

## Global Transcriptional Response of *Bacillus subtilis* to Heat Shock

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**In response to heat stress, *Bacillus subtilis* activates the transcription of well over 100 different genes. Many of these genes are members of a general stress response regulon controlled by the secondary sigma factor,  $\sigma^B$ , while others are under control of the HrcA or CtsR heat shock regulators. We have used DNA microarrays to monitor the global transcriptional response to heat shock. We find strong induction of known  $\sigma^B$ -dependent genes with a characteristic rapid induction followed by a return to near prestimulus levels. The HrcA and CtsR regulons are also induced, but with somewhat slower kinetics. Analysis of DNA sequences proximal to newly identified heat-induced genes leads us to propose ~70 additional members of the  $\sigma^B$  regulon. We have also identified numerous heat-induced genes that are not members of known heat shock regulons. Notably, we observe very strong induction of arginine biosynthesis and transport operons. Induction of several genes was confirmed by quantitative reverse transcriptase PCR. In addition, the transcriptional responses measured by microarray hybridization compare favorably with the numerous previous studies of heat shock in this organism. Since many different conditions elicit both specific and general stress responses, knowledge of the heat-induced general stress response reported here will be helpful for interpreting future microarray studies of other stress responses.**

DNA microarray technology provides a powerful tool for the analysis of global transcriptional responses elicited by various physical and chemical stresses. One challenge in this sort of analysis is to distinguish stress-specific responses from more general stress responses. For example, in *Bacillus subtilis*, many different stresses (including heat shock, osmotic stress, and energy stress) activate the large general stress response controlled by the  $\sigma^B$  transcription factor (20, 44, 45). While others have used two-dimensional protein gels to classify cellular stress responses (55, 58), DNA-based methods have several advantages: they can be rapidly adapted to new organisms, they provide greater coverage of the genome, and data processing is comparatively easy to automate. Ultimately, it may be possible to integrate both technologies, at least for well-studied model organisms (19, 41, 56).

We have initiated a series of studies to characterize the global transcriptional responses of *B. subtilis*, a model gram-positive microorganism. Here, we document the heat-induced general stress response. Heat shock was chosen for this initial study since it is arguably the best-studied stress response in this organism and includes activation of the large general stress response under the control of  $\sigma^B$  (20, 44). In addition, a subset of antibiotics that inhibit translation have been reported to induce heat shock genes in other organisms (57). It is anticipated that knowledge of transcriptional responses to antimicrobial compounds will be useful for both antibacterial discovery and characterization (47).

Analysis of the transcriptional profile of *B. subtilis* after heat

shock clearly revealed the known heat shock regulons, including the large  $\sigma^B$ -dependent general stress regulon (21, 44), together with several operons not previously anticipated to be heat inducible. Prominent among these are operons involved in arginine biosynthesis and transport and many candidate new members of the  $\sigma^B$  regulon.

### MATERIALS AND METHODS

**Strains and growth conditions.** *B. subtilis* 168 strain MO945 was obtained from Niels Frandsen (GlaxoWellcome, Verona, Italy). It was grown in Bacto Mueller Hinton Broth (Difco, Detroit, Mich.). All experiments used baffled shake flasks. A 5-ml volume of medium in a 50-ml flask was inoculated and grown overnight at 37°C on a rotary platform (250 rpm). This culture was used to inoculate 50 ml of prewarmed medium (37°C) in a 500-ml flask to an optical density at 600 nm ( $OD_{600}$ ) of 0.05. The flask was shaken on a rotary platform (250 rpm) until an  $OD_{600}$  of 1.0 was attained. Samples (zero time) were taken from the 50-ml culture, and a 20-ml aliquot was transferred to a prewarmed 250-ml flask at 48°C which was incubated in a reciprocal-shaking water bath incubator at 48°C. A parallel identical experiment was performed with a prewarmed 250-ml flask at 37°C and incubation in a reciprocal-shaking water bath at 37°C. Samples were removed from these flasks for RNA extraction.

**Sampling and RNA isolation.** Samples of the culture were rapidly removed into 2-ml tubes and centrifuged at  $14,000 \times g$  for 10 s, and the culture supernatant was rapidly removed. The tubes containing the cell pellet were placed in liquid nitrogen. The entire procedure from the start of the centrifugation to the obtaining of the frozen pellet took approximately 40 s. Total RNA was extracted from *B. subtilis* by disruption in phenol/guanidine isothiocyanate (TRIzol; Life Technologies, Rockville, Md.). Briefly, TRIzol and zirconium silica beads were added to each 2-ml tube containing frozen cell pellets. The tubes were shaken on a Mini-beadbeater-8 (BioSpec Products, Bartlesville, Okla.) for four 1-min cycles. Nucleic acid was precipitated, and residual DNA was removed with 4 U of RNase-free DNase I (Ambion, Austin, Tex.). After extraction with phenol-chloroform, precipitation and resuspension the RNA was quantitated with RiboGreen (Molecular Probes, Eugene, Oreg.).

**Generation of ORF DNA and production of microarrays.** Oligonucleotide primers for all 4,100 open reading frames (ORFs) in the *B. subtilis* genome were purchased from Eurogentec (Seraing, Belgium). Full-length ORFs were made by PCR, with the following cycling conditions: 1 min of denaturing at 95°C, 45 s of annealing at 55°C, and 3.5 min of elongation at 72°C. All PCR products were

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purified with the QIAquick 96-well purification kit from Qiagen (Valencia, Calif.). The quality of the amplified sequences was checked by electrophoresis on a 1.5% agarose gel. The gels were digitally imaged, and the band sizes were entered into a database where the expected size was compared to the observed size. Additional data describing faint and multiple bands were also collected. In 481 cases, the PCRs failed to yield satisfactory products (no product, wrong size, additional bands, or faint bands) and oligonucleotide primers for selected genes were redesigned and obtained from Operon (Alameda, Calif.) or MWG Biotech (High Point, N.C.). Finally, over 90% (3,703 ORFs) of the *B. subtilis* genome was correctly amplified. Slide preparation and printing followed the procedures described by Wilson et al. (64). Briefly, amplicons were suspended in 6× SSC–15% DMSO and spotted onto poly-L-lysine-coated slides by using Telechem (Sunnyvale, Calif.) SMP5 spotting pins and an SPH16 printhead fitted to a Genemachines Omnigrad arrayer (1× SSC is 0.15 M NaCl plus 0.015 M sodium citrate). Spot spacing was 230 μm. Slides were processed as previously described (64) and stored under N<sub>2</sub>.

**cDNA labeling and slide hybridization.** Each fluorescently labeled cDNA probe was prepared from 6 μg of DNase I-treated total RNA by random hexamer [pd(N)<sub>6</sub>; Amersham Pharmacia Biotech, Piscataway, N.J.]-primed polymerization using reverse transcriptase (Superscript II RT; Life Technologies, Gaithersburg, Md.). Concentrations of nucleotides in the labeling reaction mixture were as follows: 0.5 mM dGTP, 0.5 mM dATP, 0.5 mM dTTP, and 0.05 mM dCTP. The final concentration of Cy3-dCTP or Cy5-dCTP (Fluorolink Cy dye-labeled dCTP; Amersham Pharmacia Biotech) was 0.04 mM. The final concentration of random hexamer was 0.06 mM. Unincorporated dye-labeled dCTP was removed by washing the probe in a microconcentrator (Microcon YM-30; Millipore, Bedford, Mass.).

Microarray slides were incubated for 30 min at 42°C with prehybridization solution (1% bovine serum albumin, 0.5% L-glutamate, 4× SSC), washed three times in double-distilled H<sub>2</sub>O, and dried by centrifugation at 50 × g for 90 s. For each hybridization, cDNA probe made from RNA from untreated cells (time zero sample) was mixed with probe made from RNA from heat-shocked cells. Each microarray received ~22 μl of hybridization solution (4.2× SSC, 42% formamide, 0.17% SDS, 63 μg of salmon sperm DNA/μl) containing the two probes. The solution was applied by capillary action under a coverslip (LifterSlip; Erie Scientific Company, Portsmouth, N.H.) placed over the microarray. The whole assembly was sealed in a hybridization chamber (CMT Hybridization Chamber; Corning Incorporated, Corning, N.Y.) and submerged for 16 h in a 42°C water bath. Microarray slides were washed for 1 min in 1× SSC–0.05% SDS, 30 s in 0.06× SSC, and again for 1 min in 0.06× SSC. Slides were dried by centrifugation at 50 × g for 90 s and were immediately scanned and analyzed with a confocal laser scanner/software package (Axon GenePix 4000A/GenePix Pro 3.0; Axon Instruments, Inc., Foster City, Calif.).

**Data analysis.** For analysis, any gene feature that had <80% of pixels >2 standard deviations above the local background in both channels was rejected. Ratios for levels of RNA (heat-shocked divided by time zero sample) were calculated using a ratio of medians method. Any gene feature wherein one channel was within one standard deviation of the local background was flagged as giving a potentially inaccurate ratio (indicated in the tables by values in italics; also indicated in supplemental material S2 [http://www.micro.cornell.edu/faculty.JHelmann.html]). Data normalization was based on the premise that the ratio of measured expression averaged over the entire set of sorted genes for which data was obtained is approximately equal to 1. We used a normalization method based on the geometric mean (average of the logarithmic measures of the ratios) rather than the arithmetic mean of ratios, as the geometric mean accounts for down- as well as up-regulation. Specifically, each ratio output from the scanner was multiplied by a factor of 2<sup>–[average of log<sub>2</sub>(ratios)]</sup>. A further explanation and proof of this normalization method are given in the supplemental material (S1 [http://www.micro.cornell.edu/faculty.JHelmann.html]).

To check for reproducibility in the cDNA preparation and hybridization steps, we tested the competitive hybridization of two cDNA samples both prepared from a culture grown at 37°C. Of the 2,033 gene signals detected, the overall range of ratios was quite small (1.47- to 0.62-fold range; 96% of the ratios were between 0.75 and 1.25). All experimental data were collected by the competitive hybridization of three independent cDNA preparations from each time point against the non-heat-shocked control sample (referred to as experiments 1 to 3). A comparison of 90 genes previously assigned to the heat shock stimulus showed that for genes where all three experiments yielded valid ratios, 46% of triplicate ratios yielded a coefficient of variation (CV) of <20% and 97% yielded a CV of <40%. The entire data set for all three experiments can be found in the supplemental material (Table S2 [http://www.micro.cornell.edu/faculty.JHelmann.html]).

Initial analysis focused on three overlapping sets of genes. For the first set, the

induction profiles for all reported members of the heat shock stimulus were compiled from all three experiments. The resulting data from one hybridization experiment (no. 3) are presented in this study except where noted. This data set was chosen since this set of slides yielded a more complete data set than the other two replicates, with more than 3,000 genes detected with signals above background (compared to ~2,600 genes for experiments 1 and 2). However, the overall transcriptional response in each set of hybridizations was very similar (e.g., see Fig. 4). For the second set, the 50 most highly induced genes from each of the nine data sets (three experiments with three time points each) were tabulated. The resulting list of 450 gene signals was found to result from 143 different genes (set 2), most of which appeared, as expected, in multiple experiments. For the third set, all 405 genes induced greater than twofold at the 3-min time point in experiment 3 were analyzed further. Since the σ<sup>B</sup> regulon was so large and was induced only transiently, many potential new members of this regulon were included in set 3 but were not found in the other two sets of genes.

The lists of genes in sets 2 and 3 were analyzed to remove artifactual signals as judged by nonreproducibility in the induction. For example, in several cases, genes were initially included in set 2 (among the top 50 induced genes at one time point), but further analysis indicated that this was due to a single point which was flagged as possibly inaccurate as noted above. Moreover, there was often reliable data in the replicate experiments that clearly showed little or no induction of this same gene. By this criterion, 19 genes were removed from set 2. Of the remaining 124 genes which were reproducibly induced by heat shock, about half were members of known heat shock regulons (set 1). The remaining genes were visually analyzed to determine likely operon organization and inspected for the presence of candidate σ<sup>B</sup>-like promoter elements. A similar treatment was used for those genes in set 3. Those having a plausible match to the σ<sup>B</sup> consensus are listed in Table 4, and other heat-inducible genes of unknown regulation are included in the supplementary material (Table S3 [http://www.micro.cornell.edu/faculty.JHelmann.html]). The heat shock response of genes likely to be cotranscribed with strongly induced genes was also evaluated, and in many cases they demonstrated very similar folds induction and kinetics. This provides additional support for the observed regulation. Likely operon organization, DNA sequences, and current functional assignments were all obtained from the SubtiList database (37).

**Quantitative RT-PCR.** Taqman quantitative reverse transcription (RT)-PCR primers and probes were designed using Primer Express software (Applied Biosystems, Foster City, Calif.) and were synthesized by Applied Biosystems. 6FAM reporter dye and TAMRA quencher were affixed on the 5' and 3' ends of the probe, respectively. Primer and probe sequences were as follows: *sigB* sense primer, GATGAAGTCGATCGGCTCATAAG, antisense primer, CCCGCAC AAGCGTTTCC, and probe, TTACCAACAAAGCAAGATGAACAAGC GC; *argB* sense primer, TTGCTGAGCTTGCCAAACAC, antisense primer, CAAAAGACCGCCATCCTTACC, and probe, AATGCCCGCGCTCGCA GT; *dnaK* sense primer, TGAGCTTGGCGACGGTGTGA, antisense primer, GATGATCGATGATAACTTGGTCAA, and probe, TTCGTTCAACTGCC GCGACAA; *cisR* sense primer, CAAGGTAATTTAGAAAGAGAAGC AA, antisense primer, TTCTCGCTCTTAATTCATCACGTT, and probe, TAA TGGACGCTCAGTTTTACACATTGACTTACC. Reactions were performed using 50 ng of the DNase-treated total RNA, a 300 nM concentration of each Taqman primer, and 150 nM Taqman probe in a 50-μl volume. Controls lacking reverse transcriptase or template were used. Reactions were run on an ABI 7700 instrument (Applied Biosystems) using the following cycling parameters: reverse transcription at 48°C for 30 min, reverse transcriptase inactivation at 95°C for 10 min, 40 cycles of denaturation at 94°C for 15 s, and extension at 60°C for 1 min. Changes in expression were calculated from the displacement of the amplification curve of the heat-shocked sample from the time zero sample.

**Determination of transcriptional orientation.** Transcriptional orientation for the elucidation of proximity effects in the transcription of *yfkT* was determined by Taqman RT-PCR using oppositely oriented primers. RT was carried out for 30 min at 48°C from 50 ng of DNase-treated total RNA with either 300 nM *yfkT* sense primer (TGACCAAGATGGCGCAGAT) or 300 nM antisense primer (CCAGCGTAAATGGAAGGAACA). The Taqman probe (6FAM-TTCCTAT TTCCATTCGGCATCCTGGTC-TAMRA; 150 nM) and AmpliTaq Gold DNA polymerase (Applied Biosystems) were included in the reaction mixture. RNA was digested by the addition of an RNase A (5 μg; Roche)/RNase T1 (10 U; Ambion) mixture and incubated at 37°C for 1 h. Reverse transcriptase was inactivated, and *Taq* was activated at 95°C for 10 min. A 300 nM concentration of the opposing primer was then added, and the reaction was run on an ABI 7700 instrument with 40 cycles of denaturation at 94°C for 15 s and extension at 60°C for 1 min. Changes of expression in either orientation were calculated as described above.

TABLE 1. Induction of class I (HrcA-dependent) heat shock genes

Gene <sup>b</sup>	Microarray results (fold induction at time indicated)			mRNA levels from slot blot analysis <sup>a</sup>		
	3 min	10 min	20 min	5 min	10 min	15 min
	<i>hrcA</i>	6.0	8.6	2.6	10.5	11.0
<i>grpE</i>	3.1	5.8	2.2	7.0	7.0	2.0
<i>dnaK<sup>c</sup></i>	2.6 (2.5)	5.5 (3.1)	2.3 (2.3)	6.0	6.5	3.0
<i>dnaJ</i>	2.0	2.0	0.88	3.5	3.0	1.5
<i>yqeT</i>	1.7	1.5	0.67	3.5	3.0	1.5
<i>yqeU</i>	1.1	0.85	0.55	3.0	3.0	1.0
<i>yqeV</i>	1.4	1.1	0.79	2.5	2.5	1.0
<i>groEL</i>	3.1	6.9	4.1			
<i>groES</i>			2.7			

<sup>a</sup> Values were estimated to the nearest 0.5-fold, from a slot blot histogram (Fig. 2) (25).

<sup>b</sup> Downstream genes in operons are indented.

<sup>c</sup> Quantitative RT-PCR results are shown in parentheses.

## RESULTS AND DISCUSSION

To develop a platform for monitoring global transcriptional responses in *B. subtilis*, we have amplified, using PCR, ~90% of the ~4,100 annotated ORFs and arrayed the resulting products on glass slides. In this report, we characterize the transcriptional response elicited by shifting a growing culture from 37 to 48°C, and we compare the resulting data with those obtained in the numerous previous studies of the heat shock stimulon in this organism (reviewed in references 19, 21, and 44).

**Experimental design and array validation.** To measure gene expression under different conditions, total RNA was isolated and labeled by RT in the presence of either Cy3-dCTP or Cy5-dCTP in reactions primed with random hexamers. The resulting cDNAs were hybridized to glass slide microarrays as described in Materials and Methods. The relative hybridization of the two cDNA populations was ascertained by the relative fluorescence of the two fluorophores. The resulting data are expressed as the fold induction in the accompanying tables. While it is possible, by using appropriate normalizations, to convert fluorescence intensities to absolute transcript levels (63), we have not attempted such an analysis with these data.

Altogether, three sets of hybridization experiments were performed to measure heat shock-induced changes at 3, 10, and 20 min after shifting to 48°C (nine data sets). To control for possible variability in nucleoside incorporation, each experiment was performed at least once with the Cy3- and Cy5-labeled nucleosides reversed. As a practical matter, fold induction or repression could be confidently measured over a nearly 10,000-fold range (100-fold induction to 100-fold repression). However, for some genes, the fluorescence signal in one channel was near background and the fold induction or repression could not be confidently estimated.

In a typical experiment, hybridization signals were obtained, at levels significantly above background, for ~70% of all genes under these growth conditions. This is comparable to results reported previously for *Escherichia coli* (4, 54, 63). When these signals are mapped onto the chromosome, several large clusters of apparently silent genes map to the integrated SPβ prophage, the *skin* element, and several other proposed prophages (data not shown). In a control experiment involving competitive hybridization of two cDNA samples both prepared

TABLE 2. Heat shock induction of selected  $\sigma^B$ -dependent genes<sup>a</sup>

Gene <sup>c</sup>	Fold induction at time (min) <sup>b</sup> :		
	3	10	20
<i>bmrU</i>	18	1.6	1.9
<i>bmr</i>	1.2	0.45	0.37
<i>bmrR</i>	2.0	1.6	1.2
<i>bofC</i>	5.7	0.86	1.3
<i>csbA</i>	4.2	1.1	1.4
<i>csbB</i>	7.3	0.56	1.2
<i>yfhO</i>	4.9	0.35	0.77
<i>csbC (yxcC)</i>	16	2.2	2.4
<i>csbD (ywmG)</i>	22	14	4.8
<i>csbX</i>	6.7	1.0	1.1
<i>ctc</i>	4.1	2.0	1.4
<i>dps</i>	8.1	2.5	2.5
<i>gsiB</i>	6.7	19	7.4
<i>gspA</i>	26	9.7	7.2
<i>katB</i>	25	9.0	4.4
<i>trxA</i>	1.6	2.6	2.3
<i>yacH</i>	17	27	3.6
<i>yacI</i>	18	30	4.5
<i>yacL</i>	5.0	4.3	1.4
<i>yedF</i>	11	8.8	1.9
<i>yedG</i>	22	7.9	3.8
<i>ydaP</i>	24	6.3	4.4
<i>yfkM</i>	10	7.5	3.6
<i>yflT<sup>d</sup></i>	30	34	10
<i>yhdF</i>	23	15	6.2
<i>yhdG</i>	0.81	10	0.60
<i>yhdN</i>	14	5.2	3.7
<i>yjbC</i>	4.1	2.7	1.5
<i>yjbD</i>	2.9	4.2	3.0
<i>ykzA</i>	21	17	6.2
<i>yocK</i>	12	1.6	2.4
<i>ysdB</i>	4.3	1.9	1.4
<i>ytkL</i>	4.5	3.5	2.8
<i>ytxG</i>	8.0	2.0	1.4
<i>ytxH</i>	7.9	2.2	1.7
<i>ytxJ</i>	5.7	1.9	1.4
<i>yvyD</i>	2.3	1.3	1.4
<i>yxkO</i>	5.9	6.8	— <sup>e</sup>

<sup>a</sup> Additional known  $\sigma^B$ -dependent genes are shown graphically in Fig. 1 and 2. Three additional  $\sigma^B$ -dependent genes (*katX*, *gtaB*, and *opuE*) were not included in the arrays used in this study.

<sup>b</sup> All data are from one of three experiments (no. 3). Qualitatively similar results were obtained in each of the other two experiments except where noted. Numbers in italics are those flagged as inaccurate due to low signal in one channel.

<sup>c</sup> Downstream genes in (putative) operons are indented.

<sup>d</sup> *yflT* was assigned to the  $\sigma^B$  regulon based on proteome studies (60).

<sup>e</sup> —, insufficient signal was obtained to estimate fold induction.

from a culture grown at 37°C, no signal (of >2,000) varied by more than twofold (range, 1.47- to 0.62-fold). Thus, changes in cDNA populations well beyond this range are likely to reflect real differences in the corresponding RNA populations. The analysis described used data from one set of hybridizations (experiment 3), but similar results were obtained from the other two experiments (see Materials and Methods and supplemental material [http://www.micro.cornell.edu/faculty.JHelmann.html]), and reference is made to these results where needed. In order to independently confirm the veracity of the microarray results, the expression of four genes was also quantitated by real-time RT-PCR (22).

**Overview of the heat shock stimulon.** To obtain an overview of the heat shock stimulon, we focused our analysis on three

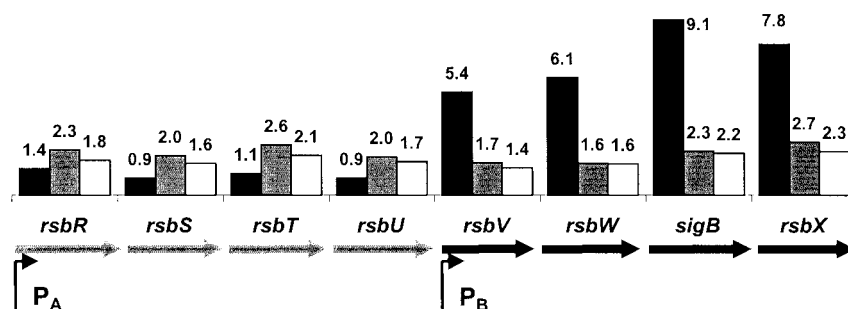


FIG. 1. Heat induction of *sigB* operon genes (*rsbR-S-T-U-V-W-sigB-rsbX*). The *sigB* operon is illustrated schematically (genes are not to scale), and the fold induction by heat shock at 3 min (black), 10 min (grey), and 20 min (white) is superimposed on the operon structure. The operon is transcribed from an upstream  $\sigma^A$ -dependent promoter (65) and from an internal, heat-inducible  $\sigma^B$ -dependent promoter (6, 27). Measurements of *sigB* induction by RT-PCR yielded values of 11.8-fold (3 min), 3.0-fold (10 min), and 3.3-fold (20 min).

overlapping sets of genes. Set 1 included all previously described heat-inducible genes, set 2 included the 50 most strongly induced genes in all nine data sets (three replicate hybridization experiments with three time points each: 450 gene signals), and set 3 included all those genes induced at least twofold at the 3-min time point in experiment 3 (which yielded the most complete data set). Since the  $\sigma^B$  regulon is very large and is induced transiently, many new candidate members of the  $\sigma^B$  regulon appeared in set 3 but not in set 2. Our analysis identified many known members of the  $\sigma^B$  and other heat shock regulons. However, we have also identified new heat shock genes, including many with candidate  $\sigma^B$ -dependent promoters.

Consistent with existing nomenclature in *B. subtilis* (20, 21, 44), heat shock genes are assigned to several discrete classes: class I is the HrcA regulon, class II genes are  $\sigma^B$  dependent, and class III genes are regulated by CtsR (and may also be regulated by  $\sigma^B$ ). We suggest that those genes for which the regulatory pathway is not yet characterized be designated class U heat shock genes (unknown regulation) rather than class IV, since the latter nomenclature will likely lead to confusion as additional regulons are defined.

**The HrcA regulon (class I).** The HrcA protein is a transcriptional repressor of class I heat shock genes (50). This repressor binds to conserved *cis*-acting regulatory sequences known as CIRCE elements (67) and responds specifically to heat induction. In *B. subtilis*, HrcA is known to regulate the expression of two operons, the complex *hrcA* operon (24, 50) and the *groEL*-

*groES* operon (34, 49). Strong and reproducible signals were not obtained for the *groEL-groES* operon in this study, so we focus our analysis on the *hrcA* operon.

Transcription initiating in the *hrcA* promoter region leads to the synthesis of an 8-kb primary transcript spanning seven genes that is rapidly processed into a complex family of smaller transcripts (24, 25). Previously, mRNA levels for all seven genes were measured at 5, 10, 15, and 30 min after the shift from 37 to 48°C (25). Our data are in reasonable agreement with those obtained previously (Table 1). We find the strongest induction for the first three genes in the operon, *hrcA*, *grpE*, and *dnaK*, with weaker effects on the downstream genes. In fact, in our studies, the three promoter distal genes were induced little if at all, with a maximal fold induction of ~2-fold. This is consistent with the slot blot analysis, which demonstrated at most two- to fourfold induction for these genes (Table 1).

**The *sigB* regulon (class II heat shock genes).** Activation of  $\sigma^B$  in response to heat stress is well documented, and it is estimated that the  $\sigma^B$  regulon includes over 200 genes (19, 44, 45). Genes belonging to the  $\sigma^B$  regulon are prominently represented among the genes of the heat shock stimulon, particularly at the 3-min time point. Because the mRNA levels for many  $\sigma^B$  regulon genes return rapidly to pre-stimulus levels (see below), members of the  $\sigma^B$  regulon were not well represented among the most strongly induced genes at later time points.

For purposes of discussion, we can divide known and puta-

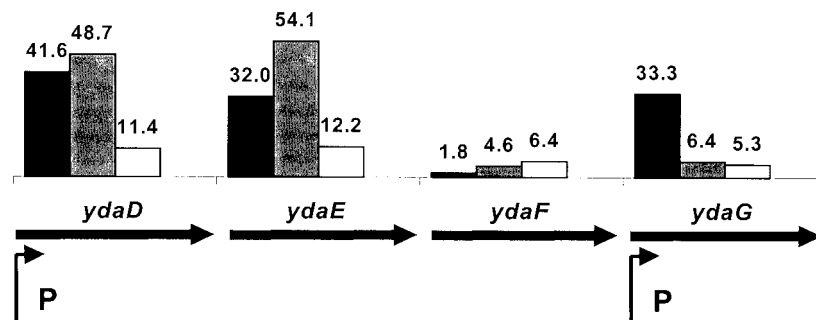


FIG. 2. Heat induction of the *ydaDEFG* gene cluster. The *ydaDEFG* cluster of genes has been previously shown to be expressed from two  $\sigma^B$ -dependent promoter elements as shown (42). Induction at 3, 10, and 20 min is shown as in Fig. 1.



TABLE 3. Fold induction of candidate  $\sigma^B$  regulon genes<sup>a</sup>

Gene <sup>b</sup>	3 min	10 min	20 min
<i>aldY</i>	17	11	18
<i>lctE</i> <sup>c</sup>	8.8	10	8.3
<i>ycnH</i> <sup>c</sup>	8.0	16	8.5
<i>ydaT</i>	26	17	4.9
<i>ydaS</i>	27	22	6.1
<i>ydhK</i>	14	1.9	1.5
<i>yfhK</i>	16	2.3	3.3
<i>yfhL</i> <sup>d</sup>	7.1	1.9	2.7
<i>yfhM</i>	7.7	1.9	2.4
<i>yflA</i>	21	3.1	3.5
<i>yjgB</i>	37	22	5.3
<i>yjgA</i>	3.6	2.8	1.7
<i>ykgA</i>	25	2.1	3.4
<i>ykgB</i>	2.6	1.7	1.5
<i>yoxC</i>	16	2.5	2.4
<i>yoxB</i>	12	1.2	1.5
<i>yoaA</i>	4.6	2.0	2.8
<i>yqhA</i>	12	0.82	1.8
<i>yqxL</i> <sup>e</sup>	17	2.3	3.8
<i>yvrE</i>	6.3	1.7	1.5
<i>ydbP</i>	1.4	1.4	1.4
<i>yoxA</i>	0.62	0.28	0.68
<i>ypuB</i>	1.9		1.2
<i>ypuC</i>	2.8	1.4	
<i>yqhQ</i>	1.3	1.1	0.99
<i>yqhP</i>	1.6	1.3	1.2
<i>yqiS</i>	0.66	0.73	0.78
<i>yrvD</i>	1.7	0.65	1.1

<sup>a</sup> Genes were previously identified by Petersohn et al. (43). No data were obtained for *yotK* and *yycD*.

<sup>b</sup> Downstream genes in (putative) operons are indented.

<sup>c</sup> Ethanol induction is not  $\sigma^B$  dependent (43).

<sup>d</sup> The *yfhLM* genes are also transcribed from a  $\sigma^W$ -dependent promoter (26).

<sup>e</sup> Data are from experiment 1 instead of experiment 3.

tive members of the  $\sigma^B$  regulon (class II heat shock genes) into three subcategories. Class IIA includes those genes for which a dependence on  $\sigma^B$  has been documented by direct start site mapping (e.g., by primer extension), genetic experiments, or both (see reference 44). Class IIB includes genes previously postulated to be members of the  $\sigma^B$  regulon, based on a promoter consensus search procedure (43). Class IIC includes additional heat shock genes identified in this study that are preceded by candidate  $\sigma^B$  promoters. Many of these same genes were independently assigned to the  $\sigma^B$  regulon based on transcriptional profiling experiments monitoring gene expression in response to ethanol stress and induction of *sigB* expression (45).

**Class IIA.** Many of the well characterized genes belonging to the  $\sigma^B$  regulon (44) are induced following heat shock (Table 2; Fig. 1 and 2). Comparison of their expression kinetics reveals a consistent pattern: in general, the  $\sigma^B$  regulon is very rapidly activated in response to temperature shift with relative RNA levels (fold induction) rapidly increasing by the 3-min time point and often, though not always, declining by the 10-min time point. The transient nature of induction in response to heat stress is consistent with previous analyses of  $\sigma^B$ -dependent transcripts, including *sigB* itself (6, 59), *ctc* (6), *gspA* (1), *katB* (13) and *trxA* (48).

The  $\sigma^B$  regulon includes *sigB* itself, which is transcribed as part of a complex operon containing eight genes (Fig. 1). As

expected, RNA levels corresponding to all four genes downstream of the internal  $\sigma^B$ -dependent promoter are rapidly elevated after heat shock (~5- to 9-fold), while RNA corresponding to the upstream genes is only slightly induced. Note that none of these genes are strongly induced, and none of them are represented among the top 50 induced genes, even at the 3-min time point. Like other members of the  $\sigma^B$  regulon, mRNA levels return to near the pre-stress level by 10 and 20 min following heat shock. Previous measurements of *sigB* mRNA levels following heat shock revealed a maximal induction of >20-fold at 5 min, followed by reduced levels at 10, 15, and 20 min following temperature stress (59).

For comparison with the array data, we independently determined the degree of induction for the *sigB* mRNA using a quantitative RT-PCR approach (22). These experiments revealed consistent kinetics of induction with a maximal induction (at 3 min) of 12-fold. Note that this is somewhat higher than the induction measured using the arrays (ninefold). In a control experiment, using cells shifted from 37°C to another flask at 37°C, we also see a much weaker (~2-fold) but still significant induction of *sigB* (data not shown). This may be a response to stresses associated with transfer of the cells. For example, removal from a well aerated flask using a glass pipette could allow a transient depletion of oxygen in the rapidly growing culture.

The *ydaDEFG* region of the chromosome has been previously analyzed (42) and found to have two  $\sigma^B$ -dependent promoters giving rise to a complex family of transcripts as determined by Northern blot analysis. Our data support the suggestion, from Northern analysis, that *ydaD* and *ydaE* are cotranscribed as both are strongly and coordinately induced (Fig. 2). In addition, we find very strong induction of *ydaG*, but *ydaF* is induced much more modestly, and with slower kinetics. This is consistent with the presence of a prominent heat-induced transcript corresponding to the *ydaG* gene, and argues that there may be limited transcriptional readthrough from *ydaDE* into *ydaF*.

Two genes considered to be members of the  $\sigma^B$  regulon that are not strongly induced by heat shock are *bmrR* and *bmr*, the two promoter distal genes in the *bmrU* operon (Table 2). However, *bmrU* is strongly and reproducibly induced. It had previously been suggested that *bmrR* and *bmr* may be cotranscribed with *bmrU*, but the published Northern blot analysis suggests that most transcription terminates after the *bmrU* gene (42). The lack of strong heat induction of the promoter distal genes, *bmrR* and *bmr*, is consistent with the idea that readthrough into these downstream genes does not greatly affect their expression.

**Class IIB.** Using a consensus search approach, Petersohn et al. (43) identified 31 additional  $\sigma^B$ -type promoters. In three cases (*yhdF*, *yacL*, *ysdB*), these promoters were confirmed by primer extension mapping, and these genes have therefore been added to class IIA (Table 2). An additional 25 sites were shown, using slot blot analysis, to be induced by ethanol. In all but four cases, this induction was apparent in the wild type but not in a *sigB* mutant strain (43). Three genes (*yabJ*, *yhaR*, and *yqhZ*) are not induced by ethanol (43), and we found that these genes did not respond to heat shock. Thus, these putative  $\sigma^B$  promoters may represent false positives generated by an imperfect search algorithm.

TABLE 4. Heat shock-induced genes associated with candidate  $\sigma^B$ -dependent promoters

Gene <sup>a</sup>	Sequence [GttTww-----gGgwAw (sp12-15)]	3 min	10 min	20 min	Function or comments <sup>b</sup>
<i>yaaI</i>	ttcaaa <b>gTTTTT</b> tcattgc-ctaaaa <b>gGCT</b> acatattaact	11	2.4	1.9	Isochorismatase homolog
<i>yaaH</i>		12	2.5	2.0	Similar to cortical fragment-lytic enzyme; <i>N</i> -acetylglucosaminidase (7)
<i>ybyB</i>	caacag <b>gTTT</b> agcaattt--ccaaaa <b>gGGAAT</b> gatacagga	4.6		5.9	84 aa; unknown function (no signal in 2 expt)
<i>ycbP</i>	aataag <b>gTTT</b> aactttt--acatttg <b>gGAAT</b> tatacataa	5.3	4.3	2.0	Membrane protein; divergent from <i>cwlJ</i>
<i>ydaB</i>	gaaatag <b>TTTT</b> caagttatctttt <b>gGTT</b> aaaaatgggtg	5.5	1.6		Predicted acid-CoA ligase ( <i>yfhL</i> paralog)
<i>ydaC</i>	aaaaat <b>gTTT</b> ccatggaa-cgctgag <b>gGAA</b> ttaactcaa	19			HC
<i>ydaJ</i>	ggctct <b>gTTT</b> cttaatg-ttcaaaa <b>gGGA</b> aaaaagcta	11	2.7	3.0	No described homolog
<i>ydaK</i>		4.8		0.81	HC; similar in one domain to <i>Myxococcus xanthus</i> ActA response regulator (17)
<i>ydaL</i>		5.3	1.1	1.2	No described homolog
<i>ydaM</i>		4.4	0.71	0.94	Similar to predicted glycosyl-transferases implicated in intercellular adhesion and biofilm formation in staphylococci (23)
<i>ydaN</i>		4.0	0.74		No described homolog
<i>ydbD</i>	tttttc <b>gTTT</b> atcttttcta--tcgat <b>gGAA</b> tataaaaaag	17	23	7.1	Identified as general stress protein (GS80) (2); 50% ID to Mn catalase from <i>Lactobacillus plantarum</i>
<i>ydgC</i>	atactc <b>gT</b> gagtaacatta---ctc <b>gTgG</b> gatatttttgg	2.5	2.5	1.4	Transcriptional regulator (TetR/AcrR family)
<i>yerD</i>	gctatt <b>gTTT</b> ggaaagtgt-tctact <b>gTgGAA</b> tggttacat	8.4	3.3	1.8	Similar to glutamate synthase (ferredoxin)
<i>yfhF</i>	aaac <b>gT</b> tttcttttatt--acaat <b>gagG</b> taaaagtatttt	11	1.9	2.4	No described homolog, <i>yfhFED</i> operon
<i>yfhD</i>		12	2.1		No signal for <i>yfhE</i> ; 36 aa
<i>yfkE</i>	tacaac <b>gTTT</b> ccaaaagcaggcaacct <b>gAAAA</b> agcctata	17	2.8	2.6	Similar to H <sup>+</sup> /Ca <sup>2+</sup> exchanger
<i>yfkD</i>		18	3.2	2.8	No described homolog
<i>yfjK</i>	atgaag <b>gT</b> ttctttttaga-gaaat <b>agG</b> gcaagaataggg	11	2.4	2.0	Similar to protein-tyrosine phosphatase
<i>yfjKH</i>		9.4	2.6	2.1	Similar to transporter (no signal for <i>yfjI</i> )
<i>yfjS<sup>c</sup></i>	aact <b>gT</b> taaggagttagaaggaaaa <b>gG</b> gatcggaaaaaca	13	1.8	2.0	50% ID to 2-oxoglutarate/malate translocator (61)
<i>ygxB</i>	ccaaat <b>gT</b> ataaataattt---cagcc <b>gG</b> gagatttcatat	16	4.4	1.5	HC
<i>yhaS</i>	cataa <b>gT</b> tttataatagtagaaaa <b>gG</b> gatactctgATGA	4.0	5.9	2.3	No described homolog
<i>yhaT</i>		4.3	6.8	3.0	YvrC paralog; HC
<i>yhaU</i>		3.0	4.9	2.4	Similar to Na <sup>+</sup> /H <sup>+</sup> antiporter
<i>yhcM</i>	tataac <b>gT</b> taatttgtct-aacgag <b>gG</b> gaaaaatgaata	16	10	4.0	No described homolog
<i>yheK</i>	ggaaa <b>gT</b> taATTgTgct-caaat <b>cgG</b> tagtagtgtgt	30	16	8.4	HC; renamed <i>nhaX</i> and proposed to be regulatory gene co-transcribed with <i>nhaC</i> encoding Na <sup>+</sup> /H <sup>+</sup> antiporter (62) (see text)
<i>yhxD</i>	aaacat <b>gT</b> tttctgctta-tgctcag <b>gG</b> taacacatcagaa	13	16	4.6	YhdF (Table 2) paralog (41% ID)
<i>yjcE</i>	tgtgcc <b>gT</b> tttacaagaa-----acac <b>gG</b> gatcgcgtgctt	17	1.7	2.1	No described homolog; note suboptimal 10-bp spacer
<i>yjgC</i>	ttgtat <b>gT</b> tttatgagtt-gttgta <b>gG</b> gaaactgaaatagg	18	11	3.5	Formate dehydrogenase; divergent from <i>yjgB</i>
<i>yjgD</i>		11	11	2.6	HC
<i>yocB</i>	agtcag <b>gT</b> ttgatcgttt-ttaagag <b>gG</b> aaaaagaaaaacta	17	8.3	4.5	No described homolog
<i>ypuD</i>	tttac <b>gT</b> ttttatcca-tgaaaa <b>gG</b> aaactcatat	8.1	19	6.8	No described homolog
<i>yqzZ</i>	taaat <b>gT</b> tttaaatgaaa-aatgatc <b>gG</b> gtagtattctac	20	1.9	2.4	HC
<i>yqhB</i>	acacat <b>gT</b> tttatgagca-ttttcag <b>gT</b> ggaatggaatgtag	31	4.3	5.6	HC, family of five paralogs similar to hemolysins
<i>ytaB</i>	tcgggg <b>gT</b> ttgatatttataagata <b>gG</b> gtaataaataca	14			HC
<i>yunG</i>	gttctag <b>gT</b> ttttaaatactcatcaac <b>gT</b> ggtatcttttttta	2.6	1.4	2.1	No described homolog (possibly an operon with <i>yunFEDC</i> based on induction data)
<i>yuzA</i>	ataact <b>gT</b> tttaataatt---catgg <b>gG</b> aggttgcaaaac	11	8.3	3.9	HC
<i>yvaA</i>	agttag <b>gT</b> tttaccattt-gatcag <b>gG</b> gtataatactctg	4.7	1.6	1.6	Putative oxidoreductase; convergent with <i>yvaB</i>
<i>yvaG</i>	caatcag <b>gT</b> ttctgtcaa-taaata <b>gG</b> aatcaaaaacgg	10	2.9	1.5	Similar to 3-oxoacyl-acyl-carrier protein reductase; possibly some transcription into downstream genes ( <i>yvaFEDCB</i> )
<i>yvaK</i>	caaaa <b>gT</b> tttttctga-ttaaact <b>gT</b> gaaaaactaaaatg	3.6	0.72	1.0	70% identical to <i>Bacillus stearothermophilus</i> carboxylesterase (33)
<i>yvaJ(mr)</i>		5.1	1.3	1.2	Exoribonuclease (40)
<i>yvbG</i>	ataaag <b>gT</b> ttaccgggaaatcgctcc <b>gG</b> taaaaggggtgga	2.3	1.1	1.2	HC
<i>yvgN</i>	ttaagc <b>gT</b> attatttggtatcgctgag <b>gG</b> aatgtgagataa	2.7	4.5	3.1	Putative plant metabolite dehydrogenase, YtbE paralog
<i>yvgO</i>	tattgag <b>gT</b> ttacaataac-attgagc <b>gG</b> gtatgctgtagt	9.2	7.1	3.0	No described homolog (divergent from <i>yvgN</i> )
<i>yvgW</i>	gttttt <b>gT</b> ttttcattgacacttctt <b>gG</b> aaaaaacatata	4.8	12	6.9	Heavy-metal ATPase (downstream of <i>yvgZYX</i> )
<i>yvgZ</i>	acaac <b>gT</b> tttgacaatc-agtataat <b>gG</b> aattaatatcat	2.0	1.6	1.5	<i>yvgY</i> and <i>yvgX</i> weakly induced
<i>ywdD</i>	tcact <b>gT</b> ttcgtctttt---tcagg <b>gG</b> aaagagttagga	2.7	4.3	2.4	Possibly an operon with <i>ywdEF ung</i> , which are all also induced between two- and fourfold at 3' end
<i>ywiE</i>	tacaag <b>gT</b> ttatcgatta-gaaaa <b>gG</b> taatacagaggt	13	1.2	2.9	Probable cardiolipin synthase; downstream genes <i>ywjA</i> and <i>ywjB</i> induced ~3-fold, suggesting a possible operon structure
<i>ywsC(pgsB)</i>	agaga <b>gT</b> ttggcttagt---cgatt <b>gG</b> gaagattatgtta	3.4	1.4		Capsular polyglutamate synthesis (5)
<i>ywtG</i>	aaaa <b>gT</b> tttaattggccgg-aaaa <b>gG</b> ctaaaagatttct	20	3.2	4.2	49% ID to CsbC
<i>yxgB</i>	tcgc <b>gT</b> ttatcactgca--catagc <b>gG</b> gaagacaaataga	22	12	8.0	YcdF paralog (glucose-1-dehydrogenase)
<i>yxJ<sup>d</sup></i>	acagc <b>gT</b> tttttttgat--ctgctc <b>gG</b> aatggtacaatg	5.6	3.2		Divergent from <i>katX</i> ; similar to DNA-3-methyladenine glycosidase
<i>yxzF<sup>d</sup></i>	tagcat <b>gT</b> ttaaaggaagaggcaatcag <b>gG</b> ATGgttgagaa	18	13	6.8	Operon with <i>yxJ</i> ; start codon capitalized
<i>yxnA</i>	ttaaag <b>gG</b> gtaagacct-tccggat <b>gG</b> gtaatgtacaaaa	11	6.2	3.5	Similar to glucose-1-dehydrogenase
<i>yxvE</i>	cttgg <b>gT</b> tttttctcatt--cgaaag <b>gG</b> aaatgacgt	2.4	3.9		No described homolog

<sup>a</sup> Downstream genes in (putative) operons are indented.

<sup>b</sup> HC, hypothetically conserved; no described homolog, a unique protein found to date only in bacilli. Functional annotations are derived from the SubtiList database (37). ID, identity.

<sup>c</sup> *yfjS* is downstream of the strongly induced *yfjT* gene, but our experiments did not detect induction of the intervening *pel* gene.

<sup>d</sup> Data shown are from experiment 1.

We found strong heat shock induction for genes proximal to 16 of the proposed promoters (43), and the kinetics of induction are comparable to those of known members (class IIA) of the  $\sigma^B$  regulon (Table 3). These results support the previous suggestion that these genes are part of the  $\sigma^B$  regulon (43). Note that the *yfhK* gene is upstream of the  $\sigma^W$ -dependent operon *yfhLM*, and this promoter may be responsible for the heat induction of those genes as well (26).

Five genes (*ydbP*, *yoxA*, *ypuB*, *yqhQ*, and *yrvD*) shown to be inducible by ethanol in a  $\sigma^B$ -dependent manner (43) were not strongly induced by heat shock in our study. Nor was heat shock induction detected for *yqiS*, a gene induced by ethanol in both the wild-type and *sigB* mutant strains (Table 3). Finally, no data were obtained for *yotK* and *yycD*, as these genes were absent from the arrays used in these experiments. Additional experiments will be required to establish whether or not the putative  $\sigma^B$ -dependent promoters associated with these eight genes are in fact functional.

**Class IIC.** By sequence inspection, we propose 44 additional candidate  $\sigma^B$ -dependent promoters (likely controlling ~70 genes) proximal to newly identified heat shock genes (Table 4). In many cases, these candidate promoters are a good match to the  $\sigma^B$  consensus (43, 44) in both the  $-35$  and  $-10$  recognition elements. Indeed, 19 of these operons were independently proposed to be candidate members of the  $\sigma^B$  regulon, based on an analysis of genes induced by ethanol or by induction of  $\sigma^B$  expression, and 11 of these same promoters were identified using a hidden Markov model (45). Thus, it is likely that many, although probably not all, of the genes we have identified represent new members of the  $\sigma^B$  regulon. In some cases, for example, the candidate promoters we propose differ in potentially significant ways from the  $\sigma^B$  consensus, and these may be nonfunctional, chance occurrences. Interestingly, several of these genes encode paralogs of known members of the  $\sigma^B$  regulon (YwtG is 49% identical to CsbC; YxbG is 34% identical to YcdF; YdaB is 33% identical to YfhL; YxD is 41% identical to YhdF). As a class, these candidate  $\sigma^B$  regulon members include many predicted membrane proteins and transporters, functions consistent with the composition of the  $\sigma^B$  regulon as a whole.

Mapping all the known  $\sigma^B$  regulon members, together with additional likely members emerging from this study, onto the *B. subtilis* genome revealed three instances of clusters of transcriptional units. As many as nine  $\sigma^B$  consensus elements are clustered around the *ydaDEFG* operon (Fig. 2). This cluster includes the  $\sigma^B$ -dependent *gsiB* and *ydaP* genes (Table 2), the *ydaTS* and *ydbD* operons (Table 3), and the heat-induced *ydaB*, *ydaC*, and *ydaJKLMN* genes (Table 4). A second cluster occurs upstream of the *comG* operon (four promoters: *yqxL*, *yqhB*, *yqhA*, and *yqgZ*). The third cluster includes the *yfkM*, *yfkJIH*, *yfkF*, and *yfkED* operons. The vast majority of the remaining  $\sigma^B$ -dependent operons are apparently isolated or are occasionally found in small clusters of two or three operons.

Finally, analysis of the *yheK* gene leads us to propose a revision to the existing genome annotation (37). This gene displays the characteristic  $\sigma^B$  induction pattern, yet the best candidate  $\sigma^B$  promoter is situated with the  $-35$  region overlapping the assigned start codon (TTG). Sequence inspection identifies an alternative start site (ATG) at codon 19 of the *yheK* ORF. Furthermore, most YheK homologs lack the 18

TABLE 5. Fold induction of class III heat shock genes

Gene <sup>a</sup>	3 min	10 min	20 min
<i>ctsR</i> <sup>b</sup>	17	26	3.2
<i>yacH</i>	17	27	3.6
<i>yacI</i>	18	30	4.5
<i>clpC</i>	13	29	4.3
<i>sms</i>	5.1	9.6	2.2
<i>yacK</i>	4.2	7.2	1.8
<i>clpP</i>	6.4	21	6.8
<i>clpE</i>	63	88	25
<i>clpC</i>	13	29	4.3

<sup>a</sup> Downstream genes in (putative) operons are indented.

<sup>b</sup> Fold induction for *ctsR* obtained by RT-PCR was 22.2-fold (3 min), 19.8-fold (10 min), and 4.6-fold (20 min).

additional amino acids that would result from initiation at the assigned TTG start codon. We therefore suggest that translation of YheK begins with the ATG codon at position 19 and that the indicated promoter element may therefore be physiologically relevant (this new translation start site was also chosen in the latest annotation of the SubtiList database; release R16.1). Note that this gene has been redesignated *nhaX* and is proposed to form an operon with the downstream gene *nhaC* (formerly *yheL*) (62). However, there are no published data to support the suggestion of an operon structure, and 130 bp separate the *yheK* (*nhaX*) and *nhaC* genes. Moreover, we did not observe heat induction for *yheL*.

**The CtsR regulon (class III).** A subset of genes regulated by  $\sigma^B$  is also controlled by another heat shock pathway under control of CtsR (10, 32). CtsR is encoded by the first gene in the *ctsR* operon, which is transcribed from both  $\sigma^B$ - and  $\sigma^A$ -dependent promoters. We noted strong induction of the *ctsR* operon in this study (Table 5), but unlike that of  $\sigma^B$ -dependent heat shock genes, transcription of the *ctsR* operon peaked at the 10-min time point. The lower level of induction of the two promoter distal genes (*sms* and *yacK*) is consistent with recent data indicating that these genes are part of a separate,  $\sigma^M$ -dependent operon (A. Moir, personal communication). CtsR also regulates *clpP*, *clpE*, and *clpC* (10), which are strongly induced at the 10-min time point. This is in agreement with previous mRNA measurements that document a peak induction of *clpP* of ~28-fold between 6 and 9 min after a shift to 48°C (15). A similar pattern of induction was observed for *clpE* (9).

**The AhrC regulon.** One of the most unexpected findings in this study was the exceptionally strong transcriptional induction of three operons involved in arginine biosynthesis and transport (Fig. 3). There was no induction at the 3-min time point, but by 10 min after heat shock, all three operons were induced at least 50-fold. Independent confirmation of *argB* induction by quantitative RT-PCR showed over 900-fold induction, suggesting that the microarray experiment may underestimate the change in expression. Since both the arginine biosynthesis operons are repressed by the AhrC arginine-sensing transcription factor (8, 53), it is possible that heat shock induced a transient arginine deprivation. Alternatively, the AhrC protein itself may be temperature labile (12). The *yqiXYZ* operon was recently shown to encode an arginine transport system (52), and it also displays the same magnitude

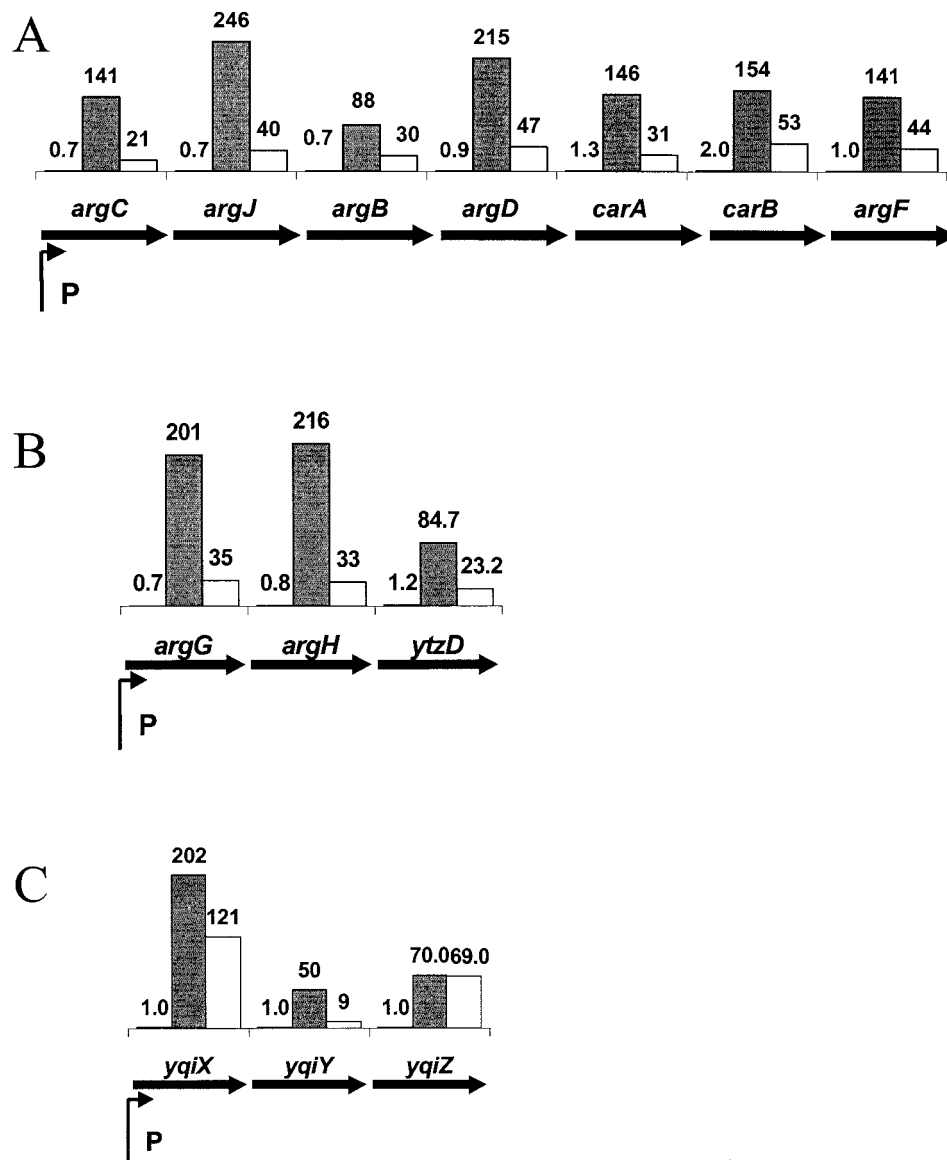


FIG. 3. Heat induction of arginine biosynthesis and uptake genes. Transcription of the *argC* biosynthetic operon (A), *argG* biosynthetic operons (B), and the *yqiX* arginine transport system (C) (52) is illustrated. The biosynthetic operons are repressed in response to arginine by the AhrC regulatory protein (8, 53). The *yqiXYZ* operon has been renamed *artPQM* to be consistent with *E. coli* nomenclature (52). Values for *argB* obtained by RT-PCR were 0.9-fold (3 min), 920-fold (10 min), and 328-fold (20 min). In the microarray studies, signals were not detected reproducibly for the 3-min points, presumably due to low message levels. However, those signals that were detected (four signals) were near 1, so these values have been shown as 1.0 for illustration purposes.

and kinetics of induction as noted for the two biosynthetic operons (Fig. 3C).

Since AhrC is also required as a positive activator of arginine catabolic genes (14, 29, 35), we also looked at the effects of heat shock on transcription of the *rocA*, *rocG*, and *rocD* operons. These operons are rapidly repressed following temperature shift and are among the most dramatically repressed genes in our analysis (mRNA levels declined by 3- to 20-fold after 10 min). This is consistent with a rapid (within 10 min) functional inactivation of the AhrC transcriptional activator combined with a short mRNA half-life for these transcripts.

**Other identified stress response genes (class U).** Many other genes have been identified as heat inducible in previous

studies but are regulated by as-yet-unknown mechanisms. Several of these genes were also found to be heat induced in our study, as shown in Table 6. For example, we detect an ~5-fold induction of both *ykda* (*htrA*) and *yvtA*, two heat-inducible HtrA paralogs regulated by unknown mechanisms from similar promoter elements (38).

An additional 66 members of the heat shock stimulon are not associated with obvious candidate promoter elements for  $\sigma^B$  or obvious recognition sites for known heat shock regulators (see supplementary material; Table S3 at <http://www.micro.cornell.edu/faculty.JHelmann.html>). All of these genes showed reproducible heat induction of at least 3.5-fold (or are cotranscribed with induced genes). Since regulatory pathways



TABLE 6. Class U (other): fold induction of known stress genes

Gene <sup>a</sup>	3 min	10 min	20 min	Comment(s) and/or reference
<i>ahpF</i>	1.5	1.6	1.8	<i>ahpC</i> not on array (3); <i>ahpC</i> not on array (3)
<i>clpX</i>	0.85	1.3	1.1	16
<i>ftsH</i>	1.1	1.5	1.4	11
<i>htrA</i>	3.2	5.8	5.5	51
<i>lonA</i>	4.9	3.1	2.9	= <i>ykdA</i> (39)
<i>yvtA</i>	1.6	2.3	1.6	46
<i>yvtB</i>	4.8	1.9	2.1	HtrA paralog (38)
<i>ywcG</i>	5.0	1.7	1.7	
<i>ywcH</i>	4.7	6.9	3.3	= <i>nfrA</i> (36)
	3.3	4.5	2.0	

<sup>a</sup> Downstream genes in (putative) operons are indented.

for these genes are not known, we assigned them to class U. Interestingly, several of these genes encode transport functions, including the *appDFABC* operon, one of two oligopeptide uptake systems in *B. subtilis* (30, 31). Other transporters induced by heat shock include a choline ABC transporter (*opuB* operon), a putative Na<sup>+</sup>/nucleoside cotransporter (*yutK*) and a multidrug efflux homolog (*yuxJ*). We also note heat induction of a subset of the S-box regulon (18) including specifically those genes implicated in methionine biosynthesis. Additional work will be required to determine the mechanism and relevance of this heat induction.

**Gene signals arising from proximity effects.** In addition to increased transcription due to heat shock, some of the signals detected in these experiments may arise from what we generically call proximity effects. For example, transcription termination at the end of operons is often less than 100% efficient, and these read-through transcripts may lead to signals corresponding to genes downstream of strongly induced heat shock genes. If the downstream gene is codirectional with the heat shock gene, these signals could be physiologically relevant. However, in some cases, the downstream gene is convergent with the heat shock gene and the transcript through this region is anti-sense. These are nevertheless detected using random hexamer priming and could give rise to spurious signals. Two likely examples that emerged in this study are the *yfkQ* operon and the *yknA* gene (Table 7). All four genes of the *yfkQ* operon showed some heat induction, but there was a clear gradient, with the largest apparent induction near the end of the operon. Since this operon is convergent with the strongly induced,  $\sigma^B$ -dependent *yflA* gene (Table 3), this pattern is consistent with read-through transcription from *yflA* giving rise to (gradually diminishing) antisense RNA through this region.

To test this model, readthrough transcription from *yflA* into

TABLE 7. Apparent heat induction of convergent operons due to readthrough transcription

Gene <sup>a</sup>	3 min	10 min	20 min	Comment
<i>yfkQ</i>	3.2	1.3	1.2	<i>yfkQ</i> operon convergent with <i>yflA</i>
<i>yfkR</i>	7.7	1.2		
<i>yfkS</i>	11	1.9	2.4	
<i>yfkT</i>	20	3.2		
<i>yknA</i>	8.9	4.2	2.3	Convergent with <i>ykzA</i>

<sup>a</sup> Downstream genes in (putative) operons are indented.

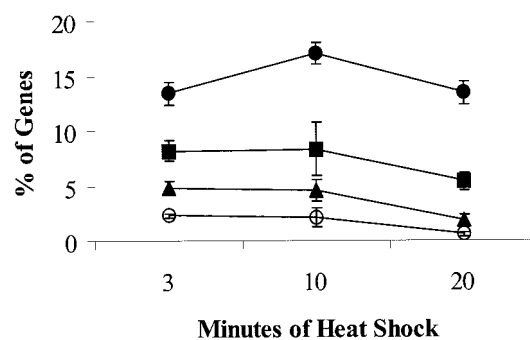


FIG. 4. Overview of heat shock stimulum. The percentage of genes induced after heat shock was calculated for all three experiments for cells sampled at 3, 10, and 20 min after the shift from 37 to 48°C. The percentage of genes ( $\pm 1$  standard deviation) that are induced by at least 2-fold (closed circles), 3-fold (squares), 5-fold (triangles), or 10-fold (open circles) is shown. Percentages were determined by dividing the number of induced genes by the total number for which mRNA was detected at a level significantly above background (the average numbers of total genes [over all three time points] for which a signal was detected were 2,528 [experiment 1], 2,489 [experiment 2], and 3,204 [experiment 3]).

*yfkT* was measured by a modification of the standard Taqman quantitative RT-PCR protocol. RT was conducted with either a sense or an antisense primer, after which time the RNA was digested and reverse transcriptase was inactivated. The opposing primer was then added and quantitative PCR was carried out. RT with the sense primer (i.e., priming off the antisense strand) showed a 29-fold induction at the 3-min time point, whereas the antisense primer showed only a threefold induction (data not shown). This result is consistent with a proximity effect whereby the apparent induction of *yfkT* by heat shock is primarily due to read-through from *yflA* with the antisense strand of *yfkT* being transcribed. Similarly, the apparent induction of *yknA* may result from the fact that this gene is convergent with the strongly induced *ykzA* gene (Table 2). Although both the *yknA* gene and the *yfkQ* operon were considered good candidates for the  $\sigma^B$  regulon on the basis of transcriptional profiling studies (45), our findings suggest that a reinterpretation of these data is in order. Similar proximity effects have been noted in microarray studies of *E. coli* (28, 66). It is possible to avoid this complication by using 3'-end, gene-specific primers for RT. However, as discussed in detail elsewhere (4), this approach does not uniformly label all mRNAs and therefore provides a more limited picture of the transcriptome.

**Summary.** The transfer of *B. subtilis* from 37 to 48°C elicits a very large transcriptional response coordinated by several distinct transcription factors (19–21, 44). In the studies described here, we document the heat induction of hundreds of genes and independently confirm the microarray data for four genes by quantitative RT-PCR. Over 5% of the transcriptionally active genes are induced at least threefold, and well over 10% of the genome displays a measurable induction in response to heat shock (Fig. 4).

Activation of the  $\sigma^B$  regulon is the single largest component of the heat shock response in *B. subtilis* (19, 44). We have measured the induction of 70 known or previously proposed members of the  $\sigma^B$  regulon (Tables 2 and 3; Fig. 1 and 2) and identified another 72 candidate  $\sigma^B$  regulon members (Table 4).

Our heat shock data provides additional support for many, albeit not all, members of the  $\sigma^B$  regulon proposed previously on the basis of consensus search procedures (43) and transcriptional profiling studies (45).

As expected, heat induction of the CtsR (Table 5) and HrcA (Table 1) regulons is apparent, and other known heat shock proteins (Table 6) are also induced. Finally, we can assign many new genes to the heat shock stimulon (Table S3 [http://www.micro.cornell.edu/faculty.JHeilmann.html]), though the factor(s) mediating their heat induction are not clear at present. Prominent among these genes are three operons involved in arginine biosynthesis and transport (Fig. 3). Induction of these genes may reflect an in vivo temperature lability of the AhrC regulatory protein, an idea supported by the decrease in expression of the AhrC-dependent arginine catabolic genes.

Our analysis provides further evidence of the power and utility of microarray approaches to defining bacterial stimulons and regulons. As we extend this work to include other stimulons, a thorough knowledge of the heat shock activated general stress response will be very useful in distinguishing specific from more general transcriptional responses.

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