ-sigG (PIIQ-sigG, BTD3007), three copies of PspIIQ-sigG (3× PIIQ-sigG, BCM791), a δsigG, δspoIIIAE double mutant that contains three copies of PspIIQ-sigG (δE, BCM816), and a δsigG, δspoIIQ double mutant that contains three copies of PspIIQ-sigG (δE, BCM814). Sporulating cells were monitored at hour 2 of sporulation. The membranes from the same field were visualized using the dye TMA-DPH (false-colored red) and merged with the GFP signal (false-colored green). (B) Late σG activity requires SpoIIIA and SpoIIQ proteins. σG activity was quantified at hour 3.5 of sporulation from the same strains as above. The total fluorescence intensity of GFP was measured in each forespore from one field (>400 forespores per strain). Background fluorescence from the same measured region was subtracted. The histogram shows the distribution of GFP intensity in BCM791, BCM814, BCM816.

Figure S8 Early σG activity is higher and more prevalent in the absence of CsbB/Gin. σG activity was assessed in single cells by microscopy using a fluorescent reporter (PspE-cfp) in a wild-type background (wt, BTD3002), in a δsigG mutant (δsigG, BTD3095), a δsigG mutant containing three copies of PspIIQ-sigG (3× PIIQ-sigG, BTD3100), and the same strain lacking CsbB (3× PIIQ-sigG, Agin, BTD3102). Sporulating cells were monitored at hour 2 of sporulation. The membranes from the same field were visualized using the dye TMA-DPH (false-colored red) and merged with the GFP signal (false-colored green). The membrane dye inefficiently traverses the lipid bilayer and therefore reports on the engulfment status of the forespore [1]. Forespores that stain weakly with TMA-DPH have been completely engulfed by the mother cell. Forespores that have not yet completed engulfment have strong signal due to the two membranes surrounding the spore. Yellow carets highlight examples of forespore that have σG activity but have not completed engulfment. The fluorescence intensities of the GFP reporter in the σG strain are ~2-fold higher than the intensities in the matched control strain. Scale bar, 1 μm.

Figure S9 A subset of cells that are blocked for engulfment have σG activity. (A) σG activity was assessed in single cells by microscopy using a fluorescent reporter (PspE-cfp) in a wild-type background (wt, BTD3002), a δspoID mutant (δIID, BTD3085), and a δspoID, spoIIIAA/D224A double mutant (δIID, A(D224A), BTD3086). Cells were visualized at hour 3 of sporulation. Forespore GFP fluorescence (false-colored green in the lower panel) and the fluorescent membrane dye TMA-DPH (false-colored red in the lower panel) are shown. (B) Large fields of
sporulating cells from the same three strains showing forespore CFP fluorescence. The percentage of sporulating cells that have σG activity is shown below the field.

### Table S1 Sporulation efficiency of the SpoIII A mutants.

<table>
<thead>
<tr>
<th>Strain</th>
<th>Sporulation Efficiency</th>
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<td>AOH</td>
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### Table S2 Strains used in this study.

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<tr>
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<td>ΔspoIIIA</td>
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<tr>
<td>MS H</td>
<td>ΔspoIIIAΔH</td>
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### Table S3 Plasmids used in this study.

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<td>pPS3</td>
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### Table S4 Oligonucleotide primers used in this study.

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<td>spoIIIB</td>
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### Text S1 Supplemental materials and methods.

#### References