The master regulator for entry into sporulation in *Bacillus subtilis* becomes a cell-specific transcription factor after asymmetric division

Masaya Fujita and Richard Losick

Department of Molecular and Cellular Biology, Harvard University, Cambridge, Massachusetts 02138, USA

Gene transcription at the onset of sporulation in *Bacillus subtilis* is governed by Spo0A, a member of the response regulator family of transcription factors. Spo0A is traditionally viewed as the master regulator for entry into development. We now report that Spo0A continues to function after the initiation phase of sporulation and that it becomes a cell-specific transcription factor when the sporangium is divided into a mother cell and forespore. We observed that (1) Spo0A and Spo0A-directed gene transcription reached high levels in the mother cell; (2) an activated form of Spo0A impaired sporulation when produced in the forespore but not when produced in the mother cell; and (3) an inhibitor of Spo0A called Spo0A-N impaired sporulation and Spo0A-directed transcription when produced in the mother cell but not when produced in the forespore. Spo0A-N, which corresponds to the NH2-terminal domain of Spo0A, was shown to compete with the full-length response regulator for phosphorylation by the phosphorelay protein Spo0B. We propose that Spo0A is the earliest-acting transcription factor in the mother-cell line of gene expression and that in terms of abundance and transcriptional activity Spo0A may function predominantly as a cell-specific regulatory protein.

*Keywords: Bacillus subtilis; sporulation; response regulator; transcription; asymmetry*

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Sporulation in the bacterium *Bacillus subtilis* is governed by a series of transcription factors that are subject to temporal and spatial regulation (Stragier and Losick 1996; Piggot and Losick 2002). The initiation phase of development is controlled by Spo0A, which is traditionally considered to be the master regulator for entry into the sporulation pathway (Hoch 1993). Spo0A is a member of the response regulator family of transcription factors (Hoch 2000). The activity of Spo0A is determined by phosphorylation, which is mediated by a phosphorelay consisting of multiple kinases and two phosphorelay proteins, Spo0F and Spo0B (Burbulis et al. 1991). Environmental and physiological signals are integrated by the relay and by dedicated phosphatases that drain phosphoryl groups from it, to determine the phosphorylation state of Spo0A [Grossman 1995; Perego and Hoch 2002]. Phosphorylated Spo0A [Spo0A–P] sets up a self-reinforcing cycle that stimulates its own synthesis and phosphorylation [Predich et al. 1992; Strauch et al. 1993; Fujita and Sadaie 1998]. Spo0A–P acts in conjunction with RNA polymerase containing the housekeeping sigma factor σ^H^ and with the alternative sigma factor σ^H^ to induce gene transcription at the start of sporulation [Stragier and Losick 1996; Piggot and Losick 2002]. This transcription is responsible for remodeling the sister chromosomes of the developing cell [the sporangium] into an elongated nucleoid known as the axial filament [Pogliano et al. 2002, Ben-Yehuda et al. 2003] and for the formation of an asymmetrically positioned [polar] septum that divides the sporangium into unequal-sized compartments known as the forespore [the small compartment] and mother cell [Levin and Losick 1996; Ben-Yehuda and Losick 2002]. Spo0A–P is also responsible for the synthesis and activation of the cell-specific regulatory proteins that set in motion the forespore and mother-cell lines of gene expression [Stragier and Losick 1996; Piggot and Losick 2002].

Here, we reexamine the role of Spo0A in sporulation. We find that Spo0A continues to function after the initiation phase of sporulation. Our results indicate that Spo0A accumulates to high levels following the formation of the polar septum, when it is present principally or exclusively in the mother cell. Cells that have been en-
engineered to produce an activated form of Spo0A in the forespore are impaired in sporulation. Likewise, cells that had been engineered to produce a newly devised inhibitor of Spo0A phosphorylation [called Spo0A-N] in the mother cell are defective in sporulation and in mother-cell-specific gene expression. Thus, Spo0A appears to have two distinct functions: as a master regulator during the initiation phase of sporulation and as a mother-cell-specific transcription factor following the formation of the polar septum. In terms of abundance and transcriptional activity, Spo0A may function predominantly as a cell-specific regulatory protein.

Results

Preferential transcription in the mother cell from promoters under the control of Spo0A

We previously presented evidence that transcription of the sporulation operon spoIIG persists after the morphological stage of polar division and that this transcription occurs preferentially in the mother cell compartment of the postdivisional sporangium [Fujita and Losick 2002]. In comparison, a constitutively active promoter P\text{spac}\text{c} which was derived from the IPTG-inducible P\text{spac} promoter by elimination of the operator for Lac repressor, [Fujita and Losick 2002] directs the accumulation of green fluorescent protein (GFP) in both the forespore and mother-cell compartments of the postdivisional sporangium. P\text{spoIIG} and P\text{spac}\text{c} are transcribed by σ^A-containing RNA polymerase, but transcription from P\text{spoIIG} is additionally dependent upon the phosphorylated form (Spo0A–P) of Spo0A. Thus, it was conceivable that Spo0A acts preferentially in the mother cell. As a test of this idea, we fused gfp to an additional sporulation promoter [that for the spoIIE operon] that is under the control of Spo0A–P. As shown in Figure 1, GFP from the P\text{spoIIE–gfp} fusion as well as from a P\text{spac–gfp} fusion preferentially accumulated in the mother-cell compart-

![Figure 1. Subcellular localization of GFP produced under the control of Spo0A during sporulation. Strains carrying a P\text{spoIIE–gfp} fusion [MF237] or a P\text{spac–gfp} fusion [MF277] were induced to sporulate, treated with the vital membrane stain FM4-64 at hour 3 of sporulation, and observed by fluorescence microscopy. A strain carrying a P\text{spac–gfp} fusion [MF339] was used as a control. Bar, 1 µm.](image-url)

ment of sporangia that had reached the stage of engulfment. In contrast, GFP from the P\text{spac–gfp} construct was uniformly distributed throughout the sporangia. These findings are consistent with the idea that Spo0A remains active after polar division when it acts preferentially in the mother cell.

Selective accumulation of Spo0A in the mother cell

In light of the above results, we wondered whether Spo0A preferentially accumulates in the mother cell. To investigate this possibility, we carried out immunofluorescence microscopy using anti-Spo0A antibodies. Consistent with its being a DNA-binding protein, Spo0A was found to colocalize with the nucleoid. In predivisional sporangia and shortly after the formation of the polar septum, the sister chromosomes of the sporangium are in the form of an elongated, serpentine-like structure known as the axial filament. As sporulation progresses, the axial filament is resolved into separate mother cell and forespore nucleoids and the two nucleoids become compact [particularly so in the forespore] and amorphous.

Early after asymmetric division [hour 2], Spo0A was found in association with the axial filament in both chambers of the sporangium but generally appeared to be enriched in the mother cell. Over time, the abundance of Spo0A seemed to increase in the mother cell relative to the forespore, and by hour 4, little Spo0A could be detected in the forespore (Fig. 2, Supplementary Table 1). Instead, a strong signal from Spo0A was detected in the mother cell that colocalized with the mother-cell nucleoid. In comparison, the fluorescent signal obtained with antibodies against the major housekeeping sigma factor, σ^A, was no less intense in the forespore than in the mother cell [Fig. 2, Supplementary Table 1].

These results suggest that Spo0A is a cell-specific transcription factor that selectively accumulates in the mother cell. Selective accumulation of Spo0A in the mother cell could be the result of persistent and selective synthesis of the regulatory protein in the mother cell or to degradation of Spo0A in the forespore, or both.

Activation of Spo0A in the forespore inhibits sporulation

As a test of the idea that Spo0A acts in a cell-specific manner after polar division, we investigated the effect of producing a constitutively active form of Spo0A in the forespore. Sad67 is a mutant form of Spo0A that is locked in an active state because of a short deletion internal to its coding sequence [Ireton et al. 1993]. Sad67 does not depend on phosphorylation in order to be active. We placed the gene for Sad67 under the control of P\text{spoIIQ}, the promoter for a sporulation gene [spoIIQ] that is transcribed in the forespore by σ^A-containing RNA polymerase. The sporulation efficiency of the P\text{spoIIQ–sad67} containing strain [MF826] was 2% that of the corresponding wild-type strain as judged by the production of heat-resistant, colony-forming units [spores] and by phase-con-
trast microscopy [Fig. 3]. That the impaired sporulation efficiency of MF826 was a result of the use of the sad67 allele was demonstrated by the construction of a strain (MF828) in which wild-type spo0A was placed under the control of PspoIIQ. In contrast to MF826, MF828 was largely unimpaired in its ability to undergo sporulation. As a control, we constructed a strain (MF825) in which sad67 was under the control of the promoter for a gene (spoIID) that is transcribed in the mother cell by αE-containing RNA polymerase. The sporulation efficiency of MF825 was similar from that of the wild type (Fig. 3). Finally, Western blot analysis carried out with antibodies against Spo0A showed that Sad67 (which can be distinguished from the wild-type protein by its smaller size) accumulated to similar levels in the PspoIIQ–sad67-containing and the PspoIID–sad67-containing strains. In toto, these findings indicate that Spo0A impairs sporulation when it is present and active in the forespore.

In other work, we sought to determine the stage of sporulation at which Sad67-producing sporangia become arrested. The results showed that cells of MF826 reached the stage of engulfment but were blocked in the expression of a gene (spoIVB) under the control of the late-acting, forespore transcription factor σF but not in the expression of genes under the control of the earlier-acting regulatory proteins Spo0A, σH, and σE (data not shown).

**Use of a truncated form of Spo0A to block activation of the full-length transcription factor**

The results with Sad67 are consistent with the view that Spo0A is not normally active in the forespore. Next, we wished to address the complementary issue of whether Spo0A is not only active in the mother cell but also required in the mother cell for efficient sporulation. To address this issue, we took advantage of the fact that Spo0A, like other response-regulator proteins, consists of
two domains: a receiver domain (the site of phosphorylation), which is located in the N-terminal portion of the protein, and a transcriptional regulatory (DNA-binding) domain, which is confined to the C-terminal region (Hoch 2000). The receiver domain is phosphorylated by means of a phosphorelay involving multiple kinases and two phosphorelay proteins, Spo0F and Spo0B. Spo0F is responsible for transferring phosphoryl groups from the kinases to Spo0B, which, in turn, transfers the phosphoryl groups to Spo0A (Burbulys et al. 1991). We reasoned that a truncated form (Spo0A-N) of Spo0A corresponding to the N-terminal domain (residues 1–142) would compete with Spo0A for phosphorylation by the phosphorelay and hence impair activation of the full-length response regulator. Because sporulation is believed to depend on the attainment of a threshold level of Spo0A activity (Chung et al. 1994), we anticipated that even a modest level of competition might be enough to block sporulation.

As a test of our reasoning, we created a strain (MF972) harboring a fusion \( P_{\text{spo0A}}-\text{spo0A-N} \) of the coding sequence \( \text{spo0A-N} \) for Spo0A-N to the promoter for \( \text{spo0A} \). MF972 produces both Spo0A-N and the full length Spo0A at the start of sporulation. The results of Figure 4A show that cells of MF972 were impaired in sporulation, exhibiting a block prior to polar septum formation.

Next, we carried out immunoblot analysis to determine the relative levels of Spo0A and Spo0A-N produced by MF972. A complication in this analysis was the likelihood that Spo0A-N, being a truncated form of Spo0A, would not be as immunoreactive as the full-length protein. Accordingly, we purified Spo0A and Spo0A-N and used the purified proteins to create standardized curves relating the intensity of the immunoblot signal to protein concentration. The results showed that Spo0A was approximately fivefold more immunoreactive than Spo0A-N on an equimolar basis (Fig. 4B). Using the standardized curves, we determined that Spo0A and Spo0A-N accumulated to similar levels in strain MF972, reaching concentrations of ~200 fmole/µg of total cellular protein, which corresponds to ~500 molecules/cell.

Overproduction of Spo0A suppresses the sporulation-inhibitory effect of Spo0A-N

We interpret the results with MF972 to indicate that Spo0A-N competes with Spo0A for phosphorylation by the phosphorelay. If this interpretation is correct, then by artificially raising the concentration of Spo0A, we should be able to overcome the inhibitory effect of Spo0A-N and restore efficient sporulation. To test this prediction, we placed wild-type \( \text{spo0A} \) under the control of the IPTG-inducible promoter \( P_{\text{spac}} \). We then constructed a strain (MF1258) that harbored the IPTG-inducible \( P_{\text{spac}}-\text{spo0A} \) fusion and the \( P_{\text{spo0A}}-\text{spo0A-N} \) fusion as well as a wild-type copy of \( \text{spo0A} \). As shown in Table 1, MF1258 sporulated poorly in the absence of in-

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**Figure 4.** Inhibition of Spo0A activation in the mother cell inhibits sporulation. (A) Whole-cell extracts from cells of strain PY79 (wild type), MF972 \( P_{\text{spo0A}}-\text{spo0A-N} \), MF973 \( P_{\text{spo0A}}-\text{spo0A-N} \), and MF974 \( P_{\text{spo0A}}-\text{spo0A-N} \) that had been collected at 1-h intervals after the start of sporulation were subjected to SDS-PAGE and immunoblot analysis using anti-Spo0A antibodies (top panel). Fluorescence microscopic images of cells that had been treated with FM4-64 at hour 4 of sporulation and sporulation efficiencies are shown for each of the strains (bottom panel). (B) Samples of purified Spo0A–His8 and Spo0A–N–His8 were subjected to SDS-PAGE and stained with Coomassie brilliant blue (lanes 1, 2) or were subjected to SDS-PAGE and immunoblot analysis using anti-Spo0A antibodies (lanes 3–5). Samples of total protein from cells of strain MF972 collected at hour 2 of sporulation were subjected to SDS-PAGE and immunoblot analysis using anti-Spo0A antibodies (lanes 6–8). (Lane 1) Spo0A–N–His8; (Lane 2) Spo0A–His8; (Lane 3) Spo0A–His8 + Spo0A–N–His8; (Lane 4) Spo0A–His8 + Spo0A–N–His8 + Spo0A–N–His8; (Lane 5) Spo0A–His8 + Spo0A–N–His8 + Spo0A–N–His8; (Lane 6) Total protein (0.5 µg); (Lane 7) Total protein (1 µg); (Lane 8) Total protein (2 µg). Note that the His-tagged proteins migrated slightly slower than the corresponding untagged proteins. Positions of molecular weight markers (kD) are indicated on the left.
ducer. However, the presence of IPTG enabled MF1258 to sporulate efficiently. This finding reinforces the view that the sporulation-inhibitory effect of Spo0A-N is a result of competition with the full-length regulatory protein.

**Spo0A-N competes with Spo0A for phosphorylation by Spo0B**

To investigate the mechanism of action of Spo0A-N further, we asked whether or not purified Spo0A-N was capable of competing with the full-length protein for phosphorylation by phosphorylase proteins. Accordingly, we purified functional, tagged versions of Spo0A-N, KinA, Spo0F, Spo0B, and Spo0A [see Materials and Methods; see Supplemental Material], and reconstituted the phosphorylase in vitro [Fig. 5A, lanes 1–4]. Similar to results obtained previously with a proteolytic fragment corresponding to the NH2-terminal portion of Spo0A [Grimsley et al. 1994], Spo0A-N was an effective substrate for phosphorylation by the phosphorylase (Fig. 5A, lane 5) and in a manner that was dependent upon the most downstream member of the phosphorylase, Spo0B [Burbulys et al. 1991]. Importantly, in mixtures of the full length and truncated proteins, phosphorylation of Spo0A was partially inhibited by an equimolar amount of Spo0A-N and inhibited further as the concentration of Spo0A-N was raised [Fig. 5A, lane 6, 5B]. At the same time, neither Spo0A-N nor Spo0A had a measurable effect on the level of phosphorylation of the phosphorylase protein Spo0F by the kinase KinA [Fig. 5A, lanes 2,7,8]. Taken together, our results indicate that Spo0A-N is an effective and specific competitor of the phosphorylation of Spo0A by Spo0B–P.

**Blocking Spo0A activation in the mother cell inhibits sporulation**

In toto, the above results indicate that Spo0A-N can be used as a specific inhibitor of Spo0A and that it does so by interfering with the phosphorylation of the full-length protein. Finally, we wished to take advantage of this specific inhibitory effect to investigate whether or not sporulation depends on the continued activity of Spo0A in the mother cell. Accordingly, we constructed a strain in which a Spo0A-N was produced in the mother cell under the control the promoter for a gene (spoIID)

**Table 1. Overproduction of Spo0A suppresses the sporulation defect of a strain producing Spo0A-N**

| Strain                        | Induction Spore/mL | Sporulation efficiency |
|-------------------------------|-------------------|------------------------|
| WT                            | 5.0 × 10^8        | 1                      |
| p_pspoIID–spo0A-N             | 2.0 × 10^7        | 0.04                   |
| p_pspoIID–spo0A-N             | –IPTG 5.5 × 10^7  | 0.11                   |
| p_pspoIID–spo0A-N             | +IPTG 5.5 × 10^8  | 1.1                    |

*MF972 (ze82::amyE::PspoIID–spo0A-N)*

*MF1258 (ze82::amyE::PspoIID–spo0A-N, amyE::PspoIID–spo0A-N)*

that is transcribed by the mother-cell transcription factor αE. The results of Figure 4A show that Spo0A-N impaired sporulation when the truncated protein was produced in the mother cell in a strain [MF973] harboring a fusion [P_spoIID–spo0A-N] of spo0A-N to the promoter for spoIID. As a control, production of Spo0A-N in the forespore under the control of the promoter for spoIIQ (strain MF974) caused little or no impairment of sporulation. Western blot analysis showed that Spo0A-N accumulated to similar levels when produced under the control of αE [P_spoIIQ–spo0A-N] or αF [P_spoIID–spo0A-N; Fig. 4A].

As an additional control, we investigated the effect of producing Spo0A-N under the control [using the promoter for the gerE gene] of a late-appearing transcription factor in the mother cell, αE [data not shown]. The results showed that Spo0A-N impaired sporulation when produced at an early time in the mother-cell line of gene expression but not when produced at a late stage of development.

**Production of Spo0A-N in the mother cell inhibits Spo0A-dependent transcription from a mother-cell-expressed promoter**

If Spo0A-N in MF973 was acting by inhibiting the phosphorylation of Spo0A in the mother cell, then transcription from a promoter under the control of Spo0A should be impaired in the mother cell. As a test of this expectation, we used lacZ fused to the promoter for an operon (spoIIG) that is under the control of Spo0A to monitor the effect of Spo0A-N on Spo0A-directed gene transcription (Fig. 6). When Spo0A-N was produced under the control of the P_spoIID promoter (MF1151), accumulation of β-galactosidase from the P_spoIIG–lacZ fusion ceased at about hour 2 of sporulation, at a time approximately corresponding to when the polar division takes place and αE [and hence the P_spoIID–spo0A-N construct] is activated. For comparison, when Spo0A-N was produced under the control of the spo0A promoter (MF1140), accumulation of β-galactosidase was inhibited from the onset of sporulation. Furthermore, when Spo0A-N was produced in the forespore under the P_spoIIQ promoter (MF1161), the pattern of β-galactosidase accumulation was indistinguishable from that of a strain [MF290] that did not produce Spo0A-N.

A complication in interpreting the results of Figure 6 is that β-galactosidase is unstable during sporulation. Hence the cessation of β-galactosidase accumulation seen at hour 2 in strain MF1151 is a composite of reduced transcription from P_spoIIG–lacZ as well as the degradation of β-galactosidase. We therefore interpret the results of Figure 6 to indicate Spo0A-N inhibited the transcription of P_spoIIG–lacZ in the mother cell, but did not necessarily block it altogether.

**Visualizing the effect of Spo0A-N on cell-specific gene expression**

To further analyze the effects of Spo0A-N on Spo0A-dependent gene expression, we sought to monitor the
influence of Spo0A-N on the spatial pattern of gene expression. To do this, we fused the coding sequence for the GFP to promoters under the control of Spo0A (PspoIIG), PspoIID, and PspoIIQ; Fig. 7). Green fluorescence from the PspoIIG–gfp reporter construct in an otherwise wild-type strain (MF237) accumulated selectively in the mother cell after 3 h of sporulation. Meanwhile, GFP from the PspoIID–gfp (MF248) and PspoIIQ–gfp (PE128) reporter constructs was, as expected, restricted to the mother cell and forespore compartments, respectively of the postdivisional sporangia. (Notice that owing to the lipophilic nature of FM4-64, the membrane dye did not stain forespores in sporangia that had completed engulfment.)

Next, we created strains harboring the above described reporter constructs in combination with fusions of spo0A-N to the promoter for spo0A (Pspo0A–spo0A-N) and promoters under the control of PspoIID–spo0A-N and PspoIIQ–spo0A-N. When Spo0A-N was produced from the Pspo0A–spo0A-N fusion at the onset of sporulation, production of GFP from all three reporter constructs was significantly impaired and many of the sporangia failed to reach the stage of polar division. When Spo0A-N was produced in the mother cell from the PspoIID–spo0A-N fusion, production of GFP from the Spo0A-dependent reporter (PspoIIG–gfp) was partially impaired (a minority of the sporangia exhibited significant levels of GFP in the mother cell) and production of GFP from the PspoIID–gfp was substantially impaired. In contrast, little or no impairment of PspoIIQ–spo0A-N fusion and no defect in polar division or engulfment was observed.

**Figure 5.** Spo0A-N competes with Spo0A for phosphorylation by phosphorelay proteins in vitro. Kinase reactions were performed as described in Materials and Methods, and the proteins were analyzed by SDS-PAGE. (A) Reaction mixtures (20 µL) contained 0.2 µM KinA (lanes 1–8), 0.2 µM Spo0F (lanes 2–8), 0.2 µM Spo0B (lanes 3–6), 2 µM Spo0A (lanes 4, 6, 7), and 2 µM Spo0A-N (lanes 5, 6, 8). Reaction mixtures were incubated at 25°C for 1 h and then stopped by adding SDS-loading buffer. Samples were subjected to electrophoresis through a 16% SDS–polyacrylamide gel, and radioactive proteins were visualized by autoradiography (right panel). Fourteen picomoles of each purified protein was subjected to electrophoresis and stained with Coomassie as a reference for electrophoretic mobility (left panel). (B) Reaction mixtures contained 0.2 µM KinA, 0.2 µM Spo0F, 0.2 µM Spo0B, and 2 µM Spo0A. Increasing amounts of Spo0A-N (0, 2, 4, and 6 µM, from left to right) were added to the reactions. The relative levels of Spo0A–P are indicated.

**Figure 6.** Inhibition of Spo0A activation inhibits expression of the spoIIG operon. Culture samples from strains MF290 (Δ, amyE::PspoIIG–lacZ), MF1161 (Δ, amyE::PspoIIG–lacZ, zei82::amyE::PspoIIG–spo0A-N), MF1151 (Δ, amyE::PspoIIG–spo0A-N), and MF1140 (Δ, amyE::PspoIIG–lacZ, zei82::amyE::Pspo0A–spo0A-A-N), were collected at 1-h intervals after the start of sporulation and analyzed for β-galactosidase activity.
Discussion

Spo0A has long been recognized as the master regulator for entry into sporulation (Burbulys et al. 1991; Hoch 1993). Mutants of the response regulator fail to undergo the hallmark morphological events (axial filament formation and polar division) characteristic of early development (Levin and Losick 1996; Ben-Yehuda and Losick 2002; Ben-Yehuda et al. 2003) or to switch on genes and operons (spoIIE, spoIIA, and spoIIG) that activate the forespore and mother-cell lines of gene expression (Stragier and Losick 1996; Piggot and Losick 2002). In modification of this traditional view, the results of the present investigation indicate (1) that Spo0A continues to function after the initiation phase of sporulation when it becomes a mother-cell-specific transcription factor, and (2) that its persistent and selective activity in the mother cell plays a critical role in development. Indeed, in terms of abundance and transcriptional activity, Spo0A arguably functions predominantly as a cell-specific regulatory protein.

These conclusions derive from the following observations. First, GFP produced under the control of two different Spo0A-controlled promoters continued to accumulate to high levels after asymmetric division and selectively in the mother cell. Second, Spo0A itself accumulated to high levels in the mother cell but not in the forespore. Third, an activated form of Spo0A (Sad67) impaired sporulation when it was produced in the forespore but not when it was produced in the mother cell. Finally, a newly devised, competitive inhibitor (Spo0A-N) of the phosphorylation of Spo0A impaired sporulation and mother-cell-specific gene expression when the inhibitor was produced in the mother cell but not when it was produced in the forespore.

Additional evidence consistent with the view that Spo0A functions persistently during sporulation comes from studies with a heat-sensitive mutant of the response regulator. F. Kawamura (pers. comm.) has observed that the mutant remains sensitive to a shift to a restrictive temperature for several hours after the start of sporulation. If, as we contend, the predominant role of Spo0A is as a cell-specific transcription factor, then what is the function of Spo0A in the mother cell? Part of the answer, we believe, involves the mother-cell regulatory protein/hibitor. The/hibitor has traditionally been thought of as the earliest-acting regulatory protein in the mother-cell line of gene expression (Stragier and Losick 1996; Piggot and Losick 2002). It is derived by proteolytic processing from an inactive proprotein precursor, pro-/hibitor. Both pro-/hibitor and its processing enzyme, SpoIIGA, are encoded by the spoIIG operon (Jonas et al. 1988), which is under the direct control of Spo0A (Satola et al. 1992). As shown previously (Fujita and Losick 2002) and confirmed here, transcription of spoIIG largely takes place after the formation of the polar septum when it is confined to the mother cell. Also, as shown previously (Fujita and Losick 2002), /hibitor and/or pro-/hibitor accumulate to high levels after asymmetric division and selectively in the mother cell. These findings are now explained by the fact that Spo0A largely acts as a mother-cell-specific transcription. Meanwhile, processing of pro-/hibitor is subject to an intercompartamental regulatory pathway involving the forespore transcription factor /hibitor that ensures that pro-/hibitor is not converted to

Figure 7. Subcellular localization of GFP synthesized under the control of Spo0A, $\sigma^f$, or $\sigma^d$ in the cells producing Spo0A-N. The strains used were MF237 ($P_{spoIIG}$-gfp), MF1146 ($P_{spoIIQ}$-gfp, $P_{spo0A}$-spo0A-N), MF1156 ($P_{spoIIQ}$-gfp, $P_{spo0A}$-spo0A-N), MF1166 ($P_{spoIIQ}$-gfp, $P_{spo0A}$-spo0A-N), MF248 ($P_{spoIIQ}$-gfp), MF1144 ($P_{spoIIQ}$-gfp, $P_{spo0A}$-spo0A-N), MF1157 ($P_{spoIIQ}$-gfp, $P_{spoIIQ}$-spo0A-N), MF1167 ($P_{spoIIQ}$-gfp, $P_{spoIIQ}$-spo0A-N), PE128 ($P_{spoIIQ}$-gfp), MF1145 ($P_{spoIIQ}$-gfp, $P_{spo0A}$-spo0A-N), MF1158 ($P_{spoIIQ}$-gfp, $P_{spoIIQ}$-spo0A-N), and MF1168 ($P_{spoIIQ}$-gfp, $P_{spoIIQ}$-spo0A-N). Cells were collected at hour 3 of sporulation, treated with the membrane strain FM4-64, and visualized by fluorescence microscopy. Bar, 1 µm.
mature α until after the polar septum is formed (Stragier and Losick 1996; Piglot and Losick 2002). Thus, the relatively low level of pro-α that is produced in the predivisional sporangium does not become active until after asymmetric division. Viewed in this light, the earliest-acting, cell-specific transcription factor in the mother-cell line of gene expression is Spo0A rather than σE.

An implication of our findings is that the phosphorelay, which is responsible for phosphorylating Spo0A, also continues to function in the postdivisional sporangium. Indeed, the effectiveness of Spo0A-N in inhibiting Spo0A-directed gene transcription in the mother cell indicates that Spo0A function depends on its continued phosphorylation in the mother cell. The phosphorelay is generally believed to be a device for integrating environmental and physiological signals into the decision to sporulate, but the relay may also play a role in the conversion of Spo0A into a cell-specific transcription factor.

An important challenge for the future will be to elucidate the mechanisms that cause Spo0A to accumulate and function in a cell-specific manner. An appealing hypothesis is based on the phenomenon of transient exclusion. Formation of the polar septum initially traps the origin-proximal one third of the chromosome that is destined for the forespore in the small chamber of the sporangium. The remaining two thirds of the chromosome is subsequently pumped into the forespore across the polar septum by the DNA translocase SpoIIIIE (Wu and Errington 1994). This results in a period of transient genetic asymmetry lasting for about 15 min when the forespore is lacking origin-distal genes and the mother cell temporarily contains an extra copy of these genes (Frandsen et al. 1999; Pogliano et al. 1999; Khvorova et al. 2000; Dworkin and Losick 2001; Zapancic et al. 2001). We note that the genes for all of the components of the phosphorelay, except Spo0F, are located outside the region of the chromosome that is initially trapped in the forespore (Stragier and Losick 1996). Thus, the temporary exclusion of phosphorelay genes from the forespore could set up an imbalance in which relay proteins are preferentially synthesized in the mother cell.

The effect of such an imbalance could be amplified by the following positive feedback loop. Spo0A–P is known to stimulate the production of σE (through an indirect pathway involving the repressor AbrB), which, in turn, directly stimulates the transcription of the gene for Spo0A and the gene for KinA, the principal kinase for the phosphorelay (Fredich et al. 1992; Fujita and Sadaie 1998). In addition, both Spo0A–P and σE stimulate the transcription of the gene for the phosphotransferase protein Spo0F (Strauch et al. 1993; Fujita and Sadaie 1998). Such a self-reinforcing cycle would lead both to preferential synthesis of Spo0A and to preferential phosphorylation of the response regulator in the mother cell. Also contributing to the selective accumulation of Spo0A in the mother could be preferential degradation of the response regulator in the forespore. Conceivably, unphosphorylated Spo0A in the forespore is subject to proteolysis to a greater extent than is phosphorylated Spo0A in the mother cell.

In summary, Spo0A plays a dual role in sporulation, governing both the initiation phase of sporulation in the predivisional sporangium and acting as a cell-specific transcription factor in the postdivisional sporangium. Viewed in this light, Spo0A offers a striking parallel to the master regulator CtrA for the cell cycle in the dimorphic bacterium Caulobacter crescentus (Jensen et al. 2002; Shapiro et al. 2002). Like Spo0A, CtrA is a response regulator whose phosphorylation is believed to be governed by a phosphorelay [although this has not yet been demonstrated]. Also, like Spo0A, CtrA functions both in the predivisional stage of the C. crescentus cell cycle as well as following division when it is restricted to only one (the swarmer cell) of the two resulting progeny cells. As more is learned about the compartmentalization of Spo0A, it will be interesting to compare and contrast the mechanisms by which Spo0A and CtrA come to act in a cell-specific manner.

Materials and methods

General methods

General methods were as previously described (Fujita and Losick 2002). All B. subtilis strains were derived from the prototrophic strain PY79 (Youngman et al. 1983) and are listed in Supplementary Table 1.

Plasmid construction

A complete description of the plasmids used can be found in Supplemental Material.

Fluorescence microscopy

Fluorescence microscopy was performed as described previously [Fujita and Losick 2002].

Immunofluorescence microscopy

Immunofluorescence microscopy was performed as described previously [Hofmeister 1998]. The rabbit polyclonal antibodies that bind to σE and Spo0A (Fujita 2000) were used at 1:10,000 and 1:5000 dilutions, respectively.

Immunoblot analysis

Immunoblot analysis was done by the procedure of Rudner et al. [1999]. Polyclonal anti-Spo0A antibodies were used for the detection of Spo0A (Fujita 2000).

Protein expression and purification

KinA, Spo0F, Spo0B, Spo0A, and Spo0A-N were produced as C-terminal His6 fusion proteins in Escherichia coli from plasmids pMF193 (see Supplemental Material), pGK10 (Burkholder et al. 2001), pMF184 (see Supplemental Material), pMF182 (modified version of pETSpo0A; Fujita and Sadaie 1998), and pMF182 (see Supplemental Material), respectively. Protein production was carried out in the E. coli BL21 (DE3) pLysS strains. Cells were grown in LB at 30°C to an OD600 of 0.6, induced by addition of IPTG to 1 mM, and harvested after 5 h. The proteins were purified by affinity chromatography on Ni-NTA agarose according to the manufacturer’s protocol [Qiagen]. Protein concentrations were determined by Bradford assay (Pierce) using BSA as a standard.

Phosphorylation reaction

Phosphorylation reaction was done by the procedure of Burbulys et al. [1991]. The reactions were initiated by addition of the ATP, stopped by addition of SDS-loading buffer, and immedi-
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ately loaded on a 16% SDS–polyacrylamide gel. Gels were exposed to X-ray film to detect phosphorylated proteins.

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Masaya Fujita and Richard Losick

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