High-Throughput Genetic Screens Identify a Large and Diverse Collection of New Sporulation Genes in Bacillus subtilis

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High-Throughput Genetic Screens Identify a Large and Diverse Collection of New Sporulation Genes in *Bacillus subtilis*

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Abstract

The differentiation of the bacterium *Bacillus subtilis* into a dormant spore is among the most well-characterized developmental pathways in biology. Classical genetic screens performed over the past half century identified scores of factors involved in every step of this morphological process. More recently, transcriptional profiling uncovered additional sporulation-induced genes required for successful spore development. Here, we used transposon-sequencing (Tn-seq) to assess whether there were any sporulation genes left to be discovered. Our screen identified 133 out of the 148 genes with known sporulation defects. Surprisingly, we discovered 24 additional genes that had not been previously implicated in spore formation. To investigate their functions, we used fluorescence microscopy to survey early, middle, and late stages of differentiation of null mutants from the *B. subtilis* ordered knockout collection. This analysis identified mutants that are delayed in the initiation of sporulation, defective in membrane remodeling, and impaired in spore maturation. Several mutants had novel sporulation phenotypes. We performed in-depth characterization of two new factors that participate in cell–cell signaling pathways during sporulation. One (SpoIIT) functions in the activation of σE in the mother cell; the other (SpoIIIL) is required for σG activity in the forespore. Our analysis also revealed that as many as 36 sporulation-induced genes with no previously reported mutant phenotypes are required for timely spore maturation. Finally, we discovered a large set of transposon insertions that trigger premature initiation of sporulation. Our results highlight the power of Tn-seq for the discovery of new genes and novel pathways in sporulation and, combined with the recently completed null mutant collection, open the door for similar screens in other, less well-characterized processes.

Author Summary

When starved of nutrients, the bacterium *Bacillus subtilis* differentiates into a dormant spore that is impervious to environmental insults. Decades of research have uncovered...
over 100 genes required for spore formation. Molecular dissection of these genes has revealed factors that act at every stage of this developmental process. In this study, we used a high-throughput genetic screening method called transposon sequencing to assess whether there were any sporulation genes left to be discovered. This approach identified virtually all of the known sporulation genes, as well as 24 new ones. Furthermore, transposon sequencing enabled the discovery of two new sets of mutants in which the sporulation process was either delayed or accelerated. Using fluorescence microscopy, we determined the developmental stage at which each mutant was impaired and discovered mutants that are delayed in initiation of sporulation, or defective in morphogenesis, cell–cell signaling, or spore maturation. Our findings exemplify the utility of transposon sequencing to uncover new biology in well-studied processes, suggesting that it could similarly be used to identify novel genes required for other aspects of bacterial physiology, such as natural competence, stationary phase survival, or the responses to cell envelope stress and DNA damage.

Introduction

The morphological process of spore formation in Bacillus subtilis has been studied for over 50 years and constitutes one of the most well-characterized developmental pathways in biology [1–4]. Its molecular dissection has contributed to our understanding of diverse biological processes, including cell fate determination, signal transduction, membrane and cell wall remodelling, subcellular protein localization, and chromosome dynamics [5–8].

Underlying this seemingly simple process is a set of highly orchestrated morphological events that are both driven by and coupled to developmental programs of gene expression. Upon nutrient limitation and in response to population density, B. subtilis enters the sporulation pathway. The first landmark event in this process is an asymmetric division, generating a large cell (called the mother cell) and a smaller cell (the prospective spore or forespore). Shortly after polar division, the mother cell membranes migrate around the forespore in a phagocytic-like process called engulfment, generating a cell within a cell. Membrane fission at the cell pole releases the forespore into the mother cell cytoplasm. The mother then assembles a set of protective layers around the forespore, while the spore prepares for dormancy. When the spore is fully mature, the mother cell lyases, releasing it into the environment.

This differentiation process takes 5–7 h to complete and is controlled by a series of stage- and compartment-specific transcription factors. Although the mother and forespore follow distinct developmental programs of gene expression, they remain linked to each other through cell–cell signalling pathways. Entry into sporulation is governed by two transcriptional regulators: the response regulator Spo0A and the stationary phase sigma factor σ^H. The proteins produced under their control prevent new rounds of DNA replication, remodel the replicated chromosomes, and shift the cell division site from a medial to a polar position. These regulators are also responsible for the expression of compartment-specific transcription factors that are held inactive until polar division is complete. Asymmetric division triggers the activation of the first forespore-specific sigma factor σ^F. σ^F controls the expression of a collection of forespore proteins, including a secreted signalling molecule that activates the first mother cell transcription factor σ^E. The proteins produced under σ^E control prevent a second division at the opposite cell pole and drive the migration of the mother cell membranes around the forespore. During this process, a transenvelope complex is assembled in the membranes that separate the mother cell and forespore. This complex, which resembles a specialized secretion system,
allows the mother to maintain forespore physiology and developmental gene expression under the control of the late-acting transcription factor σ^G. After the completion of engulfment, σ^G produces two signalling proteases that are secreted into the space between the mother and forespore membranes, where they trigger the activation of the mother cell transcription factor σ^K. A large set of proteins produced under the control of σ^E and σ^K in the mother cell are responsible for assembling a thick layer of peptidoglycan between mother cell and forespore membranes and a multi-layered proteinaceous coat on the spore exterior. σ^G in the forespore controls the expression of proteins that prepare the spore for dormancy, including DNA binding proteins that compact and protect the spore chromosome and a membrane complex that imports dipicolinic acid, a compound that facilitates spore dehydration. Finally, a set of cell wall degrading enzymes produced in the mother cell under σ^K control trigger mother cell lysis and release of the dormant spore.

Five decades of research have led to the identification and characterization of over 100 genes that are required for successful completion of this developmental process. A pigmented protein produced in the mother cell and assembled around the developing spore gives rise to opaque beige colonies on sporulation agar plates [9]. Many sporulation mutants were originally identified in classical genetic screens based on their failure to produce this pigment [10–12]. Some of these genes are absolutely required for spore formation, while most play important but nonessential roles with mutant sporulation phenotypes ranging over eight orders of magnitude [13]. Advances in the identification and characterization of sporulation genes have been punctuated by the introduction of new technologies. These include the use of the Tn917 transpose (and lacZ-containing variants) to map sporulation mutants and identify sporulation-induced genes [14,15], cytological approaches to localize sporulation proteins and monitor spore morphogenesis [16–18], and transcriptional profiling to define entire sporulation-specific regulons [19–25]. More recently, taking advantage of the groundswell of sequenced bacterial genomes, novel sporulation genes have been unearthed through phylogenetic profiling [26,27].

Here, we screened for new sporulation genes using a transposon-sequencing (Tn-seq) approach [28]. Our screen identified virtually all of the known sporulation genes and 24 new ones that we have characterized by taking advantage of the ordered B. subtilis null mutant collection. These new genes play roles in several stages of development, including initiation, morphogenesis, cell–cell signalling, and spore maturation. Furthermore, by repeating the Tn-seq screen with cells undergoing sporulation for 5 h instead of 24, we discovered that a large set of uncharacterized sporulation-induced genes are required for timely development. Finally, this second screen also identified a novel set of transposon insertions that induce premature initiation of sporulation, giving the appearance of accelerated spore maturation. Collectively, this approach identified new sporulation genes, novel phenotypes, and new pathways, and underscores the utility of Tn-seq to uncover new biology in well-studied processes.

**Results**

**A Screen for Mutants Defective in Sporulation**

To investigate whether there were additional genes required for sporulation, we performed a high-throughput screen using transposon-sequencing (Tn-seq). We generated a Mariner transposon library containing over 138,000 unique insertions (in 63% of all possible TA dinucleotides) in the B. subtilis 168 genome (see Methods). On average, each gene was disrupted at 26 unique sites. The library was grown in sporulation medium and a sample was removed at the onset of starvation (T0). The remaining culture was allowed to exhaust its nutrients and sporulate over the next 24 h (T24). The culture was then incubated at 80°C for 20 min to kill
vegetative and sporulation-defective cells and the spores were plated on lysogeny broth (LB) agar plates. More than 750,000 colonies from germination-proficient spores were pooled and the transposon insertions at T0 and T24 were mapped by massively parallel DNA sequencing (see Methods). The location and number of reads for each transposon insertion were compared at the onset of starvation and after heat-kill, germination, and outgrowth (for simplicity, referred to as sporulation) (Fig 1 and S1 Fig). We set an initial statistical significance cutoff value of $p < 0.05$ with a 2-fold reduction in transposon insertions at T24 compared to T0 (Fig 1B and S1 Data). More than 350 genes met these initial criteria. Visual inspection of the transposon insertion profiles revealed that ~20% of these genes had very few insertions at T0. An additional ~20% had a modest (2–3-fold) reduction in transposon insertions at T24, but insertions were present throughout the open reading frame. Most of these genes were not considered further (Fig 1 and S1 Fig).

Of the 111 well-characterized sporulation genes, our screen identified 98 (Fig 1 and S1 Fig and S1 Table). Among these were all the genes with strong sporulation mutant phenotypes, including sigE, spoIIGA, and sigG (Fig 1), spoIIIR, spoIIIC, and spoVB (S1 Fig). Moreover, we also identified genes like kinA, spoII, and bofA for which the sporulation efficiency of the mutants ranges from 1%–20% (S1 Fig and S1 Table) [29–31]. The 13 genes that were not identified in our screen had very mild or no sporulation/germination phenotypes under our assay conditions (S1 Table) (see below). An additional 37 less well-characterized genes have been reported to be required for efficient sporulation, many of which were originally identified by transcriptional profiling [19–24, 26, 27]. Thirty-five of these genes were identified in our Tn-seq screen (Table 1). Thus, out of the 148 previously identified genes required for sporulation, our screen identified 133 of them. The validation and characterization of 24 additional genes that had not been implicated in spore formation will be described below.

**Tn-seq Identifies Mutants That Delay Sporulation**

Two well-characterized sporulation genes that our Tn-seq screen failed to identify were racA and fisB (Fig 2A). RacA acts early during sporulation to remodel the replicated chromosomes into an elongated structure called the axial filament and anchors the origins at opposite cell poles [32, 33]. Despite the pronounced cytological phenotype in cells lacking RacA, the mutant is reduced only ~2-fold in sporulation efficiency. FisB functions in membrane fission, the final stage in the phagocytic-like process of engulfment [34]. In sporulating cells lacking FisB, the mother cell membranes migrate around the forespore but are delayed and impaired in scission and release of the forespore into the mother cell cytoplasm. Again, despite its dramatic cytological phenotype, the FisB mutant is reduced only ~4-fold in sporulation efficiency. The mild phenotypes of these mutants likely account for the recovery of transposon insertions in these genes at T24 (Fig 2A). We wondered whether we would have identified racA and fisB and additional genes in a more stringent sporulation screen that demanded timely differentiation into heat-resistant spores. Specifically, we hypothesized that transposon insertions in these genes delay development. To test this, we first assayed sporulation efficiency in wild-type cells in a time course after the onset of starvation and found that by hour 5, between 0.01% and 0.4% of the sporulating cells had differentiated into heat-resistant spores. We then repeated the Tn-seq screen but harvested and heat-selected the transposon library of sporulating cells at hour 5 (T5).

As anticipated, transposon insertions in racA and fisB were significantly under-represented at this early time point compared to time 0 and hour 24 (Figs 1C and 2A). Along with these two genes and 156 of the 158 genes we identified at T24, there were 147 additional genes in which transposon insertions delayed the formation of heat-resistant spores (Fig 1C and S2 Table and S1 Data). Out of the 15 previously identified sporulation genes that we failed to
detect at T24, seven were hits at T5 (S2 Table). Interestingly, 36 of the genes identified at T5 were previously reported to be expressed under sporulation control but deletions were not found to have sporulation defects [20–25]. An additional 38 genes identified at T5 are
### Table 1. Independently validated sporulation genes identified by Tn-seq.

<table>
<thead>
<tr>
<th>section</th>
<th>gene</th>
<th>p-valuea</th>
<th>fold changeb</th>
<th>description</th>
<th>spo (%)c</th>
<th>cytological phenotype</th>
</tr>
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<tbody>
<tr>
<td>Newly identified sporulation genes</td>
<td>araR</td>
<td>1.24 × 10⁻¹²</td>
<td>24.2</td>
<td>repressor of arabinose utilization</td>
<td>28</td>
<td>no observable phenotype</td>
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<td></td>
<td>defB</td>
<td>2.53 × 10⁻⁷</td>
<td>14</td>
<td>N-formylcysteine deformylase</td>
<td>9</td>
<td>engulfment and fission defects/reduced σG activity</td>
</tr>
<tr>
<td></td>
<td>gapB</td>
<td>1.02 × 10⁻¹²</td>
<td>16.1</td>
<td>GAP dehydrogenase</td>
<td>0.3</td>
<td>delayed entry/thin cells/fewer sporulating cells</td>
</tr>
<tr>
<td></td>
<td>gidA</td>
<td>1.87 × 10⁻⁹</td>
<td>13.8</td>
<td>5′-tRNA-modifying enzyme</td>
<td>23.9</td>
<td>longer cells/some aberrant septa and forespores—similar to ΔtrmE</td>
</tr>
<tr>
<td></td>
<td>gtaB</td>
<td>4.96 × 10⁻⁸</td>
<td>123.8</td>
<td>UDP-glucose pyrophosphorylase</td>
<td>0.1</td>
<td>smaller and wider cells with small forespores—identical to ΔpgcA</td>
</tr>
<tr>
<td></td>
<td>iolR</td>
<td>7.01 × 10⁻⁵</td>
<td>3.2</td>
<td>repressor of myo-inositol catabolism</td>
<td>38</td>
<td>no observable phenotype</td>
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<tr>
<td></td>
<td>miaA</td>
<td>1.27 × 10⁻⁶</td>
<td>2.7</td>
<td>5′-tRNA-modifying enzyme</td>
<td>16</td>
<td>ND</td>
</tr>
<tr>
<td></td>
<td>minJ</td>
<td>2 × 10⁻⁵</td>
<td>3.5</td>
<td>division site selection</td>
<td>36</td>
<td>delayed entry/aberrant forespores</td>
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<tr>
<td></td>
<td>ndk</td>
<td>7.45 × 10⁻⁵</td>
<td>4.6</td>
<td>nucleotide diphosphate kinase</td>
<td>13.4</td>
<td>ND</td>
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<td></td>
<td>nrrA</td>
<td>3.04 × 10⁻⁶</td>
<td>6.7</td>
<td>nanoRNase</td>
<td>5</td>
<td>delayed entry/some small forespores with reduced σG activity</td>
</tr>
<tr>
<td></td>
<td>pgcA</td>
<td>5.17 × 10⁻¹⁵</td>
<td>28.9</td>
<td>UDP-glucose synthesis</td>
<td>0.1</td>
<td>smaller and wider cells with small forespores—identical to ΔgtaB</td>
</tr>
<tr>
<td></td>
<td>rasP</td>
<td>3.75 × 10⁻⁷</td>
<td>26.3</td>
<td>intramembrane protease</td>
<td>1.4</td>
<td>delayed entry/thin and oblong forespores</td>
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<td></td>
<td>resA</td>
<td>6.56 × 10⁻⁴</td>
<td>4.9</td>
<td>heme translocase</td>
<td>45.6</td>
<td>delayed entry</td>
</tr>
<tr>
<td></td>
<td>rex (ydiH)</td>
<td>3.42 × 10⁻⁷</td>
<td>26.7</td>
<td>transcriptional repressor</td>
<td>9.2</td>
<td>no observable phenotype</td>
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<tr>
<td></td>
<td>smpB</td>
<td>1.2 × 10⁻³</td>
<td>9.8</td>
<td>tmRNA-binding protein</td>
<td>23</td>
<td>delayed entry</td>
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<td></td>
<td>speE</td>
<td>5.16 × 10⁻⁶</td>
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<td>spermidine synthase</td>
<td>6</td>
<td>delayed entry</td>
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<td>spoIIIL (ywmB)</td>
<td>4.76 × 10⁻¹⁰</td>
<td>5.8</td>
<td>unknown</td>
<td>38</td>
<td>abortive disporic/impaired σE activation</td>
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<tr>
<td></td>
<td></td>
<td>2 × 10⁻³</td>
<td>15.9</td>
<td>unknown</td>
<td>14</td>
<td>small spores/reduced σG activity</td>
</tr>
<tr>
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<td>trmE</td>
<td>3.48 × 10⁻⁷</td>
<td>13.7</td>
<td>5′-tRNA-modifying enzyme</td>
<td>21.9</td>
<td>longer cells/some aberrant septa and forespores—similar to ΔgidA</td>
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<td></td>
<td>ubiE (menH)</td>
<td>2.41 × 10⁻⁷</td>
<td>180.8</td>
<td>menaquinone synthase</td>
<td>12.7</td>
<td>no observable phenotype</td>
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<tr>
<td></td>
<td>yaaD (pdxT)</td>
<td>8.73 × 10⁻⁵</td>
<td>11</td>
<td>pyridoxal 5′-phosphate synthesis</td>
<td>1.2</td>
<td>ND</td>
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<tr>
<td></td>
<td>yerC</td>
<td>1.15 × 10⁻⁴</td>
<td>359</td>
<td>putative DNA-binding protein</td>
<td>10</td>
<td>no observable phenotype</td>
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<tr>
<td></td>
<td>yqhT</td>
<td>2.78 × 10⁻¹⁰</td>
<td>22.1</td>
<td>Xaa-Pro peptidase</td>
<td>1.2</td>
<td>delayed entry</td>
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<tr>
<td></td>
<td>ytxG</td>
<td>5.8 × 10⁻⁵</td>
<td>193</td>
<td>general stress protein/sigB/sigH</td>
<td>39.6</td>
<td>aberrant membrane morphologies</td>
</tr>
</tbody>
</table>

Previously identified but less well characterized sporulation genes

|         | asnO     | 1.58 × 10⁻¹⁵ | 24.2 | asparagine synthase (σE regulon) | 0 | no observable phenotype |
|         | clpC     | 7.79 × 10⁻¹⁶ | 600.7| ATPase subunit of ClpCP          | 0.4| delayed entry/engulfment defect/sm small spores/reduced σG activity |

(Continued)
<table>
<thead>
<tr>
<th>section</th>
<th>gene</th>
<th>p-value&lt;sup&gt;a&lt;/sup&gt;</th>
<th>fold change&lt;sup&gt;b&lt;/sup&gt;</th>
<th>description</th>
<th>spo (%)&lt;sup&gt;c&lt;/sup&gt;</th>
<th>cytological phenotype</th>
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<tr>
<td></td>
<td>cotE</td>
<td>$3.23 \times 10^{-2}$</td>
<td>2.3</td>
<td>coat protein (σE and σK regulons)</td>
<td>49.1</td>
<td>ND</td>
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<tr>
<td></td>
<td>dacB</td>
<td>$6.54 \times 10^{-13}$</td>
<td>39.3</td>
<td>DD-carboxypeptidase</td>
<td>19.3</td>
<td>no observable phenotype</td>
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<tr>
<td></td>
<td>dgkA</td>
<td>$2.56 \times 10^{-5}$</td>
<td>170.3</td>
<td>undecaprenol kinase/cortex synthesis</td>
<td>33.7</td>
<td>ND</td>
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<td>divIVA</td>
<td>$3.21 \times 10^{-3}$</td>
<td>2.8</td>
<td>division site selection/σF activity</td>
<td>11.1</td>
<td>ND</td>
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<td>ecsA</td>
<td>$3.51 \times 10^{-9}$</td>
<td>81.5</td>
<td>ABC transporter family</td>
<td>0.1</td>
<td>ND</td>
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<td>ecsB</td>
<td>$6.79 \times 10^{-10}$</td>
<td>19.5</td>
<td>ABC transporter family</td>
<td>13.4</td>
<td>ND</td>
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<td></td>
<td>efp</td>
<td>$3 \times 10^{-7}$</td>
<td>200</td>
<td>elongation factor P</td>
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<td>ND</td>
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<td>ftsH</td>
<td>$2.14 \times 10^{-9}$</td>
<td>139.1</td>
<td>membrane protease</td>
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<tr>
<td></td>
<td>gerM</td>
<td>$2 \times 10^{-7}$</td>
<td>96.3</td>
<td>cortex synthesis and/or germination (σE regulon)</td>
<td>0.3</td>
<td>ND</td>
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<td></td>
<td>kbaA</td>
<td>$3.73 \times 10^{-2}$</td>
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<td>regulator of KinB</td>
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<td></td>
<td>lgt (gerF)</td>
<td>$1.68 \times 10^{-8}$</td>
<td>905.7</td>
<td>lipomodification/germination</td>
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<td>mntR</td>
<td>$2.16 \times 10^{-3}$</td>
<td>11.2</td>
<td>trxn regulator of Mn+ + transport</td>
<td>8.9</td>
<td>ND</td>
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<td></td>
<td>prpC</td>
<td>$4.24 \times 10^{-4}$</td>
<td>6</td>
<td>phosphatase</td>
<td>50.7</td>
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<td>putB</td>
<td>$6.2 \times 10^{-3}$</td>
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<td>proline dehydrogenase (spo0A and σE regulons)</td>
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<td>no observable phenotype</td>
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<td>rsfA</td>
<td>$4.12 \times 10^{-3}$</td>
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<td>regulator of σF controlled genes</td>
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<td>skfE</td>
<td>$1.42 \times 10^{-7}$</td>
<td>19.1</td>
<td>export of sporulation killing factor</td>
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<td>ND</td>
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<td>skfF</td>
<td>$4.1 \times 10^{-23}$</td>
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<td>export of sporulation killing factor</td>
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<td>spoVIE (pdaB)</td>
<td>$7.23 \times 10^{-6}$</td>
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<td>polysaccharide deacetylase (σE regulon)</td>
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<td>spoVIGA (ytrH)</td>
<td>$6.39 \times 10^{-5}$</td>
<td>12.4</td>
<td>cortex synthesis (σE regulon)</td>
<td>17.2</td>
<td>ND</td>
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<tr>
<td></td>
<td>spoVIGB (ytrI)</td>
<td>$1.58 \times 10^{-5}$</td>
<td>5.9</td>
<td>spore coat/cortex synthesis</td>
<td>11.9</td>
<td>ND</td>
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<tr>
<td></td>
<td>uppP</td>
<td>$2.66 \times 10^{-8}$</td>
<td>132.7</td>
<td>minor UPP phosphatase</td>
<td>0.4</td>
<td>distended intermembrane space</td>
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<td>yabQ</td>
<td>$2.97 \times 10^{-11}$</td>
<td>2206</td>
<td>cortex synthesis (σE regulon)</td>
<td>0.1</td>
<td>ND</td>
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<td></td>
<td>ydcC</td>
<td>$3.9 \times 10^{-6}$</td>
<td>8</td>
<td>slightly reduced σG activity (σE regulon)</td>
<td>13.7</td>
<td>small forespores some with reduced σG activity</td>
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<td>ylBj</td>
<td>$2.92 \times 10^{-13}$</td>
<td>801</td>
<td>cortex synthesis (σE regulon)</td>
<td>0</td>
<td>no observable phenotype</td>
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<td></td>
<td>ymdB</td>
<td>$2.64 \times 10^{-7}$</td>
<td>14.7</td>
<td>phosphodiesterase/bistable gene expression</td>
<td>11.4</td>
<td>ND</td>
</tr>
<tr>
<td></td>
<td>yqfC</td>
<td>$5.22 \times 10^{-4}$</td>
<td>32.3</td>
<td>unknown (σE regulon)</td>
<td>72.9</td>
<td>ND</td>
</tr>
<tr>
<td></td>
<td>yqfD</td>
<td>$1.54 \times 10^{-13}$</td>
<td>3448</td>
<td>similar to UDP-glucose 4-epimerase (σE regulon)</td>
<td>0</td>
<td>some small spores with reduced σG activity</td>
</tr>
<tr>
<td></td>
<td>ytvI</td>
<td>$6.68 \times 10^{-9}$</td>
<td>15.1</td>
<td>autoinducer-2 exporter family (σE regulon)</td>
<td>14</td>
<td>no observable phenotype</td>
</tr>
</tbody>
</table>

<sup>a</sup>p-value based on Mann Whitney U test comparing T0 and T24 libraries

<sup>b</sup>fold change refers to number of transposon insertions at T0 compared to T24

<sup>c</sup>sporulation efficiency (spo) was determined 24 h after starvation and refers to heat-resistant colony forming units (CFU) of null mutant compared to wild-type

<sup>d</sup>ND, not determined.

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predicted to have sporulation-specific promoters (S2 Table) [35]. To investigate whether any of these T5 hits were indeed required for timely differentiation, we chose eight for which transposon insertions were significantly underrepresented at T5 compared to T0 and T24 (Fig 2 and S2A Fig). This set included genes transcribed by \( \sigma^F \) (yqhH, yyaC), \( \sigma^E \) (ykvU, ykvI, prpB), and \( \sigma^G \) (yrbG, ykoN, yqgE). The roles of these genes in sporulation are unknown or poorly characterized [36,37] but based on transcriptional profiling, their expression appears to be largely under sporulation control. Wild-type and the null mutants were grown in sporulation medium and samples were removed at hour 5 and 24 after the onset of starvation and incubated at 80°C for 20 min to kill vegetative cells and sporulating cells that had not yet achieved heat-resistance. In
these assays, the sporulation efficiency of wild type at hour 5 ranged from 0.01%–0.4%. By contrast, in seven out of the eight mutants examined, there was a significant (7–1,500-fold) reduction in heat-resistant spores at this early time point (Fig 2B and S1 Data). Importantly and consistent with our Tn-seq data, at hour 24 the spore titers of the mutants were similar to wild-type (Fig 2B and S1 Data). These data suggest that many of the sporulation-induced genes previously identified by expression profiling play a direct role in the efficient and/or timely differentiation into a dormant spore. Furthermore, these data raise the possibility that many additional genes predicted to have sporulation promoters are indeed induced during sporulation and similarly function in efficient spore formation.

To investigate whether these mutants were indeed delayed in spore maturation and at what stage, we analyzed one (ΔykvU) that had a strong sporulation defect at hour 5 by fluorescence microscopy. Cells lacking ykvU initiated sporulation at approximately the same time as wild-type but exhibited a marked delay in the formation of phase-grey and phase-bright spores (Fig 2C and S2B Fig). Strikingly, a subset of the sporulating cells in the ΔykvU mutant had a novel phenotype that, to our knowledge, has never been reported previously. The space between the forespore and mother cell membranes was significantly larger than wild type (Fig 2C and S2B Fig). This "distended intermembrane space" was most apparent in merged images of cytoplasmic mCherry that reports mother cell gene expression under σE control and cytoplasmic CFP that shows forespore gene expression under σG control (Fig 2C and S2B Fig). A distinct gap separated these cytoplasmic signals. Furthermore, a phase-bright ring was detectable surrounding the forespore in these sporangia (Fig 2C and S2B Fig). The nature of this phase-bright material in the intermembrane space is not known and awaits further investigation (see below). All together, our data indicate that ykvU and likely many other genes identified in our screen play important roles in timely spore maturation.

Tn-seq Identifies Transposon Insertions That Trigger Premature Sporulation

In addition to the delayed sporulation mutants described above, visual inspection of the insertion profiles from the Tn-seq screen at hour 5 revealed a second class of insertions. We identified 43 chromosomal regions (S3 Table) where transposon insertions were significantly overrepresented at this early time point in sporulation (Figs 1C, 3A and S3A). Overrepresentation suggests that insertions at these positions result in premature and/or accelerated sporulation. Mutants of this variety with a so-called "derepressed" sporulation phenotype were described in the 1960s and 1970s [38–40]. In most cases, the mutations responsible for these phenotypes were never defined genetically. The insertions in our Tn-seq screen fell into two classes: those that were upstream of a gene or operon and likely result in over-expression of the adjacent locus, and those that were inserted within a coding region and likely cause gene inactivation (S3 Table). Among the regions in the former class were transposon insertions upstream of kinA and kinB (and possibly kinD) three of the histidine kinases that activate the master sporulation transcription factor Spo0A (Fig 3A and S3 Fig) [41]. Previous work indicates that over-expression of kinA and kinB can trigger entry into sporulation in LB medium prior to nutrient exhaustion [42]. We suspect that the promoter for the spectinomycin resistance gene in the magellan6 transposon [43] similarly provides additional expression of these kinases. In support of this idea, transposon insertions were commonly found in non-essential genes in operons with essential genes downstream (S4 Fig). Insertions were also overrepresented upstream of the opp (formerly, spo0K) operon encoding an oligopeptide permease [44,45]. This transporter has been shown to import quorum-sensing oligopeptides that promote sporulation and competence [46–49]. Thus, cells over-expressing this permease likely initiate
sporulation prematurely. Similarly, we identified insertions upstream of *dtpT* encoding a putative di- and tri-peptide permease [50], which we hypothesize can transport these oligopeptides or smaller unidentified signaling molecules. Moreover, insertions within the *scoC* gene were also overrepresented. ScoC is a transcriptional regulator that has been shown to repress expression of *opp* and *dtpT* [51,52]. Accordingly, in its absence, the level of these two permeases would increase. We also identified insertions in the *sda* gene, which encodes a cell-cycle regulator of KinA and KinB that prevents entry into sporulation during the initiation of replication [53,54]. Inactivation of *sda* could also result in premature entry into sporulation. The strong enrichment of transposon insertions upstream of and within genes that are known to function...
in the initiation of sporulation suggests that the enrichment of insertions adjacent to and within uncharacterized genes could define additional factors that control entry into this developmental process.

We selected 14 loci that were among the most overrepresented at T5 to validate our findings. Insertions at seven of these sites were predicted to cause gene inactivation (\textit{sigL}, \textit{scoC}, \textit{prsW}, \textit{sigW}, \textit{ysaA}, and \textit{bmrA}); for these we used deletion mutants from the knockout collection. Insertions at the other seven loci were predicted to cause over-expression, and for these, we built IPTG-inducible promoter fusions to four (\textit{sra}, \textit{yyaJ}, \textit{nhaC}, and \textit{yhzC}); we reconstructed transposon insertions upstream of two (\textit{dtpT} and \textit{helD}); and we used a previously characterized IPTG-inducible allele of \textit{kinA} [42]. Each null mutant and over-expression strain (for simplicity, referred to as the mutants) was mixed at a 1:1 ratio with wild-type cells that constitutively expressed \textit{lacZ}, and the two strains were co-cultured in sporulation medium. At the onset of starvation (T0), samples from the mixed cultures were removed and plated on LB agar containing X-gal to assess the ratio of mutant (white colonies) to wild-type (blue colonies) cells upon entry into sporulation (Fig 3B). At hour 5 (T5), samples were removed and incubated at 80°C for 20 min and the spores were plated on LB X-gal. The ratio of mutant to wild-type heat-resistant colony forming units (CFU) was compared to the ratio of CFU at T0 (Fig 3B). All of the strains that we tested had significantly (2–1,600-fold) more spores at T5 than wild type (Fig 3C and S1 Data). Importantly, virtually all the mutants (with the exception of \textit{ΔsigL} and \textit{ΔysaA}) had growth curves that were indistinguishable from wild type (S5 Fig and S1 Data), suggesting that in most cases, early spore maturation was not a consequence of premature starvation. Thus, our Tn-seq screen identified a new class of sporulation genes whose inactivation or over-expression results in earlier sporulation.

Finally, we chose four genes (\textit{helD}, \textit{nhaC}, \textit{yhzC}, and \textit{sigL}) to investigate whether their overexpression (or deletion in the case of \textit{sigL}) resulted in premature entry into sporulation. These four genes were chosen based on the strength of their phenotypes and because they had not been previously implicated in sporulation. This analysis was performed in mixed cultures with wild-type cells that contained a constitutively expressed \textit{mCherry} allele (\textit{P\textsubscript{veg}\textendash mCherry}) and a \textit{σF}-responsive promoter (\textit{P\textsubscript{spoIIQ}}) fused to \textit{cfp} to monitor an early stage in sporulation. The strains that overexpressed \textit{helD}, \textit{nhaC}, or \textit{yhzC} or lacked \textit{sigL} harbored the same \textit{σF}-responsive promoter fused to \textit{yfp} (\textit{P\textsubscript{spoIIQ}\textendash yfp}). The mixed populations were grown in sporulation medium and then visualized at the onset of starvation and during the early stages of spore formation. By hour 1 after the onset of starvation, 4–10-fold more mutant cells had entered sporulation (yellow fluorescent protein [YFP] positive, \textit{mCherry} negative) than wild type (cyan fluorescent protein [CFP] and \textit{mCherry} positive) (Fig 3D and S3B Fig). Importantly, a mixed population of wild-type cells harboring the sporulation reporters with and without cytoplasmic \textit{mCherry} had similar percentages of sporulating cells (Fig 3E). All together, these results suggest that most of the insertions sites identified by Tn-seq define genes that when over-expressed or inactivated promote premature initiation of sporulation.

**Tn-seq Screen Identifies 24 New Sporulation Genes**

Our success in identifying most of the previously identified sporulation genes prompted us to investigate whether any of the uncharacterized genes identified in the Tn-seq screen represent new sporulation loci. To assess the sporulation phenotypes of the null mutants, we individually tested strains derived from the \textit{B. subtilis} knockout collection for sporulation efficiency 24 h after the onset of exhaustion. For these experiments, we used null mutants for which the antibiotic cassettes had been excised to reduce the chance of polar effects. Among the 59 mutants tested, 24 were reduced in sporulation greater than 2-fold, 15 of these were reduced by more
than 5-fold, and nine by 10-fold or more (Table 1). The majority of the 35 false-positive hits fell into two classes: (i) mutants that had a moderate sporulation defect (50%–80% compared to wild type) and may actually represent true sporulation genes and (ii) mutants in which insertions were underrepresented but far from eliminated at T24. All of these genes were also identified as hits at T5 (S2 Table). For comparison, we also performed sporulation assays on 30 of the 37 genes known to be required for sporulation that have not been extensively characterized. The sporulation efficiencies of these mutants are reported in Table 1. Collectively, our Tn-seq screen uncovered 24 new genes that are required for efficient sporulation. Among these, none have been reported to be under sporulation control suggesting that many also function during vegetative growth. However, 12 of these genes are predicted to have sporulation-specific promoters (S2 Table) [35], raising the possibility that they are induced during development.

A Cytological Assay to Characterize the Sporulation Mutants

To rapidly assess the stage at which these new sporulation mutants were impaired, we built a strain that reports on all four compartment- and stage-specific sigma factors (σF, σE, σG, and σK) (Fig 4). The reporter for the first compartment-specific transcription factor σF that is activated in the forespore was a promoter (PspoIIQ) fusion to yfp [55]. The PspoIID promoter that is recognized by the first mother cell-specific transcription factor σE [56] was fused to mCherry. The reporter for the late-acting forespore transcription factor σG was a promoter (PsspB) fusion to cfp [57]. Finally, the PgerE promoter, recognized by the late-acting mother cell transcription factor σK [58], was fused to yfp. σF-dependent YFP fluorescence is first observed in the forespore at hour 1.5. By the time σK is activated in the mother cell (hour 4), σF activity is largely absent and YFP fluorescence is significantly reduced (Fig 4). Accordingly, we could use this fluorescent protein twice. For these experiments, we induced spore formation by re-suspending exponentially growing cells in minimal medium [59] to provide more synchronous sporulation. We monitored differentiation at four time points (hours 1.75, 2.5, 3.5, and 5) by phase contrast, membrane staining, and the four fluorescent reporters. As can be seen in Fig 4, in addition to the activity of the four stage- and compartment-specific transcription factors, this assay allowed us to assess polar division, engulfment, membrane fission, late spore development (phase grey), and spore maturation (phase bright).

Cytological Characterization of the New Sporulation Mutants

Using our reporter strain we analyzed 21 of the newly identified sporulation genes as well as nine additional sporulation mutants that had been previously identified but not extensively characterized. The mutants were sporulated by resuspension in groups of six and monitored at the time points shown in Fig 4. Among the 30 mutants analyzed, ten had no discernable phenotype in our assay (Table 1 and S6 Fig). These mutations are either impaired at stages that were not assayed, have weak or subtle phenotypes, or are not required when cells are sporulated by resuspension. Of the remaining 20, 11 (gidA, trmE, resA, yqhT, speE, gapB, smpB, rasP, nrmA, clpC, and minJ) appeared to be impaired in sporulation initiation having many fewer cells with polar divisions and forespore-specific gene expression at early time points (Fig 5 and S7 Fig). Furthermore, six mutants (gidA, ytxG, trmE, clpC, defB, and minJ) had defects that were specifically revealed by membrane staining (Fig 5 and S7 Fig). These mutants displayed defects in polar division, membrane migration, and/or membrane fission. Several of these mutants also had subtle membrane abnormalities in predivisional cells, suggesting that similar pathologies exist during vegetative growth but are significantly enhanced during morphogenesis (S7 Fig).

Six mutants (yqgE, ydcC, yqfD, nrmA, clpC, and defB) contained a subpopulation of sporulating cells in which σG activity was reduced or absent (Fig 5 and S7 Fig). Among these, many
displayed a partially penetrant small forespore phenotype that correlates with defects in $\sigma^G$ activity (Fig 5 and S7 Fig) (see below) [60,61]. Several of the mutants had distinct cytological phenotypes. All are described in Table 1 and the primary cytological data can be found in S7 Fig.
Below, we briefly highlight four genes (\textit{nrnA}, \textit{defB}, \textit{clpC}, and \textit{uppP}) that had representative or unusual mutant phenotypes (Fig 5). \textit{nrnA} encodes a nanoRNase responsible for degradation of RNA fragments \cite{62}. Cells lacking \textit{nrnA} had an intermediate sporulation defect (5\% of wild type) and were significantly delayed in the entry into sporulation. At hour 1.75, virtually no cells had polar septa (Figs 4 and 5A). All stages of differentiation analyzed were similarly delayed in this mutant. However, activation of \(\sigma^G\) appeared to be particularly impacted by the absence of \textit{nrnA} (Fig 5A). During sporulation in wild-type cells, \(\sigma^G\) activation is coincident with the completion of engulfment (Fig 4). In the \textit{nrnA} mutant, a substantial number of engulfed forespores had reduced \(\sigma^G\)-dependent CFP fluorescence. Interestingly, some of these cells also appeared to have smaller forespores: a phenotype that correlates with reduced \(\sigma^G\) activity. The role of RNA degradation in entry into sporulation has not been actively explored but these findings raise the possibility of a novel regulatory pathway.

\textit{defB} encodes a N-formylcysteine deformylase \cite{63}. The mutant had a modest sporulation defect (9\% of wild type) (Table 1). However, cytological analysis revealed an unusual engulfment phenotype in which the mother cell membranes appeared to migrate asymmetrically around the forespore meeting at non-polar positions (Fig 5B). In addition to and possibly a consequence of this uneven migration, membrane fission, the last step in engulfment, was delayed and impaired (Fig 5B). Furthermore, in a subset of cells, \(\sigma^G\) activity in the forespore was reduced or absent. The mechanism by which a deformylase impacts membrane migration is currently unknown; however, based on the strength of the cytological phenotype, further investigation is merited.

\textit{ClpC} is the ATPase subunit of the ATP-dependent ClpC-ClpP protease \cite{64}. Previous work indicates that ClpCP degrades the anti-\(\sigma^F\) factor SpoIIAB in the forespore helping to lock \(\sigma^F\) in its active state \cite{65}. The \(\Delta clpC\) mutant has a strong sporulation defect (0.4\% of wild type); however, it had not been cytologically characterized. Interestingly, the null mutant displayed a collection of cytological phenotypes, including delayed entry, asymmetric engulfment, reduced or no \(\sigma^G\) activity and a concomitant small forespore phenotype (Fig 5C). These phenotypes suggest that the ClpCP protease plays an important role in post-translational control of a host of proteins during development.

\textit{uppP} is predicted to encode a minor undecaprenyl pyrophosphate phosphatase involved in recycling the lipid carrier undecaprenyl phosphate \cite{66,67}. Cells lacking \textit{uppP} have a sporulation efficiency of 0.4\%. The mutant entered sporulation normally but was delayed in the production of phase-grey and ultimately phase-bright spores (Fig 5D) suggesting a defect in assembly of the spore cortex and/or spore dehydration \cite{68}. Strikingly, a subset of sporulating cells in this mutant had the same distended intermembrane space phenotype observed in the \textit{ykvU} mutant (Figs 2C and 5D). Furthermore, a similar phase-bright ring was present around the forespore (Fig 5D). Although this novel phenotype has not been reported previously, we have now observed it in two separate mutants. Intriguingly, YkvU is a member of the multi-drug/oligosaccharidyl-lipid/polysaccharide (MOP) exporter superfamily and a subset of the proteins in this family transport polysaccharide precursors linked to undecaprenyl phosphate across the membrane \cite{69}. This provides a possible explanation for the shared phenotype. YkvU is produced in the mother cell under the control of \(\sigma^E\) and specifically localizes in the membranes surrounding the forespore \cite{20,36,37}. In the absence of this putative flippase, an uncharacterized polysaccharide precursor would not be transported into the intermembrane space. Similarly, in the absence of UppP, recycling of the lipid carrier is predicted to be impaired, resulting in a reduction in the transport of this hypothetical precursor (S2C Fig). In the context of this model, one possible explanation for the distended intermembrane space in the mutants is that the polysaccharide produced in this putative pathway could help hold the two membranes surrounding forespore together. Dissecting the roles of these proteins and this unusual mutant phenotype awaits future investigation.
Finally, two new sporulation genes ($yqzE$ and $ywmB$) were characterized in greater detail. These experiments defined two novel factors: one (SpoIIIL) helps maintain forespore differentiation and $\sigma^G$ activity (see Box 1) and the other (SpoIIT) functions directly in the cell–cell signaling pathway that triggers proteolytic activation of $\sigma^E$ in the mother cell (see below).
YwmB (SpoIIIT) Functions in the Cell–Cell Signaling Pathway That Activates σ^E

Among the newly identified sporulation genes, the ywmB (renamed spoIIIT) mutant had one of the most revealing phenotypes in our cytological assay. A large population of sporulating cells

Box 1. SpoIIIL Is Required for σ^G Activation

Cells lacking yqzE (renamed spoIIIL) had a sporulation efficiency of 14% compared to wild type (Table 1). Cytological analysis of the mutant revealed that a subpopulation of sporulating cells had small forespores that often had reduced or no detectable σ^G activity (Fig 6B). These phenotypes are reminiscent of the small and collapsed forespores observed in cells lacking spoIIQ or any of the genes in the spoIIIA operon [60]. The forespore protein SpoIIQ and the eight SpoIIIA (spoIIAA-AH) proteins produced in the mother cell assemble a transenvelope complex (referred to as the SpoIIIA-IIQ complex) that connects the mother cell and the developing spore [60,70,71]. This complex is required to maintain transcriptional potential in the forespore, including σ^G activity, and is also required for forespore development [60,72]. It is thought that the complex functions as a specialized secretion system, enabling the mother cell to transport proteins and/or metabolites into the forespore and thereby maintain forespore physiology [60,72,73]. To investigate whether SpoIIIL functions in the SpoIIIA-IIQ pathway, we took advantage of the ΔspoIIIAH mutant phenotype. Unlike all other members of the SpoIIIA-IIQ complex, cells lacking SpoIIIAH have a relatively mild sporulation defect (1%–10% sporulation efficiency compared to 0.01%–0.001% for mutants in any other component), and the engulfed forespores are a mixture of small and normal sizes (Fig 6B) [60]. If the function of SpoIIIL relates to that of the SpoIIIA-IIQ complex, then we would expect a ΔspoIIIL ΔspoIIIAH double mutant to display synergistic phenotypes, and this is exactly what we observed (Fig 6B). The double mutant had almost no detectable σ^G activity and virtually all forespores were of small size, comparable to a mutant lacking the entire spoIIIA operon. Furthermore, like the spoIIIA operon mutant, the ΔspoIIIL ΔspoIIIAH double mutant had a sporulation efficiency of 0.001%.

spoIIIL is the last gene in the comG operon (Fig 6A) that encodes components of the DNA uptake machinery involved in natural competence [74]. spoIIIL is a small gene predicted to encode a novel cytoplasmic protein of 43 amino acids. The comG operon, which is not required for sporulation, is expressed during competence development under the control of the transcriptional regulator ComK [75]. Since ComK is not known to be active during sporulation, we suspected there was an additional promoter controlling spoIIIL. Noirot and co-workers [35] predicted a sporulation-specific promoter just upstream of spoIIIL (Fig 6A) and we found that this region promoted expression of YFP in the forespore in a σ^F-dependent manner (Fig 6C). Furthermore, the spoIIIL gene including this upstream region was sufficient to restore sporulation to the null mutant and largely restored sporulation and σ^G activity to the ΔspoIIIAH ΔspoIIIL double mutant (S8 Fig; S1 Data). Collectively, these results suggest that SpoIIIL is a component of the SpoIIIA-SpoIIQ complex on the forespore side. Its function in this transenvelope complex awaits future investigation.
lacking SpoIT had two polar divisions (referred to as an abortive disporic phenotype) and lacked $\sigma^E$ activity (Figs 7B, S9 and S10A and S1 Data). Both phenotypes are hallmarks of a gene involved in the activation of $\sigma^E$ [76]. This mother cell transcription factor is synthesized as an inactive membrane-associated precursor called pro-$\sigma^E$ (Fig 7F) [77,78]. Proteolytic processing

Fig 6. SpoIIIIL is required for $\sigma^G$ activation. (A) The spoIIIIL genomic locus. (B) Representative images of the $\Delta$spoIIIIL mutant and WT at hour 3.5 of sporulation. Small forespores with reduced $\sigma^G$ activity are indicated (yellow carets). The small forespores are also highlighted by the cytoplasmic mCherry signal in the mother cell reporting on $\sigma^E$ activity. (C) Representative images of the intergenic region between spoIIIIL and comGG (P_spoIIIIL) fused to yfp at hour 2 of sporulation. WT and a $\Delta$spoIIAC ($\Delta$sigF) mutant are shown. (D) Synergistic defects in $\sigma^G$ activity, forespore size, and sporulation efficiency in the $\Delta$spoIIIIL $\Delta$spoIIAH double mutant. Representative images at hour 3.5 of sporulation are WT, $\Delta$spoIIIIL, $\Delta$spoIIAH, the $\Delta$spoIIIIL $\Delta$spoIIAH double mutant and $\Delta$spoIIA. Spore titer relative to wild-type spores at hour 30 are indicated at the right. Scale bar is 2 $\mu$m.

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**Fig 7. SpoIIT is required for efficient proteolytic processing of pro-σ^E.** (A) The spoIT genomic locus. Three promoters for the essential gene murAA are predicted [35] to reside in the spoIT coding sequence. (B) Cytological analysis of a nonpolar spoIIT deletion mutant that leaves the three promoters intact. Representative images of WT, ΔspoIIT, and ΔspoIIR mutants at hour 2 of sporulation. σ^F activity was monitored by P^{spoIIQ}-yfp (green) and σ^E activity by P^{spoIID}-mcherry (blue). Scale bar indicates 1 μm. Larger fields of sporulating cells and additional time points can be found in S9 Fig. (C) Transcriptional fusions identify a σ^F-responsive promoter upstream of spoIIT. Two spoIIT promoter fusions to yfp are shown: one includes the upstream gene ywzB (labeled B) and its predicted σ^A promoter and the other contains 200 bp proximal to the spoIIT start codon. Expression from each fusion was examined in WT and a ΔspoIIAC (ΔsigF) mutant at hour 1.5 of sporulation. (D) Localization of pro-σ^E-gfp at hour 2 of sporulation in WT, ΔspoIIT, and ΔspoIIR. Pro-σ^E-gfp is membrane-associated while σ^E-gfp is cytoplasmic and nucleoid-associated. (E) SpoIIT is required for reconstitution of the pro-σ^E processing pathway in vegetatively growing cells. β-galactosidase assay monitoring σ^E activity (P^{spoIID}-lacZ) in strains expressing spoIIR spoIIGA, and pro-sigE during vegetative growth in the presence and absence of spoIIT. As a negative control, a strain only expressing spoIIGA and pro-sigE was also tested. Expression of spoIIGA/sigE/spoIIR was induced by the addition of 500 μM IPTG to cells growing exponentially (OD 0.2) in rich medium (casein hydrolysate [CH]). At the indicated times after induction, a sample was removed and analyzed for β-galactosidase activity. Error bars represent standard error of the mean from two biological replicates. The raw underlying numerical data for Fig 7E can be found in S1 Data. (F) Model for the proteolytic activation of σ^E under σ^F control. SpoIIR (red) and SpoIIT (purple) are made in the forespore and secreted into the intermembrane space where they activate the processing enzyme SpoIIGA (blue).

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of pro-σE releases the mature and active transcription factor into the mother cell cytoplasm [79,80]. A signaling protein (SpoIIR) that is made in the forespore under the control of σE is secreted into the intermembrane space [81–83] and activates a protease (SpoIIGA) present in the septal membranes on the mother cell side (Fig 7F) [84–86]. Mutants in spoIIR or spoIIIGA fail to activate σE resulting in abortive disporic sporangia (Fig 7B). The spoIIIT mutant phenotype was not as penetrant as the ΔspoIIR and ΔspoIIIGA mutants. At hour 2.5 of sporulation ~15% of the sporulating cells successfully activate σE and continue down the sporulation pathway (Figs 7B and S10A and S1 Data). Although SpoIIT is not absolutely required for σE activation, the identification of a new factor that might participate in this well-characterized signaling pathway prompted us to investigate SpoIIT further.

In our initial characterization of spoIIIT, we used the insertion-deletion mutant from the B. subtilis knockout collection, which lacks the entire spoIIIT coding sequence. This mutant had the strongest phenotype among all our newly identified sporulation genes, yielding no viable spores. However, the mutant could not be complemented by the spoIIIT gene in trans. spoIIIT is part of a three-gene operon (ywzB-spoIIIT-murAA) (Fig 7A) and we wondered whether the antibiotic resistance gene that replaced spoIIIT was causing inappropriate expression of murAA, leading to the sporulation defect. We attempted to remove the erythromycin (erm) resistance cassette using Cre/loxP-mediated recombination [87], but were unable to excise it. There are three predicted promoters [35] within the spoIIIT gene (Fig 7A) that could contribute to expression of the essential murAA gene, and in the insertion-deletion mutant, the erm promoter might provide this expression. Accordingly, we generated an in-frame deletion of spoIIIT that removes the first 63 amino acids of the spoIIIT coding sequence (including its predicted signal peptide) but retains all three promoters within the spoIIIT gene. This mutant was viable and exhibited a more mild but reproducible sporulation defect (38% of wild type). Importantly, similar to the original mutant, it produced a high number of abortive disporic sporangia and failed to efficiently activate σE (Figs 7B, S9 and S10A and S1 Data). Furthermore, the mutant could be complemented by expression of spoIIIT from an ectopic locus (S10B Fig). Thus, spoIIIT is a bona fide sporulation gene and is required for efficient activation of σE. The nonpolar spoIIIT deletion was used for all the characterization presented in Fig 7, S9 Fig and S10 Fig and described below.

First, we investigated whether SpoIIT functions in the proteolytic activation of σE. To do this, we examined a pro-σE-gfp fusion during sporulation by fluorescence microscopy [80]. The pro-domain of σE is amphipathic resulting in membrane localization. Proteolytic processing releases mature σE-gfp into the cytoplasm and upon association with core RNA polymerase, it co-localizes with the nucleoid. Wild-type, ΔspoIIT, and ΔspoIIIGA mutants harboring a functional pro-σE-gfp fusion were induced to sporulate by resuspension and were examined by fluorescence microscopy (Fig 7D). Consistent with the idea that SpoIIT is required for efficient pro-σE processing, in most sporulating cells at hour 2, pro-σE-GFP remained membrane-associated in the ΔspoIIT mutant (Fig 7D). Immunoblot analysis of pro-σE processing from sporulating cells (S10C Fig), provided further support for this conclusion.

The spoIIIT gene is predicted to encode a secreted protein expressed under the control of the housekeeping sigma factor σA (Fig 7A). If SpoIIT functions, as our data suggest, in the activation of the pro-σE processing enzyme SpoIIGA it must act in the intermembrane space (Fig 7F). Because secretion into the intermembrane space from the mother cell side is thought to be inefficient [88], we wondered whether SpoIIT, like the signaling protein SpoIIR, is produced in the forespore. Although spoIIIT is not predicted to have a sporulation-specific promoter [35], we constructed yfp fusions to the genomic regions upstream of spoIIIT (Fig 7C) and analyzed compartment-specific expression during sporulation. A 774 bp fragment that encompassed the σA promoter, the ywzB gene, and the intergenic region between ywzB and spoIIIT fused to yfp displayed modest YFP fluorescence in non-sporulating and predivisional cells (Fig 7C).
Strikingly, in sporulating cells, strong forespore-specific YFP fluorescence was observed (Fig 7C). A 360 bp fragment that contained part of ywzB gene and the intergenic region between ywzB and spoIIT exclusively displayed forespore-specific gene expression. Furthermore, forespore-specific YFP fluorescence from both reporters required σF (Fig 7C). These data indicate that spoIIT is expressed at low levels during vegetative growth but is induced to higher levels in the forespore under the control of σF. Thus, SpoIIT and SpoIIR are both produced in the forespore and likely secreted into the intermembrane space. Interestingly, the spoIIT gene is located at -37° on the B. subtilis chromosome, in a region that, like spoIIR, is trapped in the forespore at the time of polar division [89]. Thus, both genes are transcribed as soon as σF is activated, ensuring timely activation of σE in the mother cell [90,91].

Finally, to address whether SpoIIT acts directly in the σF signaling pathway, we took advantage of an observation made over two decades ago in which cells artificially engineered to express pro-σE, SpoIIGA, and SpoIIR during vegetative growth were found to trigger pro-σE processing and induce σE activity (as assayed by a σE-responsive promoter (PspoIID) fused to lacZ) [81,83]. In other words, SpoIIR-SpoIIGA signaling could be reconstituted in the absence of sporulation. Since spoIIT is itself expressed vegetatively (Fig 7C), to assess its impact on σE activation in this reconstituted system, we compared σE-dependent β-galactosidase activity in the presence and absence of SpoIIT (Fig 7E and S1 Data). In support of the idea that SpoIIT is directly involved in this signaling pathway, σE activity was markedly reduced in its absence. Collectively, these data support a model in which SpoIIT functions as a co-activator with SpoIIR in triggering SpoIIGA-dependent processing of pro-σE. Bioinformatic analysis indicates that SpoIIT, like SpoIIR, is present in virtually all endospore formers (S11 Fig), lending further support to the idea that both proteins are part of a core signaling pathway.

**Discussion**

Transposon-sequencing and other variations of signature-tagged mutagenesis [43,92–94] were originally developed to study commensal and pathogenic bacteria. In most cases, the molecular genetic tools in these organisms are less developed than classical models like Escherichia coli, B. subtilis and Caulobacter crescentus, and the benefits of these approaches were immediately apparent. This study, among others [95–98], highlights the utility of Tn-seq for defining gene function in well-studied model bacteria. By combining Tn-seq and the recently completed ordered null mutant collection, we have identified and validated over 20 new genes that are required for spore formation, uncovered phenotypes for many uncharacterized sporulation-induced genes, and discovered a novel set of loci that initiate development prematurely. In a complementary approach, Koo and Gross directly screened the ordered knockout library for sporulation mutants and identified an overlapping set of genes (personal communication). Tn-seq could similarly be used to identify novel genes required for other aspects of bacterial physiology, including natural competence, stationary phase survival, swimming, swarming, cell envelope stress, and DNA damage. Driven by Tn-seq and complemented by whole-genome sequencing, we are entering a renaissance in classical genetics of model organisms. When combined with ordered deletion collections, the genetic relationships suggested by these approaches can be systematically validated and explored with unprecedented speed.

**Outlook**

The Tn-seq screens performed in this study expanded the number of genes required for sporulation to >170, but we anticipate that there are many additional factors yet to be discovered. Some of these missing genes are likely to be “condition-specific,” while others were probably missed due to limitations of the approach. Our screens were performed in Difco Sporulation
Medium (DSM) in aerated shaking flasks. We suspect that there are genes specifically required for sporulation in media of different composition, on solid surfaces, or at the air–water interface in the context of a biofilm community. Other genes might be specifically required for sporulation within a polymicrobial community. Tn-seq opens the door to identify these condition-specific sporulation factors. Furthermore, a Tn-seq survey of this variety has the potential to define a core set of genes required for spore formation under all conditions. An important limitation of the Tn-seq approach is its inability to identify sporulation genes that are essential for viability, as insertions in these genes would be absent from our reference (T0) dataset. Similarly, small sporulation genes for which there were a limited number of transposon insertions could have been missed in our screens. Finally, an additional limitation is the inability to identify sporulation genes with redundant functions. Transposon insertions in any one of the redundant genes would not have a sporulation defect and would therefore not have been underrepresented among the heat-resistant spores. By building transposon libraries in mutants with mild or undetectable sporulation phenotypes, Tn-seq could be used to identify synthetic sporulation defects. Such an approach has the potential to unmask a large collection of functionally redundant sporulation factors [27].

Our study also provides important lessons related to the recently completed ordered knockout collection. The antibiotic cassettes used to generate the mutants in this library lack transcriptional terminators to ensure expression of genes in operons. However, by avoiding one polar effect, a new one may have been introduced: over-expression of downstream genes. The contribution of over-expression to particular phenotypes can be easily assessed by Cre-mediated excision of the antibiotic cassette. Our analysis of spoIIT revealed an additional way in which null mutants can generate phenotypes. The spoIIT coding region contains three predicted promoters that are all absent in the knockout. Their absence could explain the strong sporulation phenotype of the original mutant and/or our inability to excise the antibiotic cassette. Although complementation remains the gold standard for defining the genetic basis of a phenotype, promoter predictions [35] will help inform decisions about whether or not more precise deletions should be constructed, as was done in our study.

New Sporulation Genes

Among the new mutants we identified that impaired spore-formation, the largest class were those that impacted the timing and/or efficiency of initiation. Many inputs must be integrated in the decision to enter this developmental pathway, including nutritional status, cell-cycle stage, environmental cues, and quorum signals. Inactivation of individual genes in one of these pathways is likely to have a relatively modest impact on sporulation efficiency. This could account for why many of these mutants have been missed in classical genetic screens. Furthermore, since many of these genes are unlikely to be under sporulation control, they would not have been implicated in sporulation based on transcriptional profiling. Some of the mutants we identified likely impair some aspect of growth or metabolism that only indirectly impacts initiation of sporulation. However, we suspect others will provide insights into the cues that are sensed and integrated during starvation.

Interestingly, the other large class of mutants we discovered impacted the activity of the late-acting forespore transcription factor σ^G. σ^G becomes active after the completion of engulfment, when the forespore resides as a membrane-bound protoplast in the mother cell cytoplasm and is no longer in direct contact with the extracellular environment. Recent work suggests that the loss of σ^G activity in spoIIA and spoIIQ mutants reflects a loss or reduction in metabolic activity of the spore [60,72]. These mutants have reduced transcriptional potential,
are smaller in size, and appear to collapse. In light of these findings, we speculate that the large collection of mutants containing a subpopulation of cells with small forespores and impaired $\sigma^G$ activity reflects the physiologically sensitive state of the forespore at this stage in development. After engulfment is complete, the forespore likely relies on nutrients from the mother cell and catabolism of its own macromolecules to support its developmental program and its differentiation into a dormant spore. Accordingly, mutants impaired in $\sigma^G$ activity could be affecting spore metabolism and therefore represent a particularly interesting and unexplored class of genes for future analysis.

Developmentally Delayed Mutants

One of the most exciting discoveries to emerge from our screens is the identification of 36 uncharacterized sporulation-induced genes whose inactivation appears to delay spore maturation. Many of these genes were deleted and tested for sporulation efficiency when they were originally identified [20,21,23], and almost all lacked phenotypes. Others were tested in competition experiments subjecting mixed populations to rounds of sporulation and germination [26]. Our data suggest that many of these genes have quantifiable phenotypes that can be easily assayed by examining sporulation efficiency at early time points. The specific defect in spore maturation can then be assessed by fluorescence microscopy either alone or in mixed culture (Figs 2C and 3D). The identification of known sporulation genes like racA and fisB as delayed mutants, coupled with the discovery of a novel phenotype for the ykvU mutant, suggests that a systematic analysis of the remaining mutants could uncover additional cytological phenotypes reflecting novel steps in spore morphogenesis. In addition, 42 genes that are predicted to have sporulation-specific promoters that have not been experimentally validated were also identified as delayed mutants. Our analysis raises the possibility that many of these genes could be specifically induced during spore formation. We suggest that transposon-sequencing can serve as a powerful complement to expression profiling and computational predictions in defining the complete sets of regulons.

Premature Initiators

Finally, the discovery of transposon insertions that promote premature sporulation was an unexpected outcome of the Tn-seq screen and exemplifies the power of this approach. As discussed above, entry into sporulation requires the integration of multiple inputs, and mutants deficient in any one of them frequently exhibit only modest defects in sporulation. However, inappropriate overstimulation of one of these pathways has the potential to induce premature sporulation, leading to the observed “accelerator” phenotype in our Tn-seq screen. Thus, characterization of these genes could provide insight into the triggers for entry into spore maturation. Interestingly, several of the genes predicted to cause premature sporulation due to over-expression encode transporters raising the possibility that uptake of uncharacterized molecules in the medium could induce early sporulation. Furthermore, we showed that premature initiation could be elicited by over-expression of the helD gene, encoding a DNA helicase that increases transcriptional cycling [99]. Evidence suggests that HelD is involved in transcriptional adaptation to new environmental conditions and we propose that increased expression of HelD promotes more rapid induction of the transcriptional programs associated with sporulation. Finally, mutants in envelope stress pathways, in arginine utilization and phenylalanine metabolism all appear to trigger premature sporulation suggesting certain pathways and metabolites could serve as control points for monitoring starvation or nutrient limitation.
Two Forespore Signals

Finally, we turn to the pro-\(\sigma^E\) processing pathway and our discovery that a second secreted protein produced in the forespore promotes the proteolytic activation of \(\sigma^E\) in the mother cell. Collectively, our analysis supports a model in which both SpoIIR and SpoIIT function as signaling molecules. Although it remains unclear how these proteins trigger SpoIIGA activity, both genes are present in virtually all endospore formers, suggesting important complementary roles. Interestingly, the second and later-acting signal transduction pathway between the forespore and mother cell similarly utilizes two signaling proteins. In this case, \(\sigma^G\) directs the synthesis of two secreted proteases in the forespore: SpoIVB and CtpB [100,101]. These proteases in turn activate an unrelated membrane metalloprotease that processes pro-\(\sigma^K\) in the mother cell [102,103]. SpoIVB is the primary protease, which cleaves a negative regulator of the pro-\(\sigma^K\) processing enzyme [104]. CtpB further proteolyzes this negative regulator ensuring efficient and robust activation of \(\sigma^K\) [105]. SpoIVB, like SpoIIR is essential for \(\sigma^K\) activation, while cells lacking CtpB are delayed but not blocked in \(\sigma^K\) activation [106]. Although neither SpoIIR nor SpoIIT are proteases, by analogy to the pro-\(\sigma^K\) processing pathway, we hypothesize that SpoIIR triggers SpoIIGA proteolytic activity and SpoIIT helps maintain SpoIIGA in its active state. In vitro reconstitution of this signaling pathway will provide a clearer picture of the role of both of these signaling molecules in activating pro-\(\sigma^E\) processing.

Methods

General Methods

\(B. subtilis\) strains were derived from 168 [107] or PY79 [108]. Unless otherwise indicated, cells were grown in LB or CH medium at 37°C. Sporulation was induced by resuspension at 37°C according to the method of Sterlini-Mandelstam [59] or by nutrient exhaustion in supplemented DS medium (DSMcomplete) [109]. Sporulation efficiency was determined in 24–36 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. \(\beta\)-galactosidase assays were performed as described previously [102]. Insertion-deletion mutants come from the BKE knockout collection or were generated by isothermal assembly [110] of PCR products followed by direct transformation into \(B. subtilis\). All BKE mutants were back-crossed into \(B. subtilis\) 168 before assaying and prior to antibiotic cassette removal. Antibiotic cassette removal was performed using a temperature-sensitive plasmid that constitutively expresses Cre recombinase [87]. Tables of strains (S4 Table), plasmids (S5 Table) and oligonucleotide primers (S6 Table) and a description of strain and plasmid constructions can be found in S1 Methods.

Transposon Insertion Sequencing

Transposon insertion sequencing (Tn-seq) was performed on two independently generated libraries as described previously [43,87,97]. Briefly, Magellan6x transposon library DNA was transformed into competent \(B. subtilis\) 168 cells. ~750,000 transformants were pooled, aliquoted, and frozen. An aliquot was thawed, washed in DSM, and diluted into 50 ml DSM at an OD600 of 0.05. Samples were harvested at the onset of starvation (T0) and 5 (T5) and 24 (T24) h later. The T5 and T24 samples were incubated at 80°C for 20 min, and plated on LB agar. ~750,000 colonies from germinated spores from each were pooled. Genomic DNA was extracted from all three samples and digested with MmeI, followed by adapter ligation. Transposon-chromosome junctions were amplified in 16 PCR cycles. PCR products were gel-purified and sequenced on the Illumina HiSeq platform using TruSeq reagents (Tufts University TUCF Genomics facility). Raw sequence reads are available in the Sequence Read Archive (Accession: SRP066259). Reads were
mapped to the *B. subtilis* 168 genome (NCBI NC_000964.3), tallied at each TA site, and 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. To identify genomic regions (including inter-genic regions) in which reads were statistically overrepresented, we performed a sliding window Mann Whitney *U* test using a 13 TA site window size across all TA sites which register at least one read in both the T0 and T5 experiments. Regions that had a *p*-value ≤ 0.02 and were at least 3-fold enriched in T5 reads compared to T0 reads are listed in S3 Table.

**Fluorescence Microscopy**

Sporulating cells were concentrated by centrifugation at 8 krpm for 1 min and immobilized on 2% agarose pads. Fluorescence microscopy was performed using an Olympus BX61 microscope equipped with a UplanF1 100X phase contrast objective and a CoolSnapHQ digital camera (Photometrics) or a Nikon TE2000 inverted microscope with a Nikon CFI Plan Apo VC 100X objective. Images were acquired using Metamorph software. Membranes were stained with TMA-DPH (50 μM) (Molecular Probes) and fission of mother cell membranes was assessed as previously described [34]. Exposure times were 400 ms for TMA-DPH and mCherry, 200 ms for YFP and CFP, and 1,000 ms for σ^E^-gfp. Image analysis and processing were performed in Metamorph.

**Immunoblot Analysis**

Whole-cell lysates from sporulating cells were prepared as previously described [111]. Samples were heated for 5 min at 65°C prior to loading. Equivalent loading was based on OD_{600} 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% nonfat milk with 0.5% Tween-20 for 1 h. Blocked membranes were probed with anti-σ^E^ (1:100) [112], anti-σ^F^ (1:5000) [65], anti-His (1:4000) (Genscript), or anti-σ^A^ [113] primary antibodies diluted into PBS 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.

**Supporting Information**

S1 Data. Raw numerical values for Tn-seq analysis, sporulation efficiency measurements, gene expression, and image quantitation.

(XLSX)

S1 Fig. Tn-seq identifies known sporulation genes. Examples of transposon insertion profiles for ten known and “named” sporulation loci. Transposon insertions at the onset of starvation (T0) and after 24 h (T24) of sporulation are displayed for nine regions of the genome. The height of each vertical line reflects the number of sequencing reads at this position. “VT” refers to *spoVT*, “CB” to *spoIVCB*, and “IIIC” to *spoIIIC*.

(PDF)

S2 Fig. Developmentally delayed sporulation mutants identified by Tn-seq. (A) Examples of transposon insertion profiles for six mutants delayed in sporulation. Transposon insertions at the onset of starvation (T0) and after 5 (T5) or 24 (T24) h of sporulation are displayed for six regions of the genome. (B) Additional examples of the ΔykvU mutant phenotype as described in Fig 2B. Phase contrast, membrane staining and overlays of the mother cell (red) and forespore (green) cytoplasmic fluorescent proteins. Yellow carets highlight phase bright rings, the
enlarged outer forespore membrane, and the gap between the two compartments. (C) Model for the functions of YkvU and UppP during sporulation. YkvU is a putative flippase produced in the mother cell that could transport lipid-linked precursors into the intermembrane space to synthesize an unidentified polysaccharide. UppP is an undecaprenyl pyrophosphate (UPP) phosphatase potentially involved in the recycling of the undecaprenyl pyrophosphate (UP) lipid carrier.

**S3 Fig. Transposon insertions enriched at hour 5 of sporulation.** (A) Examples of transposon insertion profiles for 13 regions of the genome at the onset of starvation (T0) and after 5 (T5) or 24 (T24) h of sporulation. The scale of maximum reads is 10,000 (or 3,000 where indicated) to highlight the enrichment of reads at T5. The genes colored pink below the insertion profiles are predicted to be the one impacted by the insertions. For yhal, yhal, scoC, ysaA, prsW, sigW, sda, yqeF, bmrA, and sigL, the insertions are predicted to cause gene inactivation. For opp, kinB, slrA, sigH, dtpT, and yyal, the insertions are predicted to cause over-expression.

(B) Premature initiation of sporulation in the ΔsigL mutant and strains over-expressing (OE) yhzC and nhaC. All three strains harbor a PspoIIQ-yfp fusion to monitor and early stage of sporulation. Strains were separately mixed 1:1 with wild type (WT) harboring constitutively expressed mCherry and PspoIIQ-cfp. Images are from hour 1 after the onset of starvation. The percentage of sporulating cells (as assayed by σF activity) in the mutant (YFP positive, mCherry negative) and the WT (CFP positive, mCherry positive) are shown for each culture.

**S4 Fig. Magellan6 insertions are non-polar.** Transposon insertion profiles for four regions of the genome in which an essential gene (pink) resides downstream of a nonessential gene (blue) in an operon. The recovery of insertions in the nonessential blue gene provides evidence that the transposon can promote expression of the downstream essential gene. The profiles depicted are derived from cells harvested at the onset of starvation.

**S5 Fig. Most mutants that sporulate prematurely enter stationary phase in a manner similar to wild type.** Growth curves of all validated prematurely sporulating mutants. Indicated mutants and over-expressing (OE) strains are shown in comparison to WT. For each growth curve, the WT is depicted in black and the mutant in red. ΔsigL and ΔysaA mutants are the only strains that enter stationary phase earlier than WT. The raw underlying numerical data for S5 Fig can be found in S1 Data.

**S6 Fig. Sporulation mutants with no cytological phenotypes through hour 5 of sporulation.** Representative images of ten mutants harboring four transcriptional fusions at hour 5 (T5) of sporulation. Phase contrast, membrane staining and the indicated fluorescent fusions are shown. Scale bars are 2 μm.

**S7 Fig. Additional sporulation mutants with cytological phenotypes.** Representative images of 14 mutants harboring four transcriptional fusions at the indicated times after the initiation of sporulation (A) At hour 1.75 (T1.75), the ΔgapB mutant has few polar septa indicative of a delay in entry into sporulation. At hour 5 (T5), very few cells have completed engulfment and most forespores lack detectable σG activity. (B) The ΔresA mutant at T1.75 has few cells with polar septa and a small number with σF activity indicative of an early delay in sporulation. At hour 5 (T5), many cells have completed engulfment but most have not become phase-grey. (C)
The \( \Delta qgT \) mutant at T1.75 has many cells with polar septa but relatively few with \( \sigma^F \) activity. At T5, many cells have completed engulfment but most have not become phase-grey. (D) The \( \Delta qfD \) mutant at hour 5. A subset of sporulating cells have small forespores (white carets) and reduced \( \sigma^G \) activity (yellow carets). The two phenotypes are not always correlated in this mutant. Enlarged examples of small forespores and forespores with reduced \( \sigma^G \) activity are shown on the right. (E) The \( \Delta speE \) mutant at T1.75 has few cells with polar septa and almost no cells with \( \sigma^F \). At T5, many cells have completed engulfment but most have not become phase-grey. (F) The \( \Delta mspB \) mutant at T1.75 has few cells with polar septa and no \( \sigma^F \) activity. At T5, most cells have completed engulfment but no forespores are phase-grey. (G) The \( \Delta minJ \) mutant at T1.75 has few cells with polar septa. At T2.5, some of the engulfed forespores lack \( \sigma^F \) activity. There are a few large cells with strongly staining membranes that have \( \sigma^F \) activity (inset). By T5, some spores are phase bright but many sporulating cells have arrested or lysed. (H) The \( \Delta pgcA \) mutant and the \( \Delta gtaB \) mutant at hour 5 have relatively normal-sized forespores and small mother cells. These phenotypes are consistent with their small size during vegetative growth [114]. (I) The \( \Delta trmE \) mutant at T1.75 has longer cells; few with polar septa and almost none with \( \sigma^F \) activity. At T3.5, membrane engulfment appears aberrant with some unusual membrane invaginations (inset). (J) The \( \Delta gidA \) mutant at hour 1.75 and 3.5. \( \Delta gidA \) and \( \Delta trmE \) have similar phenotypes; (K) The \( \Delta ttxG \) mutant at hour 1.75, 2.5, and 3.5 of sporulation. The mutant exhibits membrane defects that are most pronounced during engulfment. (L) The \( \Delta rasP \) mutant at T1.75 has few cells with polar septa and almost no cells with \( \sigma^F \) activity. (M) The \( \Delta dydC \) mutant at hour 5 has small forespores (yellow carets) some with reduced \( \sigma^G \) activity. Scale bars are 2 \( \mu \)m.

**(PDF)**

**S8 Fig.** The \( \textit{spoIII} \) gene with its \( \sigma^F \)-responsive promoter complements the \( \textit{spoIII} \) mutant. (A) Representative images of sporulating cells at hour 3.5 in the indicated genetic backgrounds, highlighting complementation of the \( \Delta spoIII \) mutant at the level of \( \sigma^G \) activity (\( P_{\text{spoB-cfp}} \)), forespore size, and the formation of heat-resistant spores after sporulation by resuspension. Scale bars are 2 \( \mu \)m. (B) Percentage of heat-resistant spores after 24 h in nutrient exhaustion medium (DSM), highlighting the complementation of \( \textit{spoIII} \). Histograms from left to right are WT, \( \Delta spoIII \) mutant, \( \Delta spoIII \), \( \Delta spoIIIAH \Delta spoIII \), the complemented \( \Delta spoIII \) mutant, the complemented \( \Delta spoIIIAH \Delta spoIII \) double mutant, the complemented \( \Delta spoIII \) mutant with an IPTG-inducible \( \textit{spoIII} \) allele, the complemented \( \Delta spoIIIAH \scrip{spoIII} \) double mutant with an IPTG-inducible \( \textit{spoIII} \) allele. The raw underlying numerical data for S8B Fig can be found in S1 Data.

**(PDF)**

**S9 Fig.** Cytological phenotype of the \( \Delta spoII \) mutant. Large fields of sporulating WT and the \( \Delta spoII \) mutant from the experiment in Fig 7B are shown at hour 3 (T3) and hour 5 (T5). Cells were stained with the membrane dye TMA-DPH (shown in red), and express both \( \sigma^F \) and \( \sigma^E \) activity reporters (green and blue, respectively). Although ~40% of the cells form dormant spores, the majority fail to complete differentiation. Scale bar indicates 1 \( \mu \)m.

**(PDF)**

**S10 Fig.** Phenotype of the \( \Delta spoII \) mutant. (A) Quantitation of the \( \Delta spoII \) mutant phenotype from the experiment shown in Fig 7B. Bar graph indicates the percentage of sporulating cells at hour 2 with \( \sigma^F \) activity, \( \sigma^E \) activity, and with two polar septa for each strain. More than 1,000 sporangia were analyzed for each strain. The raw underlying numerical data for S10A Fig can be found in S1 Data. (B) Complementation of the \( \Delta spoII \) mutant. Representative images of sporulating cells at hour 2 in the indicated genetic backgrounds, highlighting complementation of the \( \Delta spoII \) mutant at the level of \( \sigma^E \) activity (\( P_{\text{spoII}\_mCherry} \)), loss of the abortive
disporic phenotype, and the formation of heat-resistant spores. The complementing allele of spoIIT was provided in trans under the control of an IPTG-inducible (Pspank) promoter. IPTG was added upon the initiation of sporulation to 500 μM. Scale bar indicates 1 μm. (C) Immunoblot analysis of proteolytic processing of pro-σE in WT, ΔspoIIT, ΔspoIIR, and ΔsigE mutants at the indicated times after the initiation of sporulation. σF protein levels were analyzed to compare entry into sporulation. (D) Immunoblot analysis of SpoIIR-His6 levels in WT and a ΔspoIIT mutant at the indicated times after the initiation of sporulation.

S11 Fig. spoIIT is broadly conserved among endospore-forming bacteria. (A) Occurrence of spoIIT across the bacterial phylogenetic tree. Red bands indicate the presence of a predicted spoIIT ortholog in each taxon. The phylogenetic tree was constructed in PhyloT and was displayed and manually pruned in iTOL. (B) The spoIIT locus (red) and surrounding chromosomal loci in representative set of endospore-forming bacteria. Also displayed is the corresponding region in Sporosarcina newyorkensis, an endospore-forming bacterium that does not contain a predicted spoIIT ortholog.

S1 Methods. Plasmid and strain construction.

S1 Table. Sporulation genes identified at T24 by Tn-seq.

S2 Table. Genes that were underrepresented at T5 compared to T0.

S3 Table. Regions of the genome in which transposon insertions were overrepresented at T5.

S4 Table. List of strains used in this study.

S5 Table. List of plasmids used in this study.

S6 Table. List of oligonucleotide primers used in this study.

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Author Contributions

Conceived and designed the experiments: AJM CDAR DZR. Performed the experiments: AJM CDAR JB. Analyzed the data: AJM CDAR JB HCL DZR. Contributed reagents/materials/analysis tools: AJM CDAR JB HCL TGB DZR. Wrote the paper: AJM CDAR JB HCL TGB DZR.
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