



Noise in Gene Expression Determines Cell Fate in *Bacillus subtilis*

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Science **317**, 526 (2007);

DOI: 10.1126/science.1140818

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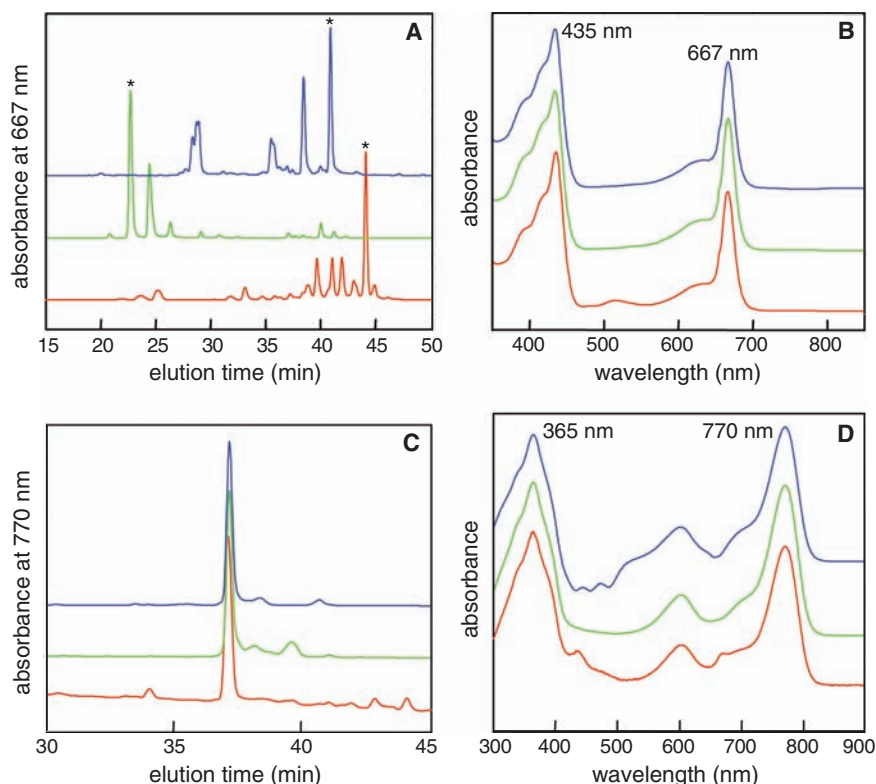


Fig. 4. HPLC elution profiles monitored at 667 nm for BChl c (**A**) and 770 nm for BChl a (**C**) for pigments extracted from a light-grown culture containing *Cab. thermophilum* and *Anoxybacillus* sp. (red line); *Chlorobium tepidum* (green line); and *Chloroflexus aurantiacus* strain Y-400-fl (blue line). The spectra of the BChl c peaks indicated by stars in (**A**) are shown in (**B**); the spectra of the BChl a peaks eluting at 37 min in (**C**) are shown in (**D**). All other peaks in (**A**) also had the absorption spectrum of BChl c.

a and c and produces chlorosomes under oxic conditions. Although chlorosomes are also found in some *Chloroflexi*, the new chlorophototroph is the only described organism outside the phylum *Chlorobi* that produces FmoA. Growth of this acidobacterium is strongly stimulated by light under photoheterotrophic conditions, but additional studies will be required to establish whether the organism is capable of autotrophic growth. Because no chlorophototroph with these properties has yet been described, we propose the name “*Candidatus* Chloracidobacterium thermophilum,” gen. nov., sp. nov., for this BChl-synthesizing, phototrophic member of the phylum *Acidobacteria*.

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25. This work was supported by NSF grant MCB-0523100 and U.S. Department of Energy grant DE-FG02-94ER20137 (to D.A.B.). The metagenomics database used in this work was created under the auspices of grant EF-0328698 from the Frontiers in Integrative Biology Program of the NSF (to D.M.W.), who also acknowledges support from the NASA Exobiology Program (NAG5-8824). D.A.B. thanks S. Schuster, J. Eisen, R. Blankenship, R. Castenholz, and S. Giovannoni for helpful discussions and comments on the manuscript. D.A.B. and D.M.W. also gratefully acknowledge the Thermal Biology Institute of Montana State University (NASA NAG5-8807) for support and hospitality while D.A.B. was a visiting fellow in summer 2005, when this work was initiated. We thank M. Melendrez and Q. Tao for preparation of the BAC library. The *pscAB-fmoA*, *recA*, and *rRNA* genes were encoded on a single BAC clone, M60-018 J19: GenBank accession number EF531339. *Roseiflexus* sp. isolate, strain RS1: GenBank accession number AAU000000000. *Synechococcus* sp. OS-A and OS-B' genome sequences: GenBank accession numbers CP000239 and CP000240.

Supporting Online Material

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30 March 2007; accepted 27 June 2007
10.1126/science.1143236

Noise in Gene Expression Determines Cell Fate in *Bacillus subtilis*

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Random cell-to-cell variations in gene expression within an isogenic population can lead to transitions between alternative states of gene expression. Little is known about how these variations (noise) in natural systems affect such transitions. In *Bacillus subtilis*, noise in ComK, the protein that regulates competence for DNA uptake, is thought to cause cells to transition to the competent state in which genes encoding DNA uptake proteins are expressed. We demonstrate that noise in *comK* expression selects cells for competence and that experimental reduction of this noise decreases the number of competent cells. We also show that transitions are limited temporally by a reduction in *comK* transcription. These results illustrate how such stochastic transitions are regulated in a natural system and suggest that noise characteristics are subject to evolutionary forces.

Variability in gene expression within a population of genetically identical cells enables those cells to maintain a diver-

sity of phenotypes, potentially enhancing fitness (1, 2). When the underlying gene network contains regulatory positive feedback loops, indi-

vidual cells can exist in different states; some cells may, for example, live in the “off” expression state of a particular gene, whereas others are in the “on” expression state (this is an example of bistable gene expression). These stochastic fluctuations in gene expression, commonly referred to as noise, have been proposed to cause transitions between these states (3–7). We apply recently developed theories of noise (8, 9) to examine how noise influences these transitions in a natural system.

An example of bistable expression with associated stochastic transitions (10–16) involves the ability of the soil bacterium *Bacillus subtilis* to develop “competence” for DNA uptake as it enters stationary growth phase, potentially allowing bacteria to increase their fitness by incorporating new genetic material. The genes needed for competence are transcribed only in the presence of ComK, the master regulator of competence. *comK* expression is subject to positive autoregulation effected by the cooperative binding of ComK to its own promoter (Fig. 1A) (17–19), resulting in bistability (5). In one state, the positive autoregulatory loop is not activated and *comK* expression is low, and in the other state, the loop is activated because the level of ComK has exceeded a critical threshold and *comK* expression is high (13, 14).

The capacity for bistability is subject to temporal regulation. While the cells are growing exponentially, the level of ComK is kept low (i) through the action of the MecA-ClpC-ClpP protease complex, which actively degrades the ComK protein, and (ii) by transcriptional repressors such as Rok, AbrB, and CodY, precluding transitions to the competent state. Upon reaching stationary phase, the accumulation of an extracellular peptide causes an increase in the expression of the ComS protein (20) (the time of the onset of stationary phase is denoted as T_0) (Fig. 1B). ComS competes with ComK for binding to the MecA-ClpC-ClpP complex (21), effectively lowering the rate of ComK degradation and allowing random fluctuations in the level of ComK to occasionally cause transitions to the competent state. Cells continue to randomly transition to competence for 2 hours, by which time (T_2) transitions have ceased to occur (16) and the 15% of the cells that have become competent remain so until diluted into fresh growth medium (Fig. 1C and movie S1). In this report, we ask why cells only transition to competence for a limited duration of time and investigate the source of the fluctuations that actuate the ComK feedback loop in a minority of cells.

To understand why cells only transition to the competent state for ~2 hours during stationary phase, we examined the dynamics of *comK* expression in noncompetent cells. Because the level of ComK in noncompetent cells is very low, we used fluorescence in situ hybridization (FISH) to count individual *comK* mRNA molecules in single cells (22–24). We achieved this level of sensitivity by using six fluorescently labeled single-stranded DNA probes, complementary to different regions of the *comK* mRNA (Fig. 2A, left). The hybridization of many fluorophores to individual mRNA molecules resulted in spots that are visible through a fluorescence microscope (Fig. 2, B and C).

During late exponential and early stationary phase in the wild-type (WT) strain, the mean number of *comK* mRNA molecules increased from 0.7 to 1 molecule per cell at T_0 , at which point transitions to competence begin to occur. Thereafter, the average number of *comK* transcripts per noncompetent cell declined to 0.3 molecules per cell at T_2 (Fig. 2, D and E).

We postulated that in early stationary phase, when the average mRNA level is elevated, the probability of transition is high because of the increased likelihood of randomly generating enough ComK to activate the positive feedback loop. Later in stationary phase, when the average is low, the probability of such an accumulation is much smaller. To test this possibility, we counted the number of *comK* mRNA molecules in a strain that cannot synthesize Rok, the transcriptional repressor of *comK*. Inactivation of *rok* does not change the temporal pattern of competence expression but markedly increases the fraction of competent

cells (14). The average number of *comK* mRNA molecules per noncompetent cell in this strain was twice as large as that in the WT strain at T_0 (Fig. 2D), in accordance with the larger number of competent cells observed in the *rok* strain (14, 25).

It is possible that the increased rate of transition to competence observed at T_0 is not caused by the increased basal rate of *comK* transcription but is rather the cause of the increased transcription rate, because of positive feedback at the *comK* promoter. This possibility was eliminated by measuring the *comK* promoter activity in a strain lacking a functional *comK* gene but instead having the *comK* promoter drive a sequence consisting of a 50-base pair (bp) motif repeated 32 times (*comK-M2*). The corresponding mRNA was detected with a single-stranded DNA probe complementary to the 50-bp sequence (Fig. 2A, right). We found that the number of these mRNAs in a strain lacking an active *comK* gene was similar to that in the WT strain (Fig. 2E), indicating that positive feedback does not play a role in *comK* expression in noncompetent cells. To further verify that the *comK-M2* construct had similar expression properties to those of the endogenous *comK* mRNA, we integrated the construct in the WT strain and simultaneously measured the abundance of transcripts from both the endogenous *comK* gene and the *comK-M2* gene, using differently colored fluorophores to label the two mRNAs. The mean and variance of the numbers of the two transcripts were almost identical (fig. S1).

To verify that the observed decrease in *comK* transcripts during stationary phase could account for a decrease in the rate of transition to competence, we constructed a simple stochastic model of

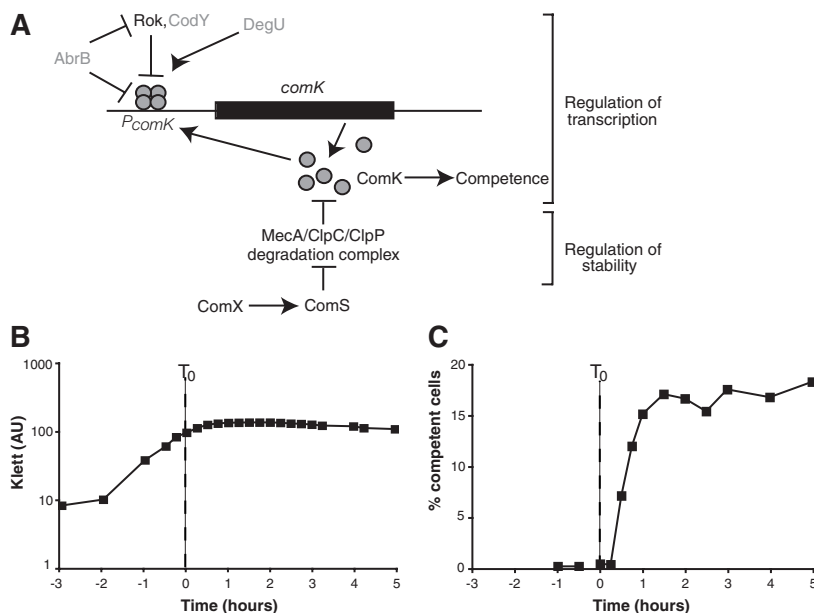


Fig. 1. The regulation of competence in *B. subtilis*. (A) The *comK* regulatory network. Arrows and perpendiculars represent positive and negative regulation, respectively. For simplicity, factors shown in gray were not considered in our modeling. (B) The kinetics of growth in competence medium [in absorbance units (AU) measured in a Klett colorimeter]. (C) Competence development, determined microscopically with strains carrying a *comK-cfp** fusion. The dashed lines in (B) and (C) represent T_0 .

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the *comK* positive feedback loop containing the salient features of the competence network, most notably the positive feedback loop [see the supporting online material (SOM)]. The model confirmed the plausibility of our conclusion that a relatively small decrease in *comK* transcription can effectively end transitions to the competent state (fig. S4).

Together, these data suggest that temporal regulation of transcription controls the frequency of transitions to the competent state and that the decline in transcription of *comK* during stationary phase effectively defines a “window of opportunity,” which explains why cells are only able to transition to competence for a limited amount of time.

Because the cells are genetically identical and grown in a well-stirred medium, the determination of which cells are selected for competence is likely due to random cell-to-cell variations in proteins involved in competence regulation. Given its critical role in the regulation of competence, we examined the role that

noise in *comK* plays in selecting cells for competence. Cell-to-cell variations in the numbers of *comK* mRNAs can come from two sources (26, 27): (i) intrinsically random events of transcription and mRNA decay (intrinsic noise) and (ii) cell-to-cell variations in regulators, polymerases, and other global factors (extrinsic noise). To gauge the relative contributions of these two types of noise to the fluctuations leading to competence, we used an approach derived from Elowitz *et al.* (26): counting the numbers of both endogenous *comK* mRNA and *comK-M2* mRNA in individual cells (Fig. 3, B and C). Because any extrinsic variations should affect both genes simultaneously, correlated variations between the mRNA numbers indicate that the variations are primarily extrinsic, whereas uncorrelated variations indicate an intrinsically stochastic origin for the fluctuations in mRNA numbers (Fig. 3A). In early stationary phase when most of the transitions occur, the numbers of mRNA

molecules from the two species were largely uncorrelated (correlation coefficient $r = 0.15$ at T_0) (Fig. 3, D and E). This finding indicates that intrinsically random fluctuations in *comK* mRNA production and degradation are likely to be a significant source of variations in ComK protein, leading to the initiation of competence (28).

To test the hypothesis that intrinsic noise is responsible for transitions to the competent state, we changed the noise in ComK protein production by altering the transcriptional and translational efficiency of *comK*. Recent studies have shown that intrinsic variations in protein expression are inversely related to the rate of transcription but are unaffected by the rate of translation (8, 9) (see SOM). Thus, increasing the rate of transcription of a gene while reducing the rate of translation by an equivalent amount would reduce noise in gene expression, despite having the same mean expression level. This reduction in noise should lead to fewer transitions to the

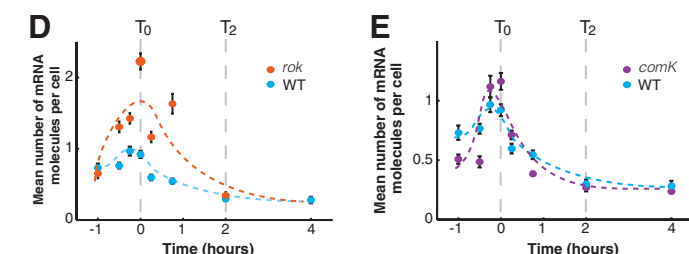
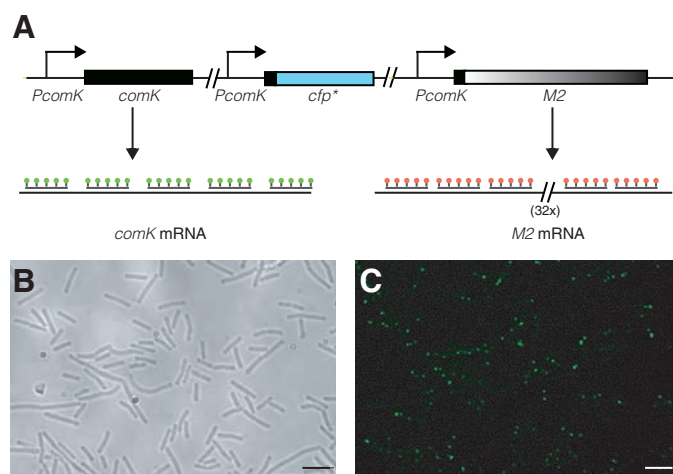


Fig. 2. Detection of single RNA molecules (*comK* and *comK-M2*) by FISH. (A) Schematic diagram depicting the endogenous *comK* (left), *comK-cfp** (middle), and *comK-M2* (right) reporters, all controlled by the *comK* promoter (*PcomK*). Multiple specific fluorescent probes bind to each mRNA molecule [*comK* (green) or *comK-M2* (red)], yielding distinct fluorescent signals. The *comK-cfp** construct identifies competent cells. *comK-cfp** designates an in-frame fusion of CFP to *comK*, and M2 designates the RNA with 32 repeat sequences. (B) Differential interference contrast (DIC) images and (C) pseudo-colored fluorescence images taken at T_0 for the WT strain, in which the *comK* mRNA was hybridized to six FISH probes [C6–tetramethyl rhodamine (C6-TMR)] that bind to the *comK* open reading frame. Dots correspond to individual mRNA molecules. Scale bars, 4 μm . (D and E) Kinetics of the population means of mRNA molecules per noncompetent cell before and after T_0 for the WT (blue circles, BD4379), the *rok* (red circles, BD4380), and the *comK* (purple circles, BD4382) strains. The WT and the *rok* strains (D) were hybridized to C6-TMR to detect *comK* mRNA molecules. The *comK* strain (E) was hybridized to a probe (PM2-Alexa 594) that binds to the M2 probe-binding sequence. Error bars were obtained by bootstrapping.

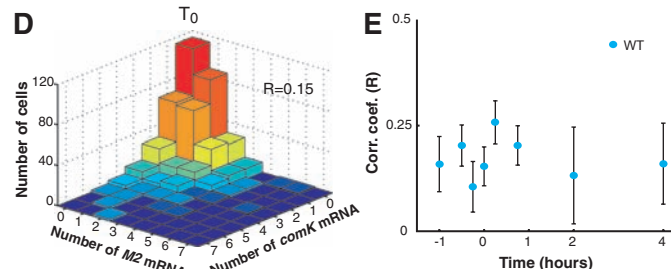
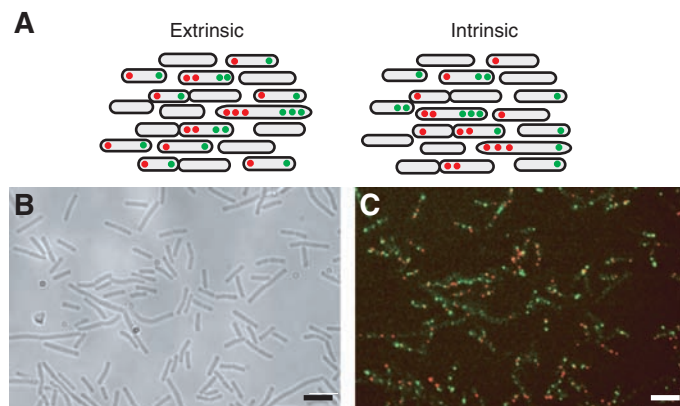


Fig. 3. Noise in *comK* transcription is mainly intrinsic. (A) Intrinsic and extrinsic noise were measured by detecting the mRNA from two coexpressed genes (*comK* and *comK-M2*) controlled by the *comK* promoter (26). Uncorrelated gene expression in individual cells is indicative of intrinsic noise. (B) DIC images and (C) pseudo-colored merged fluorescence images taken at T_0 showing hybridization to *comK-M2* (red) and *comK* (green) mRNA with the PM2-Alexa 594 and C6-TMR probes, respectively. Scale bars, 4 μm . (D) Distribution of *comK* and *comK-M2* mRNA molecules for the WT strain (BD4379) at T_0 , showing weak correlations between production of the two mRNA molecules ($r = 0.15$). (E) Correlation coefficients throughout growth for the same strain. Error bars indicate SE.

comK-M2 (red) and *comK* (green) mRNA with the PM2-Alexa 594 and C6-TMR probes, respectively. Scale bars, 4 μm . (D) Distribution of *comK* and *comK-M2* mRNA molecules for the WT strain (BD4379) at T_0 , showing weak correlations between production of the two mRNA molecules ($r = 0.15$). (E) Correlation coefficients throughout growth for the same strain. Error bars indicate SE.

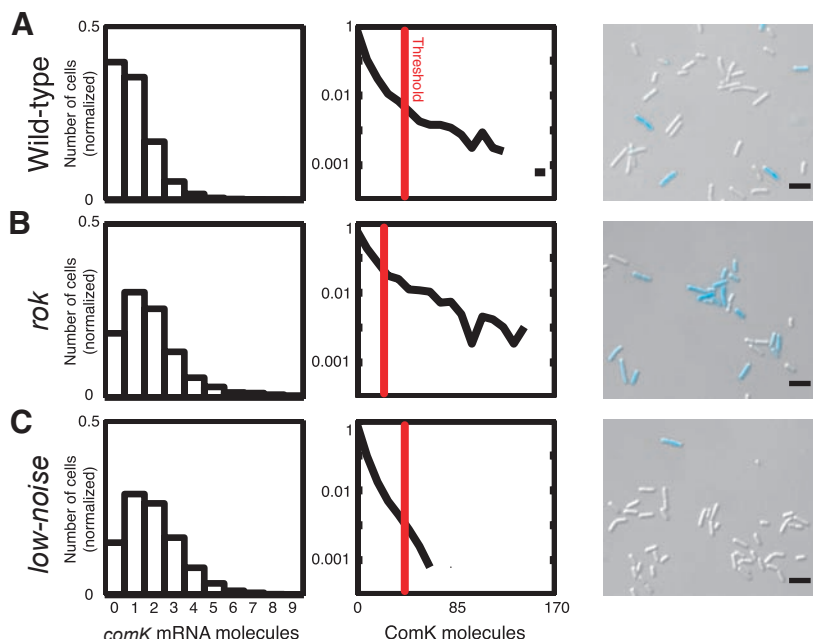


Fig. 4. Noise reduction in *comK* expression lowers the percentage of competent cells. The leftmost column depicts *comK* mRNA distributions predicted by the model for the WT (A), *rok* (B), and low-noise (C) strains at T_0 . The middle column shows ComK protein distributions at T_0 assuming a high rate of translation in the WT and *rok* strains [(A) and (B)] and a lowered rate of translation in the low-noise strain (C). The vertical red lines show the predicted threshold beyond which the positive autoregulatory loop of *comK* would be activated, resulting in competence [the threshold in the *rok* strain changes because of increased gene expression (see SOM)]. The rightmost column shows CFP fluorescence images from the three strains, taken at T_2 and overlaid on DIC images. All three strains expressed the *comK-cfp** fusion, thus fluorescing when competent. The lowest panel in the column shows a microscopic field for the low-noise strain selected to show one competent cell, although the frequency of such cells was less than 1%. Scale bars, 4 μ m.

competent state, because large fluctuations triggering activation of the positive feedback loop would become less likely (29).

To test this prediction, we used the *rok* strain, which exhibits a twofold increase in *comK* mRNA transcription over the WT strain at T_0 (Fig. 4, panels in leftmost column) (25, 30), thus decreasing the noise in ComK protein levels. To adjust the mean ComK protein level in the *rok* strain to approximate that of the WT strain, we changed the ATG initiation codon of *comK* to GTG, thereby reducing its translational efficiency (9). We verified that the mean ComK levels in the low-noise and WT strains were similar by quantifying the amount of basal ComK–cyan fluorescent protein (CFP) fluorescence in bulk culture at $T_{0.5}$ and T_0 . Despite the slightly higher mean fluorescence in the low-noise strain, the number of its competent cells at T_2 was dramatically lower than that in the WT strain, with fewer than 1% of cells being competent as compared with 15% in the WT strain (Fig. 4, panels in rightmost column). These experiments show that intrinsic noise in *comK* expression is responsible for the transitions to competence and that reducing noise can substantially alter the rate at which those transitions occur.

This result suggests that the noise characteristics of particular genes may be subject to evolutionary pressures. Indeed, the fact that the *comK* gene is weakly transcribed (12) while having a “strong”

Shine-Dalgarno sequence (GGAGG–7 bp–ATG) is suggestive. For a desired final percentage of competent cells, there must be a set fraction of cells with the level of ComK above a particular threshold, achievable either by having a basal ComK distribution with a low mean and a large variance or by having a higher mean with a lower variance. Because of the metabolic cost of maintaining a larger mean number of proteins, it is plausible that cells would opt for the former option rather than the latter, as appears to be the case for *comK*.

The temporal regulation of *comK* transcription during stationary phase defines when transitions to the competent state may occur (the window of opportunity), and intrinsic noise in *comK* expression defines the rate at which cells become competent. Our results imply that noise properties are subject to evolutionary forces and suggest how cells might alter those rates to increase fitness. Because noise has been implicated in a variety of cellular behaviors, such knowledge can help both in the understanding of natural regulatory networks (7, 31) and in the synthesis of artificial networks (4, 32).

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33. We acknowledge the assistance of S. Tyagi, S. Marras, and D. Gold for the gift of the repeat sequence plasmid, for the use of their spectrophotometer and microscope, and for the preparation of fluorescent probes. We also acknowledge D. Rudner for the gift of the codon-optimized CFP and C. S. Peskin for valuable discussions. We also thank S. Tyagi, A. van Oudenaarden, J. Gore, J. Tsang, M. B. Elowitz, G. M. Suel, and P. Mehta for comments on the manuscript, as well as an anonymous reviewer for insightful comments. This work was supported by NIH grant GM 57720. A.R. was supported by NIH grant GM 070357 and by NSF postdoctoral fellowship DMS-0603392.

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5 February 2007; accepted 6 June 2007

Published online 14 June 2007;

10.1126/science.1140818

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