Transcriptome Divergence and the Loss of Plasticity in *Bacillus subtilis* after 6,000 Generations of Evolution under Relaxed Selection for Sporulation††

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Received 4 September 2008/Accepted 16 October 2008

We used microarrays to identify the causes of sporulation deficiencies in *Bacillus subtilis* after 6,000 generations of evolution. We found that sporulation loss did not result from large-scale deletions; therefore, it must have resulted from smaller indels and/or substitutions. Transcription patterns of one strain versus its ancestor showed that sporulation was not initiated and suggested that sporulation loss may be part of an overall decline in plasticity.

Spore formation by bacteria belonging to the *Firmicutes* is an ecologically beneficial yet energetically costly developmental process; dormant spores can better withstand harsh environmental insults (e.g., heat, desiccation, radiation, and oxidative and chemical attacks [13, 14, 16, 17]) than vegetative cells, but sporulation is costly in that it requires the coordinated transcription of hundreds of genes throughout several hours (reviewed in references 4 and 22 and references therein). Spore formation is a remarkable example of phenotypic plasticity, where cells are able to tailor their gene expression to specific environments or physiological states. In environments where benefits associated with spores are absent and no longer balance the cost of constructing spores, it would be predicted that sporulation ability should be lost over time (for extensive discussion of the theoretical details, see references 7 and 8). In order to test this prediction, we performed a laboratory evolution experiment in which two sets of five *Bacillus subtilis* populations were propagated for 6,000 generations in either the presence (populations 628A through 628E) or the absence (populations 624A through 624E) of strong selection for sporulation (7, 8). We observed that indeed, when *B. subtilis* was evolved without selection for sporulation, the ability to sporulate was either severely reduced (populations 624A and 624C) or completely lost (populations 624B, 624D, and 624E) in the nonsporulating experimental populations (8) (Table 1).

It was of interest to know whether sporulation ability was lost because selection favors sporulation mutants in a constant, nutrient-rich environment or whether sporulation had simply become neutral with respect to fitness and mutations in sporulation genes had accumulated over time. These two possibilities were addressed using simulations, which suggested that only one population, 624E, lost sporulation because selection favored its loss (8). Identifying the nature of the selectively advantageous mutation(s) underlying sporulation loss in 624E, and comparing it to the mutation(s) underlying sporulation loss in the other populations, is of interest because such a comparison can facilitate our understanding of why sporulation is adaptive in some natural environments but not in others.

Two types of mutations were examined: large-scale deletions and small-scale changes. Because sporulation is no longer needed in populations 624A to 624E, genome loss via large-scale deletion might occur, in a manner similar to that observed in bacterial endosymbionts evolving in the relatively constant environment of the insect gut (11, 12). Alternatively, small-scale changes (e.g., small insertions/deletions [indels] or single-nucleotide substitutions) in critical sporulation genes could lead to a blockage of sporulation. A long-standing and extensive literature exists documenting the pleiotropic effects of simple mutations in *spo* genes (reviewed in references 4, 21, and 22 and references therein). Furthermore, we have previously shown that the rate of spontaneous mutation to rifampin (rifampicin) resistance increased during evolution in populations 624A to -E by as much as 2 orders of magnitude (7), and many of these mutations were determined to be single-nucleotide changes in *rpoB* (15; H. Maughan and W. L. Nicholson, unpublished data).

To address these two possibilities, we used DNA-DNA microarrays to characterize large-scale changes in genome structure and RNA-DNA (transcription) microarrays to assess small-scale changes leading to alterations in gene expression patterns that have occurred during the loss of sporulation proficiency.

**Large-scale deletions.** To determine whether large-scale deletions resulted in the loss of sporulation genes, we prepared chromosomal DNA from single colonies isolated from each of populations 624A to 624E, propagated for 6,000 generations under relaxed selective pressure for sporulation, and from each of populations 628A to 628E, propagated for 6,000-generations under strong selective pressure for sporulation. Fluorescently labeled probes generated from the chromosomal DNA of evolved isolates were hybridized to 4,096-open reading frame (ORF) microarray slides prepared from *B. subtilis* strain 168 (Eurogentec, Belgium), along with differentially labeled probes prepared from DNA of the corresponding ancestral strains, WN624 and WN628, respectively. The intensities of each dye were quantified, and...
genes were considered to be missing if (i) the intensities differed significantly between the ancestor and the evolved strain (as determined by a false discovery rate-corrected P value from analysis of variance [ANOVA] analyses in R/MAANOVA [http://research.jax.org/faculty/churchill/software/Rmaanova/index.html]) and (ii) the difference between intensities was greater than twofold. No large-scale deletions were detected in isolates from any of the cultures (628A to 628E) that had evolved under strong selection for sporulation (data not shown). However, isolates recovered from two different nonsporulating populations, 624A and 624B, were found by DNA microarray analysis to harbor deletions removing DNA corresponding to the \textit{ppsD} and \textit{ppsC} genes, respectively (data not shown). These genes are members of the \textit{ppsABCDE} operon, which encodes a nonribosomal machine for the synthesis of plipastatin, a lipopeptide antibiotic (24). Because the \textit{ppsABCDE} operon exhibits repeated domains within the \textit{ppsA}, \textit{ppsC}, and \textit{ppsD} cistrons (23, 24) (Fig. 1), we reasoned that the observed deletions may have resulted from recombination between these homologous domains. To test this notion, the \textit{ppsCDE} region from ancestral strain WN624 and the evolved 624A strain was amplified by PCR using primers 5'-GCCGCAGCAACCTGAAATACAGGA-3' and 5'-TGTCGAGGATGATGTCCGACATTC-3', which amplify essentially the entire \textit{ppsC}, \textit{ppsD}, and \textit{ppsE} coding sequences (Fig. 1). The resulting amplicons were digested with EcoRI and their restriction patterns compared. Analysis of the EcoRI restriction patterns revealed that the evolved 624A strain had suffered a deletion of 9.7 kbp of DNA between the repeat sequences within the \textit{ppsC} and \textit{ppsD} cistrons (Fig. 1). This deletion removed the region of DNA corresponding to the Eurogentec \textit{ppsD} microarray probe, thus explaining the observed DNA microarray result. Although we did not further analyze the deletion removing \textit{ppsC} from the 624B evolved strain, we assume that \textit{ppsC} sequences were deleted by a similar recombination event occurring between the repeat sequences located within \textit{ppsA} and \textit{ppsC} (Fig. 1).

\textbf{Mutations affecting global transcription patterns.} Because the \textit{ppsABCDE} operon is not known to affect sporulation, and large-scale deletions were detected only in this operon, the DNA hybridization data suggest that sporulation loss during evolution was not caused by large-scale deletions in sporulation genes. Therefore, small-scale point mutations or indels are likely to have caused sporulation loss in the evolving populations 624A to 624E, and if the regulation of sporulation has been altered, transcriptional profiling may enable us to identify the genes containing such genetic changes. We therefore used transcription microarrays to measure global transcription patterns in the ancestor and evolved strains during the onset of stationary phase, the time at which sporulation is normally initiated under inducing conditions. RNA was isolated as soon as exponential growth started to slow down, which corresponded to optical densities at 600 nm of 0.65 to 0.82 in sporulation-repressing medium and 0.65 to 1.65 in sporulation-inducing medium. The evolved strain chosen for study was a single colony isolate from population 624E obtained after 6,000 generations of evolution under sporulation-repressing conditions, for which simulations suggested that selection had favored the loss of sporulation (8). The 624E isolate exhibited a sporulation frequency of \(10^{-6}\), compared to the ancestral level of \(58\%\) (8, 9). (Sporulation frequencies were measured by heat shocking the culture at 80°C for 10 min.) This low sporulation frequency for the 624E isolate was consistent with phase-contrast microscopy observations where spores were not seen, and the majority of cells were in long, nonmotile filaments.

\begin{table}
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\caption{Strains used in this study}
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\textbf{Ancestor and evolved strains} & \textbf{Experimental environment} & \textbf{Sporulation frequency} \\
WN624 (\textit{trpC2 spc::amyE}) & Sporulation repressing & 0.58 \\
WN624A & & \\
WN624B & & \\
WN624C & & 4.3 \times 10^{-4} \\
WN624D & & \leq 2 \times 10^{-6} \\
WN624E & & \leq 2 \times 10^{-6} \\
WN628 (\textit{trpC2 cat::amyE}) & Sporulation inducing & 0.58 \\
WN628A & & 0.65 \\
WN628B & & 0.92 \\
WN628C & & 1.0 \\
WN628D & & 0.83 \\
WN628E & & 1.0 \\
\hline
\end{tabular}
\end{table}

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure1.png}
\caption{Organization of the \textit{ppsABCDE} operon in \textit{B. subtilis}. Shaded arrows represent cistrons. Filled bars mark the locations of PCR primers. Diagonal and horizontal dashes indicate the locations of the probes present on the Eurogentec microarray and the locations of repeat sequences in \textit{ppsA}, \textit{ppsC}, and \textit{ppsD} (5, 6), respectively. The locations of EcoRI restriction sites are marked by the letter E. Asterisks indicate EcoRI restriction fragments missing in the 624A evolved strain but present in the WN624 ancestral strain.}
\end{figure}
The first set of experiments measured transcription in the ancestral strain WN624 and the 624E evolved strain when both were grown in the glucose-rich, sporulation-repressing experimental environment (Fig. 2A); this would show us how transcription had changed in the evolved strain after evolution for 6,000 generations under sporulation-repressing conditions. Global comparison of the transcription patterns showed a correlation coefficient ($\rho$) of 0.877 (95% confidence interval [95% CI], 0.869 to 0.884) (Fig. 2A). For purposes of comparison, analysis of a control transcription microarray comparing the two ancestral strains WN624 and WN628, both grown under sporulation-repressing conditions, resulted in a $\rho$ of 0.978 (95% CI, 0.977 to 0.98) (data not shown). In addition, transcription microarrays comparing ancestral strain WN628 with a strain (628B) that had evolved for 6,000 generations under strict selection for sporulation also demonstrated a higher correlation coefficient ($\rho$ = 0.939; 95% CI, 0.935 to 0.942) (Fig. 2B). The data suggested that evolution for 6,000 generations in the absence of selective pressure for sporulation resulted in a higher degree of transcriptome divergence than when selection for sporulation was applied. This transcriptome divergence was particularly dramatic when transcription microarrays compared the ancestral strain WN628 with an evolved strain from population 624E, when both strains were grown under sporulation-inducing conditions (Fig. 2C). This particular condition resulted in the lowest correlation coefficient of all, a $\rho$ of 0.647 (95% CI, 0.629 to 0.665) (Fig. 2C).

In addition to global transcriptome patterns, the transcription levels of individual genes were analyzed for ancestor and evolved strains. Transcription differences were considered to be significant if the false discovery rate-corrected $P$ value from ANOVA analyses was less than 0.05 and if transcription levels differed by at least 1.5-fold. We also used quantitative PCR (qPCR) to verify the relative transcription levels of several genes; in all cases, results from the qPCR qualitatively agreed with the microarrays (data not shown). In the transcription microarray experiment in which ancestral strain WN628 was compared to a strain from population 628B, which had evolved for 6,000 generations with stringent selection for sporulation, and RNA was isolated from cultures cultivated in sporulation-inducing medium (Fig. 2B), only 34 out of 3,916 total genes analyzed (0.9% of the total) were significantly differentially expressed (see Table S1 in the supplemental material). Twenty-four and 10 genes were significantly up- or downregulated, respectively, and most of these were genes of unknown function. Interestingly, the gerAB gene was 2.1-fold upregulated in the evolved strain, which might be expected to give a selective advantage to the evolved strain in that it may have enhanced its germination response during daily transfer of spores into fresh medium. In support of this notion, populations that had evolved under strict selection for sporulation indeed exhibited an average spor germination/lag time of 102.5 min, compared to the 182.5 min for the ancestral strain WN628 (7). In the transcription microarray experiment comparing transcription patterns between the ancestor WN624 and the evolved 624E isolate, both grown under sporulation-repressing conditions (Fig. 2A), it was observed that 5% of the total genes spotted onto the microarray (196 out of 3,996 genes analyzed) were transcribed at significantly different levels (see Table S1 in the supplemental material). Under sporulation-repressing conditions, significant upregulation of 107 genes occurred in the 624E evolved strain, most notably genes encoding ribosomal proteins, transport/bind proteins, and metabolic proteins. Uprregulation of these functions is presumably related to the continued growth of the 624E strain in stationary phase (data not shown). In the same experiment, significant downregulation was observed for 87 genes, most notably those encoding chemotaxis, motility, and flagellar proteins (see Table S1 in the supplemental material). The decreased transcription of chemotaxis/motility genes has also been observed in Escherichia coli populations that had evolved in the laboratory for 20,000 generations (2) and is presumably associated with the lack of need for chemotaxis or motility in shaking culture, coupled with the large biosynthetic and energetic costs of producing...
and operating the flagellar apparatus (10); loss of these functions presumably results in a competitively advantageous growth rate.

When we compared the transcription of individual genes in the 624E-derived evolved strain and ancestral strain WN624, both grown under sporulation-inducing conditions (Fig. 2C), we observed that 1,083 genes out of a total of 4,025 genes analyzed (27% of the total) were transcribed at significantly different levels (Fig. 2C; see also Table S1 in the supplemental material). Many of the genes that were upregulated in the evolved strain encoded ribosomal proteins, transport/binding proteins, proteins related to the metabolism of amino acids, and proteins related to cell wall functions, again consistent with the continued growth of the evolved strain in stationary phase. In addition to changes in these functional categories, 34 sporulation-associated genes were differentially transcribed in the ancestor and the evolved strain; 5 of these were upregulated and 29 were downregulated in the evolved strain (Table 2). Notable among the upregulated genes was abrB, a key protein controlling gene expression during the transition from exponential to stationary phase (20). The upregulation of abrB likely explains the concomitant downregulation of numerous genes, such as those involved in competence and the synthesis of extracellular antibiotics and proteases (see Table S1 in the supplemental material). Interestingly, most of the downregulated sporulation-associated genes encode products known to be involved in the initiation of sporulation (4, 5, 18, 22), including one dipeptide permease (dppC), two phosphorelay kinases (kinA and kinE), six Rap phosphatase regulators (phrA, phrB, phrC, phrE, phrF, and phrG), and three protease genes (spoVS, spoOF, and spo0M) (Table 2). Collectively, these data indicate that the evolved 624E strain is unable to sporulate due to the misregulation of genes involved in the very earliest stages of the vegetative-to-stationary-phase transition state and the initiation of sporulation.
environments, is thought to be adaptive (recently reviewed in references 1). This result indicated that fewer genes in the evolved strain were differentially regulated when the strain was faced with two very different environments. In contrast, comparison of the transcription patterns of the ancestral strain WN624 (A) and the population 624E-derived strain (B). Each data point indicates the normalized expression of one gene. The intensities were normalized and log transformed; therefore, each axis has a log scale. Correlation coefficients (ρ) are given. See the text for details.

The expression microarray results also suggested that the transcription profile of the 624E-derived evolved strain in sporulation-inducing medium was quite similar to its transcription profile in sporulation-repressing medium. Because we had transcription profiles of the WN624 ancestor and the 624E-derived evolved strain grown under both sporulation-repressing and sporulation-inducing conditions, we were able to analyze how the genes transcribed by each strain differed between the two environments. Comparison of the transcription patterns of the ancestral strain WN624 grown under sporulation-inducing versus sporulation-repressing conditions resulted in a ρ of 0.7 (95% CI, 0.683 to 0.715) (Fig. 3A), indicating a substantial level of transcriptome adjustment in the ancestral strain when it was faced with the two differing environments. In contrast, comparison of the transcription patterns of the evolved strain derived from population 624E, also grown under sporulation-inducing versus sporulation-repressing conditions, resulted in a higher ρ of 0.866 (95% CI, 0.859 to 0.874) (Fig. 3B). This result indicated that fewer genes in the evolved strain were differentially regulated when the strain was faced with two very different environments.

The similarity of the transcription patterns from the evolved strain when grown in sporulation-repressing versus sporulation-inducing environments shows that the ability of this strain to respond to different environments has declined since its divergence from the ancestor. Phenotypic plasticity, the ability of one organism to express different phenotypes in different environments, is thought to be adaptive (recently reviewed in reference 1) but is also predicted to be lost in constant environments (6). We consider the inability of the 624E-derived evolved strain to tailor its transcription to differing environments to be strong evidence that this strain has lost phenotypic plasticity. Furthermore, the loss of sporulation initiation may be part of a larger decrease in the ability of this strain to respond to different environments.

The 624E population from which the evolved strain was isolated showed a clear fitness advantage over the ancestor when they were competed in sporulation-repressing medium (7). Furthermore, evidence from simulations suggested that the loss of sporulation in this population was selectively advantageous (8). Monitoring the environment and coordinating gene expression to match environmental change is beneficial for cells living in an ever-changing environment, but this monitoring may become costly when cells find themselves in a constant environment (1, 3). Given this evidence, an important question remains: was the loss of phenotypic plasticity also selectively advantageous?

Microarray data accession number. Complete microarray data and additional methodological details have been deposited in Gene Expression Omnibus (http://www.ncbi.nlm.nih.gov/geo/) under accession number GSE4469.

We thank Ryan Sprissler of the Genomic Analysis and Technology Core facility at the University of Arizona for nucleic acid labeling and microarray hybridization and scanning; Helen Dunbar for assistance with qPCR; the laboratory of Howard Ochman and Nancy Moran for use of their LightCycler and for discussions concerning this work; and Carlos Machado, Leland S. Pierson III, Michael Nachman, and Nancy Moran for reading an earlier version of this article. We also thank two anonymous reviewers for their suggestions to improve this paper. This work was funded by two grants from Sigma Xi and an NSF Doctoral Dissertation Improvement Grant (DEB-0412650) to H.M. and by grants from the USDA (FLAMCS-04602) and NASA (NNX08AO15G) to W.L.N.

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