An epigenetic switch governing daughter cell separation in *Bacillus subtilis*

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Growing cells of *Bacillus subtilis* are a bistable mixture of individual motile cells in which genes for daughter cell separation and motility are ON, and chains of sessile cells in which these genes are OFF. How this ON/OFF switch is controlled has been mysterious. Here we report that a complex of the SinR and SlrR proteins binds to and represses genes involved in cell separation and motility. We also report that SinR and SlrR constitute a double-negative feedback loop in which SinR represses the gene for SlrR (*slrR*), and, by binding to (titrating) SinR, SlrR prevents SinR from repressing *slrR*. Thus, SlrR indirectly derepresses its own gene, creating a self-reinforcing loop. Finally, we show that, once activated, the loop remains locked in a high SlrR state in which cell separation and motility genes are OFF for extended periods of time. SinR and SlrR constitute an epigenetic switch for controlling genes involved in cell separation and motility.

*Keywords*: Epigenetic; bistability; biofilm; *Bacillus subtilis*

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Biofilm formation is governed by a regulatory circuit that involves three dedicated regulatory proteins: SinI, SinR, and SlrR (Kearns et al. 2005; Chu et al. 2008; Kobayashi 2007). SinI is produced under the control of SpoOA, a global regulator of post-exponential-phase gene expression in *B. subtilis* (Gaur et al. 1988; Sonenshein 2000; Piggot and Hilbert 2004; Fujita et al. 2005; Chai et al. 2008). SinI is an anti-repressor that binds to and inhibits the SinR repressor, a protein that is produced constitutively (Bai et al. 1993; Kearns et al. 2005). SinR, in turn, binds to the promoters for matrix operons *yqxM* and *eps* (exopolysaccharide) and the gene for SlrR *slrR*, blocking their transcription [Fig. 1A; Kearns et al. 2005; Branda et al. 2006; Chu et al. 2006, 2008]. SlrR is a hybrid protein, containing domains that are homologous to both the SinI anti-repressor and the SinR repressor [Chu et al. 2008].

We report that SinR and SlrR constitute a double-negative feedback loop involving protein–protein [SlrR-mediated inhibition of SinR] and protein–DNA [SinR-mediated repression of *slrR*] interactions. We show that SlrR binds to SinR, and that the formation of the SinR•SlrR complex inhibits the capacity of SinR to bind to the *slrR* promoter [and other SinR targets]. Thus, the binding of SlrR to SinR causes derepression of its own gene, thereby creating a self-reinforcing loop. At the same time, the SinR•SlrR complex, but neither SinR alone nor SlrR alone, is a potent repressor of autolysin genes. Finally, evidence indicates that the double-negative feedback loop is an epigenetic switch that locks growing cells in alternative states of cell chaining and motility.

**Results**

*SlrR controls cell chaining during biofilm formation*

The starting point for this investigation was the question of how cell chaining is regulated during biofilm formation. Cells in the biofilm are predominantly in the form of long chains, which are bundled together by an extracellular matrix [Fig. 2A; Branda et al. 2001; Kearns et al. 2005; Kobayashi 2007]. Such chains can be seen after growth in biofilm-inducing medium on agar, and in the pellicles that form at the air/liquid interface of standing cultures [Supplemental Fig. S1]. We found that a mutant of *slrR* (YC131) is defective in cell chaining under both conditions [Fig. 2A]. Furthermore, the effect of the *slrR* mutation was complemented by providing a copy of *slrR* [under the control of its native promoter] at the *amyE* locus on the chromosome [data not shown]. Kobayashi [2007] has independently observed that an *slrR* mutant is blocked in chaining.

To investigate these observations further, we introduced into an *slrR* mutant a construct that contained an IPTG-inducible copy of *slrR* [under control of the *hy-spank* promoter P*hy*-slrR]. The results of Figure 2B show that cell chaining occurred in an IPTG-inducible manner in the engineered strain. That the chains [and other chains observed in this study] were composed of linked cells that had undergone septation but had not separated from each other was confirmed by staining with the vital membrane stain FM4-64 [data not shown]. SlrR is known to stimulate the production of the matrix protein TasA [Chu et al. 2008; Murray et al. 2009]. We wondered whether the effect of *slrR* induction on cell chaining was indirectly due to increased TasA production, which we investigated using the P<sub>amy</sub>-slrR construct. The results of Figure 2C show that induction of chaining by the addition of IPTG was not dependent on TasA. It was also not dependent on *eps*, as judged by use of a mutation (∆*epsH*) in the *eps* operon [Fig. 2D]. Therefore, cell chaining is not dependent on extracellular matrix production, and SlrR evidently has two distinct roles: stimulating matrix production, and promoting chaining.

*SlrR represses autolysin genes*

The separation of daughter cells after division is governed by peptidoglycan-degrading enzymes called autolysins [Blackman et al. 1998; Smith et al. 2000; Vollmer et al. 2008]. How does SlrR control cell chaining? The simplest hypothesis is that SlrR is a repressor of the autolysin genes. Two autolysin genes that are known to play a key role in cell separation during vegetative growth are *lytC*...
and lytF (Ohnishi et al. 1999; Chen et al. 2009). The lytC and lytF genes are transcribed under the direction of the alternative σ factor σ^D, which also directs the transcription of genes for motility and chemotaxis (Margot et al. 1999; Chen et al. 2009). To test the hypothesis that SlrR represses these σ^D-dependent genes, we fused the promoters for the lytABC operon, which contains lytC, and lytF to the lacZ gene, creating P_{lytA-lacZ} and P_{lytF-lacZ}, respectively. We then introduced the fusions into the wild type and into strains mutant for slrR (ΔslrR), or the σ^D-encoding gene sigD (ΔsigD), or both slrR and sigD (ΔslrR ΔsigD). The results shown in Figure 3, A (P_{lytA-lacZ}) and B (P_{lytF-lacZ}), demonstrate that expression of both fusions was somewhat higher in the slrR mutant than in the wild type (~120% and ~35% higher for P_{lytA-lacZ} and P_{lytF-lacZ}, respectively) in the early stationary phase in biofilm-inducing medium. The results also show that expression of lytC and lytF indeed depended on σ^D, because both fusions showed little activity in the absence of the alternative σ factor (Fig. 3A,B). The above results are consistent with the idea that SlrR negatively regulates lytC and lytF. However, the effect of the absence of SlrR seemed too small to account for the strong effect of the mutation on cell chaining (Fig. 2A).

A possible explanation comes from the fact that slrR is under the direct negative control of SinR (Chu et al. 2008; Kobayashi 2008). SinR-controlled genes, such as the eps and yqxm operons, are expressed in a bimodal pattern in shaking culture, with a small number of ON and a large number of OFF cells in the population (Chai et al. 2008; Vlamakis et al. 2008). We wondered whether expression of slrR also occurs in only a subpopulation of cells. If so, this bimodality could explain why only a mild effect was seen on lytC and lytF expression when slrR was mutant, as we were measuring β-galactosidase synthesis averaged among all cells in the population. To investigate whether slrR was ON in only a subpopulation of cells, we constructed a fusion of the promoter for slrR to the gene for the green fluorescent protein (GFP), creating P_{slrR-gfp}. We integrated the fusion into the chromosome at the amyE locus and examined P_{slrR-gfp} expression by fluorescence microscopy. As shown in Figure 3C, only a small subpopulation of cells actively expressed P_{slrR-gfp}. Indeed, this subpopulation corresponded to cells that were in the form of chains. We infer that SlrR does repress lyt genes strongly, but only in the subpopulation of cells in which slrR is expressed at high levels. If so, then we should see strong repression of lyt gene expression using P_{lytA-slrR}, which should lead to slrR expression in all cells. Indeed, when this construct was used, overall expression of P_{lytA-lacZ} was markedly repressed in the presence of IPTG during early stationary phase under shaking culture conditions (Fig. 3D).

Cell chaining depends on both SlrR and SinR

The slrR gene lies downstream from sinI and sinR in the regulatory circuit that controls both matrix production and chaining (Fig. 1; Chu et al. 2008). We predicted that sinI and sinR should also have strong effects on cell chaining. In fact, and as shown in Figure 4A, a null mutation in sinI, which greatly decreased slrR expression (Chu et al. 2008), caused an almost complete loss in cell chaining.
An epigenetic switch

Figure 4. Chaining phenotype of mutants grown in shaking culture in biofilm medium. [{A}, top row] A ΔsinI mutant (RL3853) was impaired in chaining. Overexpression of sinI (YC227, a functional gfp-sinI translational fusion was used) promoted extensive cell chaining, and a ΔslrR mutation reversed the chaining phenotype caused by overexpression of sinI (YC589). {Bottom row} A ΔsinR mutant (RL3856) showed little chaining (the strain contained a ΔepsH mutation to prevent aggregation), a ΔsinR mutation reversed the chaining phenotype caused by overexpression of sinI (YC228), a ΔsigD mutant showed extensive cell chaining (RL4169), and the chaining phenotype was not reversed by ΔslrR (YC205). {B,C} Induction of slrR [from PPy−slrR] promoted chaining in shaking culture (strain YC280, B) but not in a strain that with a ΔsinR mutation (strain YC284, C). {D,E} Shown are point mutants of sinR that promoted chaining [D shows strains YC606, YC607, and YC608, respectively, from top to bottom panels]. {E} Chaining caused by the point mutations was reversed by the presence of a ΔslrR mutation (strains YC620, YC621, and YC622, respectively, from top to bottom).

chaining in shaking culture in biofilm-inducing medium. [Note that, under shaking culture conditions in biofilm-inducing medium, only a modest proportion of wild-type cells form chains (Fig. 4A) as compared with cells embedded in biofilms in which a high proportion of cells are in chains (Fig. 2A).] On the other hand, overexpression of sinI from an IPTG-inducible promoter caused extensive cell chaining throughout the cell population (Fig. 4A). Furthermore, this extensive chaining phenotype was almost completely reversed by introducing an slrR mutation into that strain (Fig. 4A), confirming that slrR lies downstream from sinI in the pathway controlling cell chaining.

Because SinR is a repressor of slrR, and because slrR expression is derepressed in a sinR-null mutant (Chu et al. 2008), we might have expected that a sinR-null mutation would [like sinI overexpression] promote a high level of chaining. However, as observed previously (Kearns et al. 2005), a null mutant of sinR exhibited the opposite phenotype: little cell chaining (Fig. 4A). Furthermore, a ΔsinR mutation reversed the extensive cell chaining phenotype caused by overexpression of sinI (Fig. 4A).

The above experiments were performed using cells harboring a mutation in the eps operon (ΔepsH) to prevent cell aggregation. The ΔepsH mutation itself had no measurable effect on cell chaining [Fig. 2D]. In cells that are wild type for eps, the absence of SinR causes extensive aggregation (Kearns et al. 2005). In these aggregates, the cells are bundled tightly together but are separated from each other along their long axis and do not form chains (Kearns et al. 2005). In toto, these findings indicate that both SinR and SlrR are required for cell chaining.

That chaining depends on both SinR and SlrR is further indicated by the following experiment. As we showed, overexpression of slrR from an IPTG-inducible promoter dramatically enhanced cell chaining during biofilm formation [Fig. 2B]. This was also seen when cells were grown in shaking culture conditions [Fig. 4B]. However, when SlrR was overproduced in a sinR-null mutant strain, little cell chaining was seen [Fig. 4C].

Certain point mutations of sinR promote cell chaining

We showed that a null mutant of sinR is impaired in cell chaining [Fig. 4A]. Historically, however, the sinR gene was called flaD (for flagellaeless), because the original mutants exhibited a flagellaeless and filamentous phenotype [Pooley and Karamata 1984]. These filamentous [or hyperchaining] mutants were mapped to sinR [flaD1 and flaD2] and were found to be point mutations that caused single-aminoacid substitutions in SinR [Margot et al. 1996]. We infer that the flaD1 and flaD2 alleles impart the opposite phenotype [hyperchaining] to that of the null mutation [little chaining].

To investigate the hypothesis that certain single-amino-acid substitutions in SinR can cause a hyperchaining phenotype, we devised a screen for mutant alleles of sinR that promote increased chaining [see the Materials and Methods]. We recovered eight different single-amino acid substitution mutants of SinR that caused conspicuous hyperchaining phenotypes [Fig. 5]. Two substitutions were located in the DNA-binding domain of SinR [V26 and A27], and six [D55, E67, E79, D84, and A85; two different substitutions were recovered at D84] were clustered in the domain believed to be responsible for SinR–SinR interactions [Fig. 5; Lewis et al. 1996].

We picked three mutants for further study: V26G, a substitution in the DNA-binding domain; A85T, a substitution in the SinR–SinR interaction domain; and D84G, another substitution in the SinR–SinR domain that corresponds to the classic flaD1 mutation [Margot et al. 1996]. All three mutants exhibited extensive chaining when grown in biofilm-inducing medium under shaking culture conditions [Fig. 4D]. Also, and as expected, all three mutations caused overexpression of slrR as well as of matrix genes [Fig. 5; Supplemental Fig. S2].

Finally—and to characterize further the cell chaining function of the V26G, A85T, and D84G mutant proteins—we introduced an slrR-null mutation (ΔslrR) into cells harboring the three mutant alleles of sinR.
To investigate whether SlrR and SinR form a heteromeric complex, we carried out the following pull-down experiments. We mixed a lysate from Escherichia coli cells that were engineered to produce a His6-SinR fusion protein with a lysate of a B. subtilis strain that was deleted for sinR and hence overproduced SlrR. We applied the mixture to Ni-NTA agarose beads (New England Biolabs). Using antibodies that cross-react with both SinR and SlrR (Chai et al. 2009), we detected His6-SinR as well as SlrR in the eluate from the beads [Fig. 6A]. Indeed, SlrR was retained almost quantitatively on the beads, with relatively little present in the flow-through. In a control experiment with an E. coli lysate from cells lacking His6-SinR, little or no SlrR was found in the eluate from the beads [Fig. 6A]. These results indicate that SlrR forms a complex with SinR.

A heteromeric complex of SlrR and SinR binds to the promoters for autolysin genes

To investigate whether the SlrR•SinR complex binds to the promoter region of autolysin genes, we carried out electrophoretic mobility shift assays (EMSA). We initially used a radiolabeled DNA probe that contained the promoter region for the lytABC operon. The EMSA experiments were carried out with purified SinR and a purified SlrR fusion protein, glutathione S-transferase (GST)-SlrR, in which SlrR was joined to GST. First, we tested SinR alone, which was unable to cause a mobility shift, even at the highest concentration of protein used [1000 nM] [Fig. 6B]. Next, we tested GST-SlrR alone, which did cause the appearance of two species with retarded mobility [Fig. 6C]. However, even at the highest concentration of protein [1000 nM] not all of the probe was shifted [Fig. 6C, top panel, right lane].

We then investigated whether SinR would stimulate the ability of GST-SlrR to bind to the promoter sequence. To do this, we carried out the EMSA with increasing amounts of GST-SlrR, but this time in the presence of a fixed amount of SinR [Fig. 6D]. The results show that the presence of SinR [500 nM], which by itself had not shifted the DNA probe [Fig. 6B, top panel, fifth lane], greatly stimulated the DNA-binding capacity of GST-SlrR, almost all of the DNA probe was shifted, even at the very lowest concentration of GST-SlrR [63 nM] [Fig. 6D].

Conversely, we carried out a gel shift experiment in which we added increasing amounts of SinR in the presence of a fixed but low amount of GST-SlrR [63 nM], which by itself had shifted only a small amount of the DNA probe [Fig. 6C, top panel, second lane]. As shown in Figure 6E, increasing amounts of SinR caused a substantial proportion of the probe to be shifted when the low amount of GST-SlrR was used [Fig. 6E]. In other words, much more of the probe was shifted by the low amount of GST-SlrR when increasing amounts of SinR were added.

Similar results to those described above were obtained when we used as a probe DNA containing the lytF promoter (Supplemental Fig. S3A). Finally, and as a negative control, we used as a probe DNA that contained the

**Figure 5.** Amino acid substitutions in SinR that promote chaining are clustered in two domains. Two residues (V26 and A27) are located in the DNA-binding domain, whereas the other five (D55, E67, E79, D84, and A85) are clustered in the multimerization domain that is responsible for SinR–SinR interactions.

**Figure 4E** shows that the resulting, doubly mutant cells had lost the robust chaining phenotype of the parent strains [Fig. 4D]. We conclude that, like wild-type SinR, the capacity of the mutant SinR proteins to promote chaining depends on SlrR.

Both the SinR–SinR interaction and the DNA-binding domains of SinR are believed to be needed for SinR to adhere to its operators (Lewis et al. 1996, 1998; Kearns et al. 2005). Thus, a simple explanation for the classic filamentous phenotype of certain sinR alleles is that the amino acid substitution mutant proteins are impaired in SinR–SinR interaction or in recognition of the slrR promoter and are therefore defective in repressing slrR (which results in overexpression of slrR), but are not defective in promoting cell chaining. In other words, we hypothesize that SinR has two functions: repression of slrR (and the yqxm and eps operons), and a separate function that promotes cell chaining, presumably by contributing to the repression of autolysin genes. Evidence presented below indicates that this second function is to bind to SlrR to create a heteromeric complex that represses autolysin genes.

**SlrR interacts with SinR**

*In toto,* the results so far indicate that both SlrR and SinR are required for chaining. How are we to explain this requirement for both proteins? We speculated (1) that SlrR and SinR bind to each other to form a heteromeric complex, and (2) that such a complex [but neither SlrR nor SinR alone] is needed for repressing autolysin genes. In this section, we investigate whether the proteins bind to each other, and in the next two sections, we investigate whether the hypothesized complex binds to the promoter region of autolysin genes.
neither SlrR nor SinR alone, nor a mixture of two proteins, was able to shift the control DNA at all of the concentrations tested (Figs. 6B–D, bottom panels). We conclude that SlrR binds specifically to the promoter regions for the two autolysin genes tested, and that this binding is substantially stimulated by the presence of SinR. In toto, the results are consistent with the idea that the repressor for *lytABC* and *lytF* is an SlrR·SinR heteromeric complex.

An SlrR·SinR heteromeric complex covers the promoter for *lytABC*.

Next, we carried out a DNase I footprinting experiment to map the binding site for the heteromeric SlrR·SinR complex in the *lytABC* promoter region. When equal molar concentrations of GST-SlrR and SinR were incubated with the DNA probe, we observed bands that were protected from DNase I and that were hypersensitive to the nuclease over a large region (~47 base pairs [bp]). These bands extended from about the transcription start site to ~46 bp upstream of the start site (indicated by bars and asterisks in Fig. 6F). Thus, the complex almost entirely covered the promoter for *lytABC*. In comparison, no significant protection was observed when footprinting was carried out with GST-SlrR alone (Fig. 6F, middle two lanes) or SinR alone (data not shown).

Within the region protected by GST-SlrR and SinR, we observed two putative SinR operator-like sequences (indicated as SinR box) and two identical repeats. Within the protected region, two SinR operator-like sequences (indicated as SinR box) and two identical repeats are labeled. Also labeled are the −35 and −10 regions and the +1 start site for the αD-dependent promoter. (G) EMSAs using a radiolabeled DNA probe for the intergenic control region for *slrR* and *eps* (*P*~slrR/eps~), and purified GST-SlrR fusion protein and SinR. In the left panel, increasing amounts of GST-SlrR were added [0, 31.2, 62.5, 125, 250, and 500 nM]. In the middle panel, increasing amounts of SinR were added [0, 15.6, 31.2, 62.5, 125, and 250 nM]. In the right panel, a fixed amount of SinR (125 nM) was mixed with increasing amounts of GST-SlrR [0, 31.2, 62.5, 125, 250, 500 nM].
results is that two molecules of SlrR bind to the repeated AATATAA sequences, and two molecules of SinR bind to the two SinR-like binding sequences.

SlrR inhibits the binding of SinR to the control regions for slrR, eps, and yqxM

We showed that, by binding to SinR, the SlrR protein confers a new property on SinR, namely, the capacity to bind to and repress autolysin genes. We wondered whether the binding of SlrR to SinR might also alter the affinity of SinR for the operators that it otherwise binds to on its own, namely, those for slrR, eps, and yqxM. To investigate this, we carried out EMSAs using as a probe DNA containing the ~200-bp intergenic region between eps and slrR. The eps operon and the slrR gene are divergently transcribed from a common control region that contains multiple SinR-binding sites (Kearns et al. 2005). Increasing amounts of GST-SlrR (from 0 to 500 nM) was not able to shift the DNA probe [Fig. 6G, left panel], whereas increasing amounts of SinR (from 0 to 250 nM) by itself were highly effective in retarding the mobility of a DNA probe that contained the control region for slrR and eps, causing the formation of higher-order, slow-migrating species [Fig. 6G, middle panel]. However, when increasing amounts of GST-SlrR (from 0 to 500 nM) were added to a fixed amount of SinR (125 nM) that by itself was able to fully cause the formation of the higher-order complex, the gel shift was reversed with the partial reappearance of a free probe and the elimination of the slow-migrating species [Fig. 6G, right panel, right two lanes]. Thus, SlrR inhibits the capacity of SinR to bind to the control region for its own gene (slrR) as well as for eps. We wondered whether SlrR would likewise antagonize the binding of SinR to the yqxM promoter. Indeed, as shown in Supplemental Figure S4A, the addition of SlrR reversed the binding of SinR to DNA containing the yqxM promoter (Supplemental Fig. S3C).

The above results suggest that SinR, SlrR, and slrR are components of a self-reinforcing, double-negative feedback loop in which SlrR antagonizes (titrates) SinR, thereby stimulating derepression of slrR. Derepression would result in yet more SlrR production, thereby further enhancing derepression of slrR as well as the matrix operons eps and yqxM. Indeed, and in support of this hypothesis, we found that, in a strain that contains both a PslrR-lacZ fusion and an IPTG-inducible copy of slrR, the activity of PslrR-lacZ was higher in the presence of the inducer than in its absence [Supplemental Fig. S4]. These findings also explain the previous observation that overexpression of slrR increases the expression of eps and yqxM (Kobayashi 2008; Murray et al. 2009).

As reported previously, biofilm formation is triggered by the production of the anti-repressor SinI, which inhibits SinR (Kearns et al. 2005; Chai et al. 2008). We now propose that SinR is subject to anti-repression by the successive action of SinI and then SlrR. In other words, SinI sets in motion a chain of events that leads to the appearance of an additional anti-repressor, SlrR, which augments the action of SinI.

SinR and SlrR control cell chaining during exponential phase growth

Cells in the mid-exponential phase of growth are a mixture of two cell types: motile cells, and chains of sessile cells (Kearns and Losick 2005). We wondered whether the SlrR•SinR complex is also responsible for chaining under these conditions. We therefore examined the chaining phenotype of growing cells during exponential phase for the wild type and for δsinI, δsinR, and δslrR mutants. We performed the analysis in two genetic backgrounds: that of 3610, an undomesticated strain used for biofilm studies, and that of PY79, a laboratory strain. 3610 exhibits a high level of σD activity and is biased toward motile cells, whereas PY79 (due to a mutation in swrA) exhibits lower σD activity and produces a relatively high proportion of chains (Kearns and Losick 2005). We also introduced a PbgaD-gfp fusion [green in Fig. 7] into the strains as a reporter for σD-directed transcription. The cells were stained with the dye FM4-64 to visualize membranes (red in Fig. 7).

As reported previously (Kearns and Losick 2005), both wild-type strains exhibited a bimodal pattern of cell types: single cells that were ON for PbgaD-gfp, and chains of cells that were OFF [Fig. 7]. Also, as expected, the proportion of chains was higher for PY79 (wild type in Fig. 7A) than for 3610 (wild type in Fig. 7B). Importantly, the proportion of cell chains was markedly reduced or completely absent in all three mutants in both the 3610 and PY79 backgrounds [Fig. 7]. Thus, SinR and SlrR control cell chaining during exponential-phase growth as well as during biofilm formation.

We draw two inferences from these results. First, lytABC and lytF are not the only targets of the SlrR•SinR complex.
repressor. The results of Figure 7 suggest that the σD-controlled hag gene, which encodes flagellin (LaVallie and Stahl 1989), is also under the negative control of SlrR•SinR. In confirmation of this inference, hag gene expression was decreased drastically throughout the population in the cell chaining-promoting amino acid substitution mutants of SinR described above [data not shown], and EMSAs showed that the SlrR•SinR complex binds to the promoter region for hag [Supplemental Fig. S3B]. In toto, these findings raise the possibility that the SlrR•SinR complex is a repressor of additional, and perhaps many, members of the σD regulon.

The second inference is that the bimodal distribution of σD-directed transcription is governed by the SinR SlrR double-negative feedback loop [Fig. 1B]. We propose that the loop similarly creates a bistable switch that is responsible for switching σD-transcribed genes ON or OFF. The switch can be biased in one direction or the other by σD levels, as demonstrated by the effects of mutations of swrA, a gene that stimulates transcription of the operon that encodes σD (Kearns and Losick 2005). Nonetheless, swrA itself is not the basis for bistability in σD-directed transcription. Instead, we propose that the self-reinforcing cycle of SlrR synthesis contributes to, or is entirely responsible for, determining whether σD-directed transcription of autolysin and motility genes is ON or OFF.

The double-negative feedback loop exhibits hysteresis

If the SinR SlrR loop is a bistable switch, then it should exhibit hysteresis (Veening et al. 2008); that is, once activated, the circuit should remain on for long periods of time in the absence of the original stimulus. To address this question, we introduced the IPTG-inducible time in the absence of the original stimulus. To address this question, we introduced the IPTG-inducible

As shown in Figure 8, before induction, YC281 cells exhibited a bimodal distribution of single cells and chains, with a strong bias toward single cells [Fig. 8A [top], G [column a]]. YC280 cells showed little chaining before induction due to the Δslr mutation [Fig. 8A [bottom], H [column a]]. After growth in the presence of the inducer for 60 min, both strains showed greatly increased levels of chaining [Fig. 8B [top and bottom], G,H [columns b]]. The cells were then washed to remove IPTG and diluted into fresh LB medium at hourly intervals to maintain exponential-phase growth. At 120 and 180 min after washing (two fourfold dilutions into fresh medium), the YC281 (slrR+) cells showed even more extensive cell chaining than before washing [Fig. 8D, E [top], G [columns d,e]]. Moreover, and with additional rounds of dilution and growth in fresh medium, a high level of chaining could still be seen for as long as 5 h after washing [Fig. 8F [top], G [column f]], representing >10 generations of growth. In contrast, chaining by the control strain YC280 (ΔslrR) was not sustained after the removal of the inducer, with the cells returning to the short chain or single-cell state they were in before treatment with the inducer [Fig. 8D-F [bottom], H [columns d-f]]. These results indicate that, once activated by a pulse of SlrR synthesis, the double-negative feedback loop remains locked in a state in which cell separation (autolysin) genes remain OFF for multiple rounds of cell division, that is, the loop exhibits hysteresis.

Discussion

The principal contribution of this investigation is the discovery of an epigenetic switch that controls σD-directed transcription of genes for cell separation and motility during growth [Fig. 1]. The switch consists of the proteins SinR and SlrR and the gene for SlrR. Results with mutants of SinR and SlrR and with cells engineered to produce SlrR in response to an inducer indicate that the switch is both necessary and sufficient to explain the bistability of

Figure 8. The feedback loop exhibits hysteresis. (A, top panel) Cells of an undomesticated [swrA+] strain bearing an IPTG-inducible copy of slrR (YC281) were largely in the form of single cells when grown in LB to mid-exponential phase in the absence of an inducer. (Bottom panel) A derivative of the above strain that was mutant for slrR (YC280) contained few chains under the same conditions. (B) After treatment with 100 μM IPTG for 60 min to induce SlrR production, the proportion of chains markedly increased for both strains. (C–E) The cells were washed to remove IPTG, and samples were withdrawn and diluted fourfold into fresh LB medium at hourly intervals. At 120 and 180 min after suspension, cells of YC281 (top panels in D,E) were still largely in the form of chains, whereas cells of YC280 (bottom panels in D,E) had largely reverted back to single cells. (F, top panel) After further rounds of growth and suspension in fresh LB medium, the YC281 cells were still largely in the form of chains. (G,H) Quantitative results showing the ratio of chains versus single cells for YC281 (G) and YC280 (H) cells before and after induction shown in A–F.
of cell separation and motility in growing cells. We propose that the SinR SlrR switch exists in two alternative stable states: a state in which SlrR levels are high ([SlrR HIGH state]), and a state in which SlrR levels are low ([SlrR LOW state]) [Fig. 1B]. Autolysin and motility genes are OFF in the SlrR HIGH state and are ON in the SlrR LOW state.

Like the classic epigenetic switch of phase λ in which the CI repressor represses the gene for Cro and the Cro repressor represses the gene for CI (Oppenheim et al. 2005), the SinR SlrR switch is a double-negative feedback loop. Unlike the λ switch, however, the SinR SlrR switch involves protein–protein (SinR-mediated inhibition of SlrR) and protein–DNA (SinR-mediated repression of slrR) interactions. Also, like the well-studied ComK bistable switch, which governs entry into the state of genetic competence in B. subtilis (Maamar and Dubnau 2005; Maamar et al. 2007), the SinR SlrR switch is self-reinforcing in that SlrR inhibits the capacity of SinR to repress the gene for SlrR. (In effect, SlrR titrates SinR, thereby derepressing slrR.) The ComK switch is, however, a positive feedback loop in which ComK directly stimulates its own synthesis, as opposed to a double-negative loop in which one component (SlrR) indirectly stimulates its own synthesis (Maamar and Dubnau 2005). Finally, the SinR SlrR switch differs from these other examples of bistable switches in that it consists of three components (SinR, SlrR, and the SinR target slrR), whereas the phase λ switch has four components (CI, Cro, and the DNA-binding sites for the two repressors) and ComK has two components (ComK and its binding site). A virtue of the three-component switch is that it not only creates bistability, but also repurposes one of the components: SinR.

During biofilm formation, SlrR synthesis is triggered by the synthesis of the anti-repressor SinI, which inhibits SinR, as we discuss below (Kearns et al. 2005; Chu et al. 2008). How, then, is SlrR synthesis triggered during growth in rich medium, when the levels of SinI are low? We propose that the switch is activated stochastically by noise in the expression of these same components. For example, under conditions of exponential-phase growth in which Spo0A−P levels are low, sinI is largely silent in the cell population as a whole. Yet sinI must contribute to the switching because a sinI mutant is locked in the SlrR LOW (autolysin and motility genes ON) state in exponential-phase cells. We therefore hypothesize that stochastic and transient expression of sinI is a feature of exponentially growing cells. In only a subpopulation, enough SinI accumulates to cause derepression of slrR, thereby triggering self-reinforcing synthesis of SlrR. This process could be aided by noise in the expression of slrR (that is, escape of slrR from SinR-mediated repression), which could push some cells to the brink of triggering self-reinforcing synthesis of SlrR. Triggered by stochastic events, self-reinforcing synthesis of SlrR would switch the SinR SlrR loop into the SlrR HIGH state [autolysin and motility genes OFF] and lock a subpopulation of cells in the form of chains [Fig. 1B].

We began this investigation by attempting to elucidate the mechanism that governs cell chaining during biofilm formation. We discovered, however, that the same circuitry that governs cell chaining in biofilms also controls cell chaining in growing cells, but with one important difference. During biofilm formation, the switch is not activated stochastically. Rather, it is driven into the SlrR HIGH state by the programmed synthesis of the anti-repressor SinI [Fig. 1B], that is, in biofilm-inducing medium, the gene for the SinI anti-repressor is turned on by Spo0A−P, a global regulator of post-exponential-phase gene expression (Fujita et al. 2005; Chai et al. 2008). Spo0A−P-directed synthesis of SinI results in inhibition of SinR, and hence depression of the gene for SlrR [Fig. 1A]. Thus, the activation of Spo0A−P in stationary phase sets in motion a cascade in which SinI is successively inhibited by SinI and then by SlrR, resulting in derepression of SinR targets (i.e., slrR, eps, and yqxm) and repression of SlrR targets (i.e., lytABC and lytF) through the formation of the SinR•SlrR complex. In other words, during biofilm formation, expression of slrR is not left to chance; it is a downstream consequence of the activation of Spo0A−P.

Fascinatingly, SinI, SinR, SlrR, and SlrA are members of a family of homologous proteins that are linked to each other through intricate circuitry. We did not consider SlrA here because it does not contribute importantly to biofilm formation or cell chaining under laboratory conditions (unless it is artificially overexpressed) (Chai et al. 2009). Its synthesis is believed to be governed by an unknown environmental signal (Kohayashi 2008; Chai et al. 2009). Nonetheless, it is a paralog of SinI, and, like SinI, it is an anti-repressor for SinR (Chai et al. 2009). SlrR is itself a hybrid protein, with domains that are homologous to SinI and SlrA on the one hand, and with domains that are homologous to SinR on the other hand (Chu et al. 2008). In fact, then, SinR is subject to inhibition by three anti-repressors: SinI, SlrA, and, as we observed, SlrR. Meanwhile, SinR, on its own, is a repressor of a particular set of promoters [slrR, eps, and yqxm], but can be repurposed by SlrR to bind to an entirely new set of promoters [lytABC, lytF, hac], and, perhaps, those for additional ρ^− transcribed genes] by forming a SinR•SlrR heterocomplex.

In summary, a single, simple regulatory circuit is deployed by B. subtilis to control gene expression under different biological circumstances. Under conditions [stationary phase in poor medium] that promote the formation of architecturally complex communities, the SinR SlrR switch is part of a regulatory cascade that culminates in the formation of long chains of cells and in the production of an extracellular matrix that binds them together. Conversely, under conditions of exponential-phase growth in rich medium, the same circuit is an epigenetic switch that is activated stochastically to lock cells in alternative states of motility and cell chaining.

Materials and methods

General methods for media, strain construction, β-galactosidase assays, protein production, and purification

See the Supplemental Material for general methods for media, strain construction, β-galactosidase assays, protein production, and purification.
Strains
See Supplemental Table S1 for a list of strains.

Colony and pellicle formation
For colony formation, cells were first grown to exponential growth phase in LB broth, and 3 μL of these cultures were spotted onto solid MSgg medium containing 1.5% Bacto agar. The plates were incubated at 23°C. Images of the colonies were taken using a SPOT camera [Diagnostic Instruments]. For pellicle formation, cells were grown to exponential phase, and 9 μL of cultures were mixed with 9 mL of liquid medium in a six-well microtiter plate (VWR). Plates were incubated for 2–3 d at 23°C. Images of the pellicles were recorded similarly.

Microscopy analysis
For assays of cell chaining during pellicle formation, cells were cultured as described above. Cells were collected from pellicle-forming wells after 48 h of incubation and were washed twice with cold PBS buffer. Cells were suspended in 50 μL of cold PBS buffer and were analyzed using phase-contrast microscopy. For assays of cell chaining in shaking culture conditions, cultures were collected in either early stationary phase when grown in MSgg medium, or mid-exponential growth phase when grown in LB medium at 37°C. Cells were treated as described above for microscopy analysis. To induce expression of slrR from the hy spank promoter, 500 μM IPTG (unless otherwise indicated) was added to cells grown to early exponential phase in broth medium, and the cells were allowed to continue to grow with shaking for one more hour at 37°C before being harvested.

For fluorescence microscopy analysis, cells were grown in either MSgg broth to early stationary phase or LB medium to mid-exponential phase. One milliliter of the culture was harvested and washed with 10 mL of cold phosphate buffer. Cells were suspended in 50 μL of PBS buffer. One microliter of diluted membrane-staining dye FM4-64 was mixed with suspended cells. Three microliters of FM4-64-treated cells were dropped to the center of an agar-coated microscopy slide. Cover slides were pretreated with poly-l-lysine (Sigma). Samples were examined using an Olympus workstation BX61. Images were taken using an automated software program [SimplePCI] and analyzed with MetaMorph [Universal Imaging Corporation].

Screen for chaining-promoting mutants of SinR
B. subtilis cells mutant for ymcA are severely impaired for biofilm formation [Brandt et al. 2004]. We discovered previously that, after prolonged incubation, a ΔymcA mutant was able to form pellicles due to the appearance of suppressor mutations [Kearns et al. 2005]. Frequently, the suppressor mutations were mapped to sinR. We also discovered that, often, those suppressors were rescued not only for matrix production, but also for cell chaining. A strain (YC581) was constructed as described above that contained a null mutation in ymcA (ΔymcA) and a kanamycin resistance gene was inserted into a chromosomal locus immediately downstream from sinR. The purpose of introducing the kan gene was to facilitate the transfer of suppressor mutations in sinR to another genetic background. The kan gene had no effect on biofilm formation and cell chaining (data not shown). After prolonged incubation of YCS81 cells in standing MSgg medium, we were able to pick suppressors that formed pellicles. Among the suppressors obtained, 12 contained mutations in the sinR gene. Nine of the 12 suppressor mutations were found to rescue both matrix production and chaining, while the remaining three rescued matrix production, but not chaining, and they were not further characterized. All nine suppressor mutants contained point mutations that caused single-amino-acid substitutions in the SinR protein (Fig. 5; A85T substitution was obtained independently twice).

Pull-down experiment
The cleared lysate containing His6-SinR fusion proteins was prepared from the E. coli cells as described above, and was mixed in equal volume with cleared lysate from a B. subtilis ΔsinR mutant (RL3856). Preparation of cleared lysate from B. subtilis RL3856 was as follows. Cells were inoculated into 100 mL of MSgg broth and grown at 37°C to early stationary phase. Cells were harvested and washed with 10 mL of cold phosphate buffer [20 mM sodium phosphate, 200 mM NaCl, 10% glycerol, 1 mM PMFSF at pH 7.4]. Cell pellets were suspended in 1 mL of cold phosphate buffer supplemented with 500 μg mL−1 of freshly made lysozyme solution and were incubated for 30 min on ice. Cells were further disrupted using sonication on ice. Cell lysate was centrifuged at 5000 rpm for 5 min to remove cell debris, and were further centrifuged at 14,000 rpm for 30 min at 4°C. The mixture of the E. coli and the B. subtilis lysates was incubated for 30 min at 4°C, applied to Ni-NTA agarose beads, and incubated for one more hour at 4°C with rotation. Proteins were eluted afterward, following the same procedures that were described above. Immunodetection of SinR and SlrR proteins was performed following a protocol that was published previously [Chai et al. 2009]. The control experiment was performed similarly except that the E. coli lysate does not contain His6-SinR.

EMSA
The DNA probes for the promoter sequences of lytABC, lytF, and slrR were generated by PCR using chromosomal DNA of 3610 and primers PlytA-F1 and PlytA-R1, PlytP-F1 and PlytP-R1, and PlylR-F1 and PlylR-R1, respectively. See Supplemental Table S2 for a description of the primers. The DNA probes were digested with EcoRI, gel-purified, and filled in using klenow [exo−], DTT, and [α-32P]-dATP [New England Biolabs]. EMSA was conducted following a protocol that had been described previously [Chai et al. 2009]. When both SinR and GST-SlrR were applied in the assay, proteins were mixed and incubated for 15 min on ice first before the DNA probe was added.

DNase I footprinting
The DNA probe containing the promoter region for the lytABC operon was amplified and end-labeled as described above. Radiolabeled DNA was mixed with purified SinR and or GST-SlrR at indicated concentrations, and the mixture was incubated in 50 μL of binding buffer [20 mM Tris buffer at pH 8.0, 5 mM MgCl2, 0.1 mM DTT, 50 μg mL−1 BSA, 5 μg mL−1 poly-dI–dC, 10% glycerol] for 20 min on ice. The mixture was warmed up for 10 min at room temperature. To each mixture, 50 μL of digestion mix—which contains 0.05 U of DNase I [amplification grade, Invitrogen], 10 μL of 10× DNase I digestion buffer, and 0.3 mM DTT—was added and quickly mixed. The mixture was incubated for 30 sec at room temperature. The reaction was stopped by addition of 5 μL of 200 mM EDTA. DNAs were precipitated by addition of 2.5 vol of cold ethanol and 10% of 3 M sodium acetate [pH 5.3], and were kept for 1 h at −80°C. DNAs were resuspended in 10 μL of formamide loading buffer, and 5 μL of the sample were loaded to 6% TBE sequencing gel [National Diagnostics].
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An epigenetic switch
An epigenetic switch governing daughter cell separation in *Bacillus subtilis*

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