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Gene Conservation among Endospore-Forming Bacteria Reveals Additional Sporulation Genes in *Bacillus subtilis*

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The capacity to form endospores is unique to certain members of the low-G+C group of Gram-positive bacteria (*Firmicutes*) and requires signature sporulation genes that are highly conserved across members of distantly related genera, such as *Clostridium* and *Bacillus*. Using gene conservation among endospore-forming bacteria, we identified eight previously uncharacterized genes that are enriched among endospore-forming species. The expression of five of these genes was dependent on sporulation-specific transcription factors. Mutants of none of the genes exhibited a conspicuous defect in sporulation, but mutants of two, *ylxY* and *ylaA*, were outcompeted by a wild-type strain under sporulation-inducing conditions, but not during growth. In contrast, a *ylmC* mutant displayed a slight competitive advantage over the wild type specific to sporulation-inducing conditions. The phenotype of a *ylaA* mutant was ascribed to a defect in spore germination efficiency. This work demonstrates the power of combining phylogenetic profiling with reverse genetics and gene-regulatory studies to identify unrecognized genes that contribute to a conserved developmental process.

The formation of endospores is a distinctive developmental process wherein a dormant cell type (the endospore) is formed inside another cell (the mother cell) and ultimately released into the environment by lysis of the mother cell (1, 2). Endospores are metabolically inactive and highly resistant to environmental stresses, such as heat, radiation, chemicals, and desiccation (3). At the same time, these spores monitor the environment and are capable of rapidly resuming growth when conditions are favorable (4). Endospore formation is unique to the low-G+C group of Gram-positive bacteria (*Firmicutes*). For the most part, it is restricted to the family *Bacillaceae* and the class *Clostridia*, but members of the less well-studied family *Veillonellaceae* (e.g., *Acetonea longum* [5, 6]) also produce endospores. The last common ancestor of *Clostridium* and *Bacillus* predates the initial rise of oxygen in the atmosphere (7) approximately 2.3 billion years ago, and yet, remarkably, orthologs of signature sporulation genes are shared between the genomes of these distantly related bacteria (2, 8). We wondered whether gene conservation among endospore formers could be exploited to discover previously unrecognized genes involved in sporulation.

Sporulation has been most extensively studied in the model organism *Bacillus subtilis*. Entry into sporulation is governed by the master regulator Spo0A, which is activated by phosphorylation through a multicomponent signal transduction pathway (9). Phosphorylated Spo0A (Spo0A~P) directly regulates (activates or represses) the expression of 121 genes (10) and significantly influences the expression of over 500 genes (11). Sporulation is initiated when Spo0A~P levels reach a threshold (12). Sporulating cells undergo several successive morphological changes, a hallmark of which is the formation of a two-compartment sporangium consisting of forespore and mother cell compartments. As development proceeds, the forespore is wholly engulfed by the mother cell to create a cell within a cell. The inner cell becomes the dormant spore and is released from the mother cell by lysis (2). Upon release, the mature spore can remain dormant for long periods or, in response to germinants, give rise to a vegetative cell. The developmental program of sporulation is governed in part by

the successive actions of four compartment-specific sigma factors (appearing in the order σ^F , σ^E , σ^G , and σ^K), whose activities are confined to the forespore (σ^F and σ^G) or the mother cell (σ^E and σ^K) (13).

Traditional approaches of forward genetics have identified many, if not all, genes that are essential for sporulation (*spo*). These approaches rely on conspicuous phenotypes for the identification of target genes. Complementary approaches, such as the identification of sporulation-specific proteins (e.g., SASP and coat proteins) and transcriptome analysis (14, 15), revealed additional genes under sporulation control, including genes that contribute to efficient sporulation and spore resistance properties. Here, we report the application of phylogenetic profiling in an effort to discover additional genes involved in sporulation that might have gone undetected in previous approaches. We therefore sought to identify previously unrecognized genes involved in sporulation in *B. subtilis* on the basis of their conservation among endospore-forming bacteria. Using phylogenetic profile analysis for the initial identification and transcriptional and mutational analyses, we discovered previously overlooked genes under sporulation control, including two genes whose mutants caused a small but detectable developmental defect.

MATERIALS AND METHODS

Bacterial strains and culturing conditions. *Escherichia coli* strain DH5 α was used for propagating plasmids and was grown and transformed using standard procedures (16). The *B. subtilis* strains used in this work are

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TABLE 1 Strains used in this study

Strain ^a	Genotype	Source or reference
PY79	Prototrophic derivative of <i>B. subtilis</i> subsp. <i>subtilis</i> 168	20
RL5360	<i>amyE::P_{hyperspank}-lacZ spc</i>	This study
RL5361	<i>bkdR::spc</i>	This study
RL5362	<i>bkdR::erm</i>	This study
RL5363	<i>buk::erm</i>	This study
RL5364	<i>ylmC::erm</i>	This study
RL5365	<i>ylmC::erm amyE::ylmC spc</i>	This study
RL5366	<i>ymxH::spc</i>	This study
RL5367	<i>ylxY::spc</i>	This study
RL5368	<i>ylxY::spc amyE::ylxY cat</i>	This study
RL5369	<i>ymfB::spc</i>	This study
RL5370	<i>yteA::erm</i>	This study
RL5371	<i>ylyA::erm</i>	This study
RL5372	<i>ylyA::erm sacA::ylyA kan</i>	This study
RL5373	<i>yocK::tet</i>	This study
RL5374	<i>amyE::P_{bkdR}-lacZ cam</i>	This study
RL5375	<i>amyE::P_{bkd}-lacZ cam</i>	This study
RL5376	<i>spo0A::spc amyE::P_{bkd}-lacZ cam</i>	This study
RL5377	<i>bkdR::erm amyE::P_{bkd}-lacZ cam</i>	This study
RL5378	<i>spo0A::spc bkdR::erm amyE::P_{bkd}-lacZ cam</i>	This study
RL5379	<i>amyE::P_{ylmC}-lacZ cam</i>	This study
RL5380	<i>sigF::kan amyE::P_{ylmC}-lacZ cam</i>	This study
RL5381	<i>sigE::erm amyE::P_{ylmC}-lacZ cam</i>	This study
RL5382	<i>sigG::kan amyE::P_{ylmC}-lacZ cam</i>	This study
RL5383	<i>spo0A::spc amyE::P_{ymxH}-lacZ cam</i>	This study
RL5384	<i>sigF::kan amyE::P_{ymxH}-lacZ cam</i>	This study
RL5385	<i>amyE::P_{ylxY}-lacZ cam</i>	This study
RL5386	<i>sigF::kan amyE::P_{ylxY}-lacZ cam</i>	This study
RL5387	<i>sigE::erm amyE::P_{ylxY}-lacZ cam</i>	This study
RL5388	<i>sigG::kan amyE::P_{ylxY}-lacZ cam</i>	This study
RL5389	<i>amyE::P_{ylyA}-lacZ cam</i>	This study
RL5390	<i>amyE::P_{ymfB}-lacZ cam</i>	This study
RL5391	<i>sigF::kan amyE::P_{ymfB}-lacZ cam</i>	This study
RL5392	<i>sigG::kan amyE::P_{ymfB}-lacZ cam</i>	This study
RL5393	<i>spoVT::spc amyE::P_{ymfB}-lacZ cam</i>	This study
RL5394	<i>amyE::P_{yteA}-lacZ cam</i>	This study
RL5395	<i>sigG::kan amyE::P_{yteA}-lacZ cam</i>	This study
RL5396	<i>spoVT::spc amyE::P_{yteA}-lacZ cam</i>	This study
RL5397	<i>amyE::P_{ylyA}-lacZ cam</i>	This study
RL5398	<i>sigG::kan amyE::P_{ylyA}-lacZ cam</i>	This study
RL5399	<i>spoVT::spc amyE::P_{ylyA}-lacZ cam</i>	This study
RL2242	<i>spo0A::spc</i>	11
RL1265	<i>sigF::kan</i>	17
RL1061	<i>sigE::erm</i>	19
RL4962	<i>sigG::kan</i>	18
RL3873	<i>spoVT::spc</i>	15

^a All strains are isogenic with PY79 unless otherwise indicated.

listed in Table 1. All strains were derived from the prototrophic laboratory strain PY79 (20). Transformation of *Bacillus* strains with double-stranded PCR fragments, or plasmid or genomic DNA, was done as previously described (21). Sporulation was induced by exhaustion in Difco sporulation (DS) medium or, in the case of β -galactosidase activity assays (see below), by resuspension in Sterlini-Mandelstam (SM) medium (22).

Plasmid construction. The oligonucleotides used for PCR in this study are listed in Table 2. Fragments of the upstream regions of candidate genes, including the respective ribosome binding sites (RBS) and start codons, were amplified by PCR and cloned into pAH124 (23) using the appropriate restriction sites (Table 2). In this way, *lacZ* reporter constructs with the start codon of the gene fused directly to the *lacZ* gene were obtained. The constructs were introduced into PY79 by transformation.

TABLE 2 Oligonucleotides used to construct reporter and complementation constructs

Primer ^a	Sequence (5'–3')
bkdR –296E	CTGGAATTTCGATGAATCCTGACAACCCTTG
bkdR +3H	CTGAAGCTTCATCCCGATACCCCTTTGTAT
Bkd –298E	CTGGAATTCGAAGCGGAAAAGCTGTCTGT
Bkd +3H	CTGAAGCTTCATCTGTTACCACCTTTCTTG
ylmC –300E	CTGGAATTCAGTGAAACGGGAGTGTCCA
ylmC +3H	CTGAAGCTTCATCCATCACGTCCTTTTTC
ylmC +362B	CTGAGGATCCCTTATTTTACCACATCTTACTG
ymxH –282E	CTGAGAATCCATGTACCCTTAGAAAGCAC
ymxH +3H	CTGAAAGCTTCATGTCTGTCAACCCCTTG
ylxY –330E	CTGGAATTCCTCGGGGCTTTCGTTGAAATT
ylxY +3E	CTGGAATTCATGTTCTGTCCCCCTCAC
ylxY +1076B	CTGAGGATCCATCGCAACAGAACGGACTGTC
ylzA –380E	CTGGAATTCATCAAAGAATGGACTGAAGACG
ylzA +3E	CTGGAATTCATCTTCTACGTTCCCTCTGT
ymfB –234E	CTGGAATTCAAACATCAAATGTCGAAATGGTC
ymfB +3H	CTGAAGCTTCATAATGCTGTCTTCCGATC
yteA –351E	CTGGAATTCCTGGCTTTATGTAATGCATGTAG
yteA +3E	CTGGAATTCATGTGATCGCCTCGTTTCT
ylyA –613E	CTGGAATTCGTTGCTCATTAAACATTTGTTG
ylyA +3E	CTGGAATTCATCTTCCAACTCCCTGCTC
ylyA +514B	CTGGATCCCTGCAATAAAGTAGTGCAATC

^a The numbers refer to the 5' nucleotide position relative to the first nucleotide of the start codon (+1) of the respective gene. B, BamHI; E, EcoRI; H, HindIII.

Complementation constructs for the *ylmC*, *ylxY*, and *ylyA* mutants were made by PCR amplification of fragments carrying the gene promoters and entire open reading frames (ORF). These PCR fragments were cloned, using the appropriate restriction enzymes, into pDG1662 (24), in the case of *ylmC* and *ylxY*, or pSac-Cam (25), in the case of *ylyA*. The constructs were introduced into the respective mutant strains by transformation.

Phylogenetic profile analysis. For each predicted gene product in *B. subtilis* subsp. *subtilis* 168, its presence or absence in 626 complete archaeal and bacterial genomes that were available at the time of the initial phylogenetic analysis was determined by asking whether a putative ortholog was present (1) or absent (0) in that species, similar to previous descriptions (26). This analysis included 46 genomes of endospore-forming bacteria belonging to the family *Bacillaceae* and the class *Clostridia*, namely, *Bacillus amyloliquefaciens* FZB42, *Bacillus anthracis* strain Ames, *B. anthracis* strain Ames Ancestor, *B. anthracis* strain Sterne, *Bacillus cereus* ATCC 10987, *B. cereus* ATCC 14579, *B. cereus* E33L, *B. cereus* subsp. *cytotoxicus* NVH 391-98, *Bacillus clausii* KSM-K16, *Bacillus halodurans* C-125, *Bacillus licheniformis* ATCC 14580, *Bacillus pumilus* SAFR-032, *Bacillus thuringiensis* serovar konkukian strain 97-27, *B. thuringiensis* strain AI Hakam, *Bacillus weihenstephanensis* KBAB4, *Geobacillus kaustophilus* HTA426, *Geobacillus thermodenitrificans* NG80-2, *Oceanobacillus ihelyensis* HTE831, *Alkaliphilus metalliredigens* QYMF, *Alkaliphilus oremlandii* OhILAs, *Caldicellulosiruptor saccharolyticus* DSM 8903, *Carboxydotherrmus hydrogeniformans* Z-2901, *Clostridium acetobutylicum* ATCC 824, *Clostridium beijerinckii* NCIMB 8052, *Clostridium botulinum* A strain ATCC 3502, *C. botulinum* A strain ATCC 19397, *C. botulinum* A strain Hall, *C. botulinum* F strain Langeland, *Clostridium difficile* 630, *Clostridium kluyveri* DSM 555, *Clostridium perfringens* ATCC 13124, *C. perfringens* SM101, *C. perfringens* strain 13, *Clostridium phytofermentans* ISDg, *Clostridium tetani* E88, *Clostridium thermocellum* ATCC 27405, *Desulfotobacterium hafniense* Y51, *Desulfotomaculum reducens* MI-1, *Moorella thermoacetica* ATCC 39073, *Symbiobacterium thermophilum* IAM 14863, *Syntrophomonas wolfei* subsp. *wolfei* strain Goettingen, *Thermoanaerobacter pseudethanolicus* ATCC 33223, *Thermoanaerobacter tengcongensis* MB4, *Pelotomaculum thermopropionicum* SI, and *Thermoanaerobacter* sp. X514. Proteins were then grouped by their distribution patterns across species.

In this way, a set of 58 genes, many of which are signature sporulation genes (see below) and all of which are highly enriched among endospore-forming species, was obtained.

β -Galactosidase activity assays. Samples were collected from shaking cultures in duplicate at various time points after sporulation was induced by resuspension in SM medium. β -Galactosidase activity was measured in a Synergy 2 plate reader (BioTek), as previously described (23). The experiment was repeated to ensure reproducibility. β -Galactosidase activity is reported in arbitrary units (AU) as the rate of *o*-nitrophenyl- β -D-galactopyranoside (ONPG) conversion (i.e., V_{max} , with units of optical density at 420 nm [OD₄₂₀] per minute) divided by the OD₆₀₀ of the sample at the time of collection, as previously described (23).

Competition experiments. A wild-type reference strain was competed against strains mutant for candidate genes, as previously described (27), with a few differences. Typically, starting from overnight LB cultures, the wild-type reference strain (RL5360), which carried an isopropyl β -D-1-thiogalactopyranoside (IPTG)-inducible *lacZ*, was mixed with an excess mutant strain in 5 ml DS medium. In competition assays using *ylmC* mutants, which displayed an advantage under sporulation-inducing conditions, we started with an excess of the wild-type strain. Cultures were grown and allowed to sporulate at 37°C in DS medium for 24 h. The cultures were then heat treated at 80°C for 20 min, briefly cooled at room temperature, and diluted in 5 ml fresh DS medium. At appropriate intervals, dilutions were plated on agar plates containing 0.008% (wt/vol) 5-bromo-4-chloro-3-indolyl- β -D-galactopyranoside (X-Gal) and 1 mM IPTG. The blue and white colonies, reflecting the ratio of wild-type reference to mutant strains, respectively, were counted. Similar competition experiments were done in LB to establish if observed phenotypes were specific to sporulation-inducing conditions. In these experiments, we grew the cocultures at 25°C to minimize the time the culture was in stationary phase. In short, wild-type reference and mutant strains were grown at 25°C in LB for 24 h and subsequently diluted in 5 ml fresh LB for another round of growth. The heat treatment step was omitted in these experiments.

Germination of purified spores. Spores were purified essentially as previously described (22). In short, the strains were induced to sporulate by growing in DS medium for 48 to 72 h. Cells were pelleted by centrifugation, and the pellet was resuspended in ice-cold water. The cells were washed twice with ice-cold water and stored overnight at 4°C. The next day, the cells were pelleted by centrifugation and washed another 6 to 8 times with ice-cold water. In this way, preparations that were more than 95% phase-bright spores, as judged by phase-contrast microscopy, were obtained.

For germination assays, spores at an OD₆₀₀ of 10 were activated by heat treatment at 80°C for 20 min and cooled on ice for 2 min. Activated spores were germinated at an OD₆₀₀ of 0.5 in LB medium. Germination was recorded as a loss of optical density in a Synergy 2 plate reader (BioTekin).

RESULTS

Candidates for uncharacterized sporulation genes. We used phylogenetic profile analysis to search for genes in the genome of *B. subtilis* subsp. *subtilis* 168 (here simply referred to as *B. subtilis*) that are specifically conserved among endospore-forming, low-G+C, Gram-positive bacteria. A similar method was previously used to identify signature sporulation genes in the thermophilic firmicute *Carboxydotherrmus hydrogenoformans* (26). Phylogenetic profiling works by grouping genes according to their distribution patterns in different species. For each predicted gene product in *B. subtilis*, its presence or absence in all complete archaeal and bacterial genomes available at the time of this analysis (626 genomes in total) was determined by asking whether an ortholog was present in that species. This analysis included 46 genomes of endospore-forming bacteria belonging to the family *Bacillaceae* and the class *Clostridia*. The orthologs were then grouped by their

TABLE 3 Candidates for uncharacterized sporulation genes among genes conserved in endospore-forming bacteria

Gene	No. of orthologs in:		Predicted product
	All bacterial/archaea ^a	<i>Bacillaceae</i> / <i>Clostridia</i> ^b	
<i>bkdR</i>	87	43	DNA-binding transcriptional regulator
<i>buk</i>	62	34	Butyrate kinase
<i>ylmC</i>	45	45	Hypothetical protein; PRC barrel domain
<i>ymxH</i>	25	25	Hypothetical protein; PRC barrel domain
<i>ylxY</i>	42	38	Polysaccharide deacetylase
<i>ylzA</i>	73	43	Hypothetical protein
<i>ymfB</i>	44	44	ClpP-like protease
<i>yteA</i>	51 (33) ^c	41 (23)	DksA-like regulator
<i>ylYA</i>	11 (29)	11 (29)	DksA-like regulator

^a 626 bacterial and archaeal genomes were considered.

^b 46 *Bacillus* and *Clostridia* genomes were considered.

^c The numbers in parentheses are the numbers of orthologs for *yteA* and *ylYA* corrected for similarity in gene synteny with *B. subtilis ylyA*.

distribution patterns across species. In this way, 58 genes that were highly and specifically conserved among endospore-forming bacteria were identified (see Fig S1 in the supplemental material). Many of these are signature sporulation genes with well-studied roles in spore formation, such as *spoIIR* and *spoIIGA*, which mediate the activation of the mother-cell-specific transcription factor σ^E (28); *spoIID*, *spoIIM*, and *spoIIP*, required for forespore engulfment by the mother cell (29–31); and *spoIVA*, which encodes a morphogenetic protein required for coat assembly (32). Five genes (i.e., *bkdR*, *ylmC*, *ymxH*, *ylzA*, [formerly designated *remA*], and *ymfB* [formerly designated *tepA*]), however, had no previously documented role in spore formation, and an additional three genes (i.e., *ylxY*, *yteA*, and *ylYA*) had previously been shown to be under sporulation control, but no roles in sporulation had been described (15, 33). We hypothesized that these genes might play previously unrecognized or overlooked roles in sporulation.

Six of the eight genes were widely conserved among the 46 endospore-forming *Bacillaceae* and *Clostridia* genomes examined. In the cases of *ymxH* and *ylYA*, however, only 25 and 11 orthologs, respectively, were identified. *ymxH* orthologs are present in almost all endospore-forming *Bacillaceae* species but are missing from the majority of endospore-forming *Clostridia* species. Nonetheless, *ymxH* exhibits significant sequence similarity to one of the other genes on our list, *ylmC*, which is abundant among endospore-forming bacteria. Meanwhile, a close look at *ylYA*, which is homologous to *yteA*, revealed that several genes identified as *yteA* orthologs in endospore-forming bacteria share similar gene synteny with *B. subtilis ylyA*. *B. subtilis ylyA* is flanked by several well-characterized genes, including *divIVA*, encoding a cell division protein; *ileS*, encoding an isoleucyl-tRNA synthetase; *lspA*, encoding a type II signal peptidase; and *rluD*, encoding a pseudouridylylate synthase, many of which are found in the vicinity of *ylYA* orthologs in *Bacillaceae* and *Clostridia*. This suggests that these genes are actually *ylYA* orthologs, raising the number of orthologs to 29 (Table 3). Thus, all eight genes are widely conserved among endospore-forming bacteria or are homologous to genes that are.

Finally, two additional genes, *yvjA* and *buk*, that had orthologs both in endospore-forming bacteria and in all four species of *Lis-*

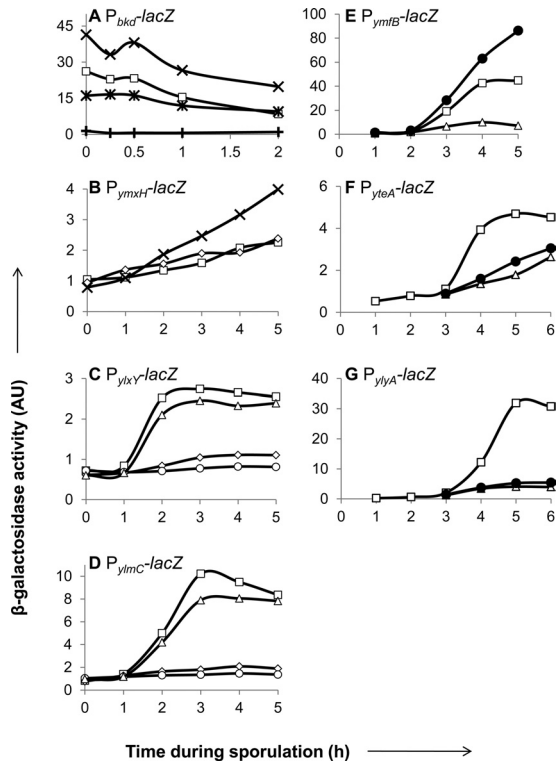


FIG 1 Transcription and regulatory-gene dependence of candidate sporulation genes. Strains carrying *lacZ* transcriptional reporter constructs for *bkd* (operon including *buk*) (A), *ymxH* (B), *ylxY* (C), *ylmC* (D), *ymfB* (E), *yteA* (F), and *ylyA* (G) were induced to sporulate, and β -galactosidase activity was monitored for samples taken at the indicated time points after sporulation induction. Activity (AU) was assayed in the wild type (\square) or strains mutant for *spo0A* (\times), *sigF* (\diamond), *sigE* (\circ), *sigG* (\triangle), *spoVT* (\bullet), *bkdR* ($+$), or *spo0A* and *bkdR* (*).

tertia included in this analysis were identified (see Fig S1 in the supplemental material). *Listeria* species are closely related to *B. subtilis* but do not form spores and lack almost all signature sporulation genes (34). Because of their presence in non-endospore-forming bacteria, we considered *yyjA* and *buk* unlikely candidates for unrecognized sporulation genes. Nonetheless, as a control, we retained one of these genes, *buk*, in our analysis. Thus, a total of nine genes were carried forward for further investigation.

Transcription under sporulation-inducing conditions. We next asked if the nine candidate genes are transcribed under conditions that induce sporulation. For this purpose, we built transcriptional reporter constructs, typically cloning a 300- to 400-bp fragment directly upstream of the gene and fusing its start codon to that of the *lacZ* gene. In the case of *buk*, which is the third gene in a seven-gene (*bkd*) operon that is involved in branched-chain amino acid utilization (35), we instead cloned an approximately 300-bp fragment upstream of *ptb*, the first gene of the operon. *B. subtilis* strains carrying these constructs integrated at the *amyE* locus were induced to sporulate by resuspension in SM medium (22), and samples taken at various times were analyzed for β -galactosidase activity. Seven of the nine reporters were expressed during sporulation, six of which were induced at various times after the induction of sporulation (Fig. 1). We did not observe activity for the *bkdR* and *ylzA* reporters under the tested conditions (data not shown).

The *bkd* reporter was active from the time of the induction of sporulation, decreasing only slightly during the time it was monitored (Fig. 1A). As previously reported, expression of the *bkd* operon depends on the alternative sigma factor σ^L and BkdR (35). Indeed, in a *bkdR* mutant, expression was abolished (Fig. 1A). We next measured expression in a strain mutant for *spo0A*. The *spo0A* gene encodes the master regulator for entry into sporulation, Spo0A, which is active in its phosphorylated form, Spo0A~P (2, 9). Activity was approximately 2-fold higher than that of the wild type at the times tested. Interestingly, in a strain doubly mutant for *spo0A* and *bkdR*, expression levels were similar to that of the wild type (Fig. 1A), showing that in the absence of Spo0A, BkdR is not required for expression. That is, a *spo0A* mutation is epistatic to a *bkdR* mutation. This suggests that BkdR antagonizes Spo0A~P to activate transcription from the *bkd* operon promoter. We found a potential Spo0A binding site (GTCGAAA [see Fig S2 in the supplemental material]) with high similarity to the consensus binding sequence (TTTGTCGAAA [10]) located immediately downstream of the σ^L -dependent transcriptional start site (35). Just upstream of the promoter are tandem sequences previously shown to be important for BkdR-mediated activation (35). *In toto*, these observations suggest that the binding of BkdR upstream of the promoter overcomes the repressive effect of the binding of Spo0A~P just downstream of the start site.

The *ymxH* reporter was expressed from an early time, increasing slightly but constantly during the first 5 h (Fig. 1B). We tested the activity in strains mutant for *spo0A* and *sigF*, which encodes the first forespore-specific sigma factor, σ^F (2). Expression in a *spo0A* mutant was upregulated, steadily increasing from 1 h after sporulation was induced. In contrast, activity in a *sigF* mutant was unchanged compared to that of the wild type (Fig. 1B). These findings suggest that *ymxH* is directly or indirectly under the negative control of Spo0A~P but is not otherwise under sporulation control.

Expression from the *ylxY* and *ylmC* reporters was induced between hours 1 and 2 of sporulation (Fig. 1C and D). Previous work indicated that *ylxY* expression is σ^E dependent and under the negative control of the mother-cell-specific regulator SpoIIID (14, 33). We tested the activity of the *ylxY* and *ylmC* reporters in strains mutant for *sigF*, *sigE* (which encodes the mother-cell-specific sigma factor σ^E), or *sigG* (which encodes the late-appearing, forespore-specific sigma factor σ^G). Activity for both was abolished in the *sigF* and *sigE* mutants but reached wild-type levels in the *sigG* mutant (Fig. 1C and D). Thus, these genes are transcribed in a σ^E -dependent manner.

Finally, three reporters were induced between 2 and 3 h after sporulation was induced, namely, those for *ymfB*, *yteA*, and *ylyA* (Fig. 1E to G). Previous transcriptome analyses indicated that *yteA* and *ylyA* are indeed part of the σ^G regulon (15), whereas *ymfB* was not known to be under sporulation control. The activities of all three reporters were abolished in a *sigG* mutant (Fig. 1E-G). In addition, *ylyA* was previously shown to be under the control of SpoVT, a modulator of σ^G -dependent transcription (15, 36). We tested the activities of all three reporters in a strain mutant for *spoVT*. The activities of the *yteA* and *ylyA* reporters were markedly reduced in a *spoVT* mutant, whereas the activity of the *ymfB* reporter increased in a *spoVT* mutant (Fig. 1E to G).

In summary, we conclude that five of the nine genes in our investigation are under sporulation control, with two, namely,

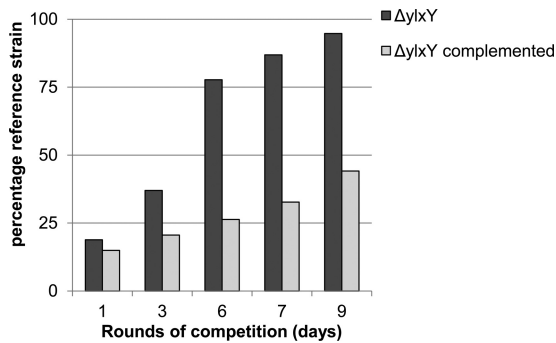


FIG 2 A *ylxY* mutant exhibits a competition deficit under sporulation-inducing conditions. A wild-type reference strain carrying an IPTG-inducible *lacZ* gene (RL5360) was competed in DS medium against a *ylxY* mutant, starting with approximately 20% wild-type strain. Cultures were incubated at 37°C for 24 h, heat treated at 80°C, and diluted in fresh DS medium. After the indicated rounds of competition, dilutions of the culture were plated on agar plates containing IPTG and X-Gal, and blue (wild type) and white (mutant) colonies were counted. The bars indicate the percentages of the wild-type reference strain. The wild-type strain was competed against a *ylxY* mutant and a complemented strain carrying a copy of *ylxY* at the ectopic *amyE* locus.

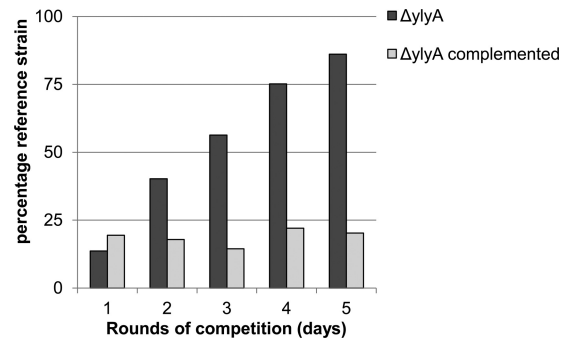


FIG 3 A *ylyA* mutant exhibits a competition deficit under sporulation-inducing conditions. A wild-type reference strain carrying an IPTG-inducible *lacZ* gene (RL5360) was competed in DS medium against a *ylyA* mutant, starting with approximately 20% wild-type strain. Cultures were incubated at 37°C for 24 h, heat treated at 80°C, and diluted in fresh DS medium. After the indicated rounds of competition, dilutions of the culture were plated on agar plates containing IPTG and X-Gal, and blue (wild-type) and white (mutant) colonies were counted. The bars indicate the percentages of the wild-type reference strain. The wild-type strain was competed against a *ylyA* mutant and a complemented strain carrying a copy of *ylyA* at the ectopic *sacA* locus.

ylxY and *ylmC*, under the control of σ^E and three, *ymfB*, *yteA*, and *ylyA*, under the control of σ^G .

Competition-based analysis of candidate gene mutants. We constructed mutant strains for eight of the nine genes by deleting and replacing their ORF with antibiotic resistance cassettes. We were unable to obtain a mutant for the ninth gene, *ylzA*. Previously, others obtained transposon insertions directly upstream of *ylzA*; however, no report was made of transposon insertions internal to the *ylzA* coding sequence (37).

None of the eight mutants had a conspicuous phenotype, as judged by colony morphology or spore formation (data not shown). Thus, if any of these genes represent previously uncharacterized sporulation genes, their contributions to spore formation must be subtle. To test for such a subtle role, we carried out competition experiments in which mutant strains were competed for several rounds of sporulation against a wild-type reference strain marked by an IPTG-inducible *lacZ* gene. Typically, mutant cells were severalfold in excess of the wild type at the start of the experiment. Cocultures of the mutant and wild type were grown and allowed to sporulate at 37°C in DS medium for 24 h. The culture was then heat treated at 80°C and diluted in fresh DS medium for another round of sporulation. At appropriate intervals, dilutions were plated on agar plates containing X-Gal and IPTG, and blue (wild-type) and white (mutant) colonies were counted. Mutant strains that were outcompeted (or in one case slightly undercompeted) by the wild type were next subjected to competition experiments in growth medium to determine if the observed competition phenotype was an indirect consequence of a growth defect rather than a defect in sporulation. In these experiments, cocultures were grown at 25°C in LB medium for 24 h and subsequently diluted in fresh LB medium for another round of growth. The heat treatment step was omitted in these experiments.

The mutant strains fell into three categories: those that did not have a competition phenotype under sporulation-inducing conditions, those that had a phenotype under sporulation-inducing but not under growth conditions, and lastly, those that had a phenotype under both conditions. The first category comprised the

bkdR, *ymxH*, *ymfB*, and *yteA* mutants. In a competition experiment in which the wild-type reference strain was competed against an unmarked wild-type strain, the percentage of the wild-type reference strain remained constant during several rounds of competition (see Fig S3 in the supplemental material), indicating that the inducible *lacZ* construct did not affect the fitness of the reference strain. Similarly, in competition experiments with strains mutant for *bkdR*, *ymxH*, *ymfB*, or *yteA*, the percentage of the wild-type reference strain remained constant (see Fig S3 in the supplemental material).

The second category comprised the *ylxY*, *ylyA*, and *ylmC* mutants. Strains mutant for *ylxY* and *ylyA* had clear competition deficits under sporulation-inducing conditions. Over the course of nine and five rounds of competition, respectively, the percentage of wild-type reference strain increased from approximately 20% to 90% of the population (Fig. 2 and 3). Genetic complementation by reintroducing a copy of the respective gene at an ectopic locus restored competitiveness to the mutant strains (Fig. 2 and 3), confirming that the observed phenotypes resulted from deletion of the genes. In contrast to the apparent deficit of *ylxY* and *ylyA* mutants, the *ylmC* mutant exhibited a slight competitive advantage under sporulation-inducing conditions (Fig. 4). As before, genetic complementation by reintroducing a copy of *ylmC* at the ectopic *amyE* locus reversed this phenotype (Fig. 4). All three competition phenotypes were found to be specific to competition experiments under sporulation-inducing conditions, because no changes from the starting ratio were observed in competition experiments during growth (Fig. 5).

Finally, the third category contained only one strain, the *buk* mutant, which had a clear competition deficit under sporulation-inducing and growth conditions (see Fig S4 in the supplemental material).

A *ylyA* mutant is defective in germination. We do not know the precise step in sporulation at which the *ylxY* mutation impedes spore formation or the step at which the *ylmC* mutation confers a competitive advantage. However, in the case of *ylyA*, we can, at least in part, attribute the competitive disadvantage to impaired spore germination. As shown in Fig. 6, spores from a *ylyA* mutant

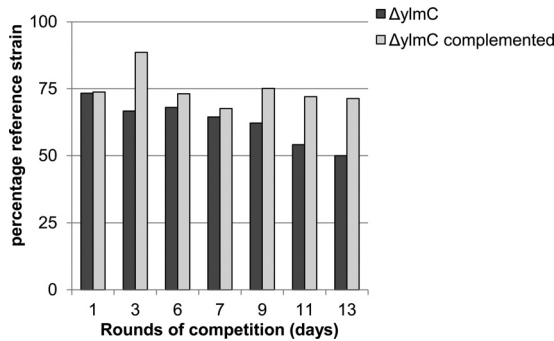


FIG 4 A *yImC* mutant exhibits a slight competitive advantage under sporulation-inducing conditions. A wild-type reference strain carrying an IPTG-inducible *lacZ* gene (RL5360) was competed in DS medium against a *yImC* mutant, starting with approximately 75% wild-type strain. The cultures were incubated at 37°C for 24 h, heat treated at 80°C, and diluted in fresh DS medium. After the indicated rounds of competition, dilutions of the culture were plated on agar plates containing IPTG and X-Gal, and blue (wild-type) and white (mutant) colonies were counted. The bars indicate the percentages of the wild-type reference strain. The wild-type strain was competed against a *yImC* mutant and a complemented strain carrying a copy of *yImC* at the ectopic *amyE* locus.

are slower to germinate than either the wild-type parent or a *ylyA* mutant strain harboring a wild-type copy of the gene at the ectopic *sacA* locus.

DISCUSSION

Using phylogenetic profiling, we identified eight genes (i.e., *bkdR*, *yImC*, *ymxH*, *ylxY*, *ylzA*, *ymfB*, *yteA*, and *ylyA*) that are widely conserved among endospore-forming species of *Bacillaceae* and *Clostridia* but were not previously reported to be involved in sporulation or, in the cases of *ylxY*, *yteA*, and *ylyA*, not well characterized. *ymfB*, whose product is homologous to ClpP-like proteases (Table 3), was previously suggested to be involved in translocation and processing of the α -amylase AmyQ (reference 38, where *ymfB* was named *tepA*). Researchers from the same laboratory, however, later reported that they were unable to replicate the initial results with a clean knockout of *ymfB* (39). *ylzA*, which encodes a hypothetical protein with no clear homology to

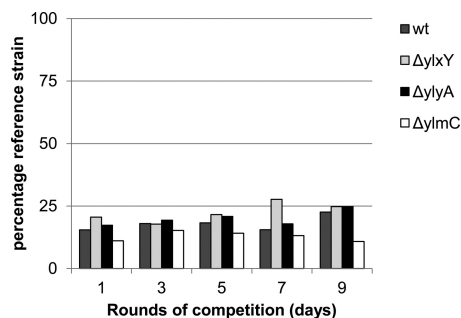


FIG 5 *ylxY*, *ylyA*, and *yImC* mutants do not exhibit a significant competitive deficit during growth. A wild-type reference strain carrying an IPTG-inducible *lacZ* gene (RL5360) was competed in LB medium against an unmarked wild type (wt) and strains mutant for *ylxY*, *ylyA*, and *yImC*, starting with approximately 20% wild-type strain. The cultures were incubated at 25°C for 24 h and diluted in fresh LB. After the indicated rounds of competition, dilutions of the culture were plated on agar plates containing IPTG and X-Gal, and blue and white colonies were counted. The bars indicate the percentages of the wild-type reference strain.

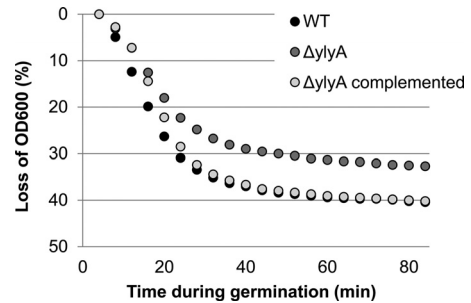


FIG 6 A *ylyA* mutant is delayed in germination. Purified spores were heat treated and cooled on ice. Activated spores were germinated at an OD₆₀₀ of 0.5 by dilution in LB medium. Every 4 min after germination induction, the OD₆₀₀ was measured, and germination is reported as the percent drop in optical density.

known proteins (Table 3), was previously shown to be involved in the regulation of extracellular-matrix components during biofilm formation in *B. subtilis* (reference 37, where *ylzA* is designated *remA*). Five of the eight genes were found to be under the control of sporulation-specific transcription factors, with *yImC* and *ylxY* under the control of the mother-cell-specific sigma factor σ^E and *ymfB*, *yteA*, and *ylyA* under the control of the forespore-specific factors σ^G and SpoVT. Inactivation of *ylxY*, *ylyA*, and *yImC* resulted in measurable changes in competitiveness under sporulation-inducing conditions, but not under growth conditions.

We included *buk* as a control in our investigation. Like the other eight genes, orthologs of *buk* are enriched among endospore-forming bacteria but are also found in some non-endospore-forming species, most notably *Listeria*, a close relative of *B. subtilis* that is asporogenic. The *buk* gene, which codes for a butyrate kinase, is part of a seven-gene operon in *B. subtilis* that is involved in the utilization of branched-chain amino acids as a nitrogen source (35). Orthologs of other members of the operon are widespread among bacteria, much more so than *buk* itself. In *B. subtilis* the operon is under the control of the alternative sigma factor σ^L and the transcription activator BkdR (35), which, as we have shown, is itself highly conserved among endospore-forming bacteria. Inactivation of *buk* resulted in defects that were not specific to sporulation, and the *bkd* operon promoter was constitutively active under the tested conditions. Interestingly, we found a putative Spo0A binding site adjacent to the predicted transcriptional start site, and our results indicate that BkdR antagonizes Spo0A~P to activate expression. The presence of BkdR orthologs in almost all endospore-forming species analyzed (Table 3), but not *Listeria* species, supports the idea that BkdR is conserved among endospore-forming bacteria to counteract the effect of Spo0A~P on transcription.

yImC and *ymxH* are paralogs that code for small hypothetical proteins that resemble a motif known as a photosynthetic reaction center (PRC) beta-barrel domain (Table 3). The PRC domain, which is itself widespread among photosynthetic and nonphotosynthetic bacteria, archaea, and plants, is thought to mediate protein-protein interactions (40). Deletion of *yImC* resulted in a strain that had a slight competitive advantage over the wild type under sporulation-inducing conditions. It is unclear what causes this unexpected phenotype. A double mutant of *yImC* and *ymxH* had essentially the same phenotype as the *yImC* single mutant (data not shown). We infer that YImC must confer some fitness

advantage during spore formation under unknown environmental conditions (e.g., a spore resistance property) but that production of the protein evidently imposes a slight cost on spore formation that impedes development.

Deletion of *ylxY* and *ylaA* resulted in clear competitive deficits specific to sporulation-inducing conditions. *ylxY* encodes a probable polysaccharide deacetylase (Table 3) and exhibits some similarity to two *B. subtilis* genes under sporulation control, namely, *pdaA* and *pdaB*, mutants of which display defects in spore cortex maturation (41–43). The nature of the competition deficit of the *ylxY* mutant is currently unknown. In contrast, we have determined that the competition defect of the *ylaA* mutant stems, not from a defect in spore formation *per se*, but rather from impaired germination of the mutant spores. Krasny and Gourse previously reported that YlyA and YteA share some sequence similarity to the transcription factor DksA (44). DksA inhibits the transcription of rRNA genes by direct interaction with RNA polymerase (45). Mutation of *ylaA* or *yteA*, however, seemingly did not affect the activity of the P1 promoter of the rRNA gene *rrnB* under any of the tested growth conditions (44). It will be interesting to see whether YlyA similarly modifies RNA polymerase activity and, if so, whether this modification influences the expression of genes involved in spore germination.

Our results reinforce the view that phylogenetic profiling, in combination with reverse genetics and gene-regulatory studies, can be a powerful tool for the discovery of genes that play a subtle role in a complex developmental process and whose contributions might otherwise be overlooked by traditional approaches of forward genetics.

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