DnaK, DnaJ, and GrpE heat shock proteins negatively regulate heat shock gene expression by controlling the synthesis and stability of \( \sigma^{32} \)

David Straus, William Walter, and Carol A. Gross

Department of Bacteriology, University of Wisconsin, Madison, Wisconsin 53706 USA

The *Escherichia coli* DnaK heat shock protein has been identified previously as a negative regulator of *E. coli* heat shock gene expression. We report that two other heat shock proteins, DnaJ and GrpE, are also involved in the negative regulation of heat shock gene expression. Strains carrying defective *dnaK*, *dnaj*, or *grpE* alleles have enhanced synthesis of heat shock proteins at low temperature and fail to shut off the heat shock response after shift to high temperature. These regulatory defects are due to the loss of normal control over the synthesis and stability of \( \sigma^{32} \), the alternate RNA polymerase \( \sigma \)-factor required for heat shock gene expression. We conclude that DnaK, DnaJ, and GrpE regulate the concentration of \( \sigma^{32} \). We suggest that the synthesis of heat shock proteins is controlled by a homeostatic mechanism linking the function of heat shock proteins to the concentration of \( \sigma^{32} \).

[Key Words: Heat shock proteins; DnaK heat shock gene expression; \( \sigma^{32} \)]

Received August 9, 1990; revised version accepted September 12, 1990.

The induction of heat shock proteins following an abrupt increase in growth temperature has been observed in every cell type examined, including examples from eubacterial, archaebacterial, and eukaryotic organisms [Schlesinger et al. 1982; Craig 1985]. Heat shock proteins are also induced by a variety of other stresses, including exposure to ethanol, UV irradiation, oxidative agents, viral infection, and the presence of abnormal proteins (for review, see Neidhardt and VanBogelen 1987). Recent work suggests that heat shock proteins provide functions that are essential in the control of protein folding [Chirico et al. 1988; Deshaies et al. 1988; Goloubinoff et al. 1989a; Gaitanaris et al. 1990], assembly [Goloubinoff et al. 1989b], and disassembly [Alfano and McMacken 1989; Dodson et al. 1989; Zyliec et al. 1989]. It is likely that heat shock proteins carry out similar functions in most cells, as there is a high degree of sequence homology between heat shock proteins from widely divergent organisms [Bardwell and Craig 1984, 1987; McMullin and Hallberg 1988].

When *Escherichia coli* are shifted from low to high growth temperature, ~17 heat shock proteins are induced [Neidhardt and VanBogelen 1987]. This induction is transient, peaking at 5–15 min after upshift and the dropping to a new steady-state rate of synthesis, which is characteristic of the new growth temperature. The expression of heat shock proteins, both at low temperature and after temperature upshift, depends on the function of the *rpoH(\( \sigma^{32} \))* gene [Neidhardt and VanBogelen 1981; Yamamori and Yura 1982; Zhou et al. 1988]. The product of this gene has been identified as a \( \sigma \)-factor, \( \sigma^{32} \), which enables RNA polymerase to recognize the promoters for heat shock genes [Grossman et al. 1984, Cowing et al. 1985; Fujita et al. 1987]. In addition to directing RNA polymerase to the heat shock genes, \( \sigma^{32} \) also regulates their expression. A rapid and transient increase in the level of \( \sigma^{32} \) is observed following temperature upshift, which can account for the induction of heat shock proteins [Lesley et al. 1987; Skelly et al. 1987; Straus et al. 1987]. Additionally, artificial induction of \( \sigma^{32} \) without a temperature shift results in the induction of heat shock proteins [Grossman et al. 1987]. The increase in \( \sigma^{32} \) level that accompanies a temperature upshift occurs as a result of changes in both the stability and synthesis of \( \sigma^{32} \) [Straus et al. 1987]. It is not known how an abrupt change in temperature causes an increase in the synthesis and stability of the alternate \( \sigma \)-factor.

Tilly et al. (1983) observed that a mutation in the *dnaK* heat shock gene resulted in enhanced heat shock gene expression at low temperature and an extended heat shock response after shift to high temperature, suggesting that DnaK is a negative regulator of the heat shock response. We have examined the phenotype of other heat shock gene mutants besides *dnaK* and found that *dnaj* and *grpE* mutants have a similar effect on heat shock gene expression. We have determined the mechanism for this effect; all three mutants are defective in

\(^{a}\)Present address: Howard Hughes Medical Institute, University of California School of Medicine, San Francisco, California 94143 USA.
the control of $\sigma^{32}$ synthesis and stability. These results indicate that heat shock gene expression is controlled by a negative feedback loop in which heat shock proteins regulate their own synthesis by controlling the level of $\sigma^{32}$. We suggest that temperature changes are sensed by this homeostatic regulatory mechanism through alteration in the function of DnaK, DnaJ, and GrpE.

**Results**

**Expression of heat shock genes is enhanced in dnaI and grpE mutants**

The regulation of heat shock protein synthesis is altered in two ways in strains carrying the dnaK756 allele [Tilly et al. 1983]. First, synthesis of heat shock proteins in these strains is increased during growth at 30°C. Second, the shutoff phase of the heat shock response is delayed. To determine whether this effect is specific to the dnaK mutant, we examined the synthesis of heat shock proteins in strains carrying mutations in other heat shock genes. Our results indicate that of the strains analyzed, only mutations in dnaI or grpE have a significant effect on heat shock protein synthesis.

Heat shock protein synthesis in the mutant strains was analyzed by two-dimensional gel electrophoresis of extracts from cells that had been pulse-labeled with radioactive amino acids at 30°C or at various times after shift to 42°C. As compared to the parental strain, the dnaK756, dnaI259, and grpE280 strains all showed a two- to fourfold increase in the 30°C synthesis rate of the four heat shock proteins analyzed. This effect was specific to the heat shock proteins, as the synthesis of EF-Tu [Table 1] and other non-heat shock proteins [data not shown] was unaltered in the mutants. As is the case for dnaK756, the dnaI259 and grpE280 mutants were also defective in the shutoff phase of the heat shock response following shift from 30 to 42°C. As exemplified by the induction kinetics of the GroE heat shock protein, each of the mutants show only a gradual drop in heat shock protein synthesis [Fig. 1]. GroEL synthesis remains elevated for at least 60 min after temperature shift in the mutants. In wild-type cells, induction of GroEL is transient, peaking at ~5 min after temperature upshift. The shutoff of synthesis of the other heat shock proteins followed similar kinetics in the mutant strains [data not shown]. These findings indicate that dnaK is not the only heat shock gene involved in the regulation of heat shock protein synthesis. Mutations in the dnaI and grpE heat shock genes also result in altered synthesis of heat shock proteins at both 30°C and during shutoff of the heat shock response.

Synthesis of heat shock proteins in strains carrying mutations in other heat shock genes was also examined. The lon146::Tn10, groEL140, and groES30 alleles had only small effects on heat shock protein synthesis. These strains showed an increase of ~50% in heat shock protein synthesis at 30°C and exhibited only a slight delay in shutting off the heat shock response. Data for the lon146::Tn10 strain is shown in Figure 2.

The induction of heat shock protein synthesis following temperature upshift is the result of increased heat shock gene transcription [Yamamori and Yura 1982; Taylor et al. 1984; Cowing et al. 1985]. To determine whether altered heat shock gene expression in the mutants could be explained by altered rates of transcription, we examined derivatives of dnaK756, dnaI259, and grpE280, which carried a plasmid in which the expression of galactokinase is driven by the groE promoter. In these strains the rate of galactokinase synthesis indicates the rate of transcription initiation at the groE pro-

**Table 1. Heat shock protein synthesis is enhanced in heat shock gene mutants at 30°C**

<table>
<thead>
<tr>
<th></th>
<th>DnaK</th>
<th>GroEL</th>
<th>HtpG</th>
<th>HtpM</th>
<th>EF-Tu</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild type</td>
<td>1 ± 0.03</td>
<td>1 ± 0.05</td>
<td>1 ± 0.01</td>
<td>1 ± 0.11</td>
<td>1 ± 0.06</td>
</tr>
<tr>
<td>dnaK756</td>
<td>1.93 ± 0.04</td>
<td>1.93 ± 0.02</td>
<td>2.28 ± 0.15</td>
<td>1.59 ± 0.12</td>
<td>0.95 ± 0.01</td>
</tr>
<tr>
<td>grpE280</td>
<td>2.54 ± 0.20</td>
<td>2.07 ± 0.03</td>
<td>2.46 ± 0.06</td>
<td>1.82 ± 0.20</td>
<td>0.98 ± 0.01</td>
</tr>
<tr>
<td>dnaI259</td>
<td>4.40 ± 0.33</td>
<td>2.59 ± 0.06</td>
<td>4.03 ± 0.20</td>
<td>2.90 ± 0.41</td>
<td>1.00 ± 0.01</td>
</tr>
</tbody>
</table>

The relative synthesis rate of the DnaK, GroEL, HtpG, and HtpM heat shock proteins and EF-Tu were determined in isogenic derivatives of C600, as described in Materials and methods. The rates for each protein are normalized to the rate of synthesis in the wild-type strain. The values are the average of triplicate determinations ± the standard error.

![Figure 1. Shutoff of the heat shock response is defective in the dnaK756, grpE280, and dnaI259 mutants. The relative synthesis of GroEL in wild-type (●), dnaK756 (○), grpE280 (■), and dnaI259 (□) strains was determined at 30°C or at 10, 20, and 60 min after shift to 42°C by pulse-labeling and two-dimensional gel electrophoresis, as described in Materials and methods. The synthesis rates are normalized to the 30°C wild-type rate. Shutoff of DnaK, HtpG, and HtpM synthesis was also defective in the mutants, and the relative magnitude of the defects in the different mutants was the same as that observed for GroEL synthesis.](genesdev.cshlp.org)
The relative synthesis of GroEL in wild type (●) and lon146::Tn10 (▲) was determined at 30°C or at 10, 20, and 60 min after shift to 42°C, as described in Fig. 1. DnaK, HtpG, and HtpM synthesis was similar to that shown for GroEL.

We measured the amount of σ^32 directly to confirm that the level of σ^32 increases in the mutant strains. As predicted from the stability measurements, the increase in σ^32 concentration in the mutants is actually greater than the increase in heat shock gene expression. Western blot analysis indicates that the level of σ^32 in the mutants is 10- to 20-fold greater at 30°C than in the wild-type strain (data not shown), whereas the synthesis of heat shock proteins in the mutants is only 2- to 4-fold greater than in the wild-type strain (Table 1). We have found that the apparent activity of σ^32 is decreased when there is an excess of heat shock proteins [Straus et al. 1989]. The difference between the increase in σ^32 level and the increase in the heat shock gene expression in the mutants is probably due to negative regulation of σ^32 activity. Initially, the enhanced stability of σ^32 in the mutants would cause a proportionate overproduction of heat shock proteins, but under steady-state conditions the activity of σ^32 would be decreased such that the increase in heat shock protein synthesis would be only a fraction of the increase in the level of σ^32.

Changes in both the stability and synthesis of σ^32 contribute to the altered heat shock response in the mutant strains

Changes in both the rates of degradation and synthesis of σ^32 normally contribute to the turn-on and shutoff of the heat shock response. We therefore determined

null phenotype, because neither a dnaK nor a dnaJ deletion mutant showed an increase in σ^32 synthesis at 30°C [data not shown]. These data indicate that the increased transcription of heat shock genes observed in the dnaK, dnaJ, and grpE strains at 30°C results solely from their inability to degrade σ^32 rapidly.

Figure 3. Transcription from the groE heat shock promoter is increased in the dnaK, grpE, and dnaJ mutants. Cultures of wild-type (●), dnaK756 (□), grpE280 (■), and dnaJ259 (□) strains carrying plasmid pDC441 were pulse-labeled with [35S]methionine for 1 min either at 30°C or at 5, 10, 20, 40, and 60 min after shift to 42°C. The synthesis of galactokinase relative to total protein synthesis was determined by immunoprecipitation, as described in Materials and methods.
DnaK, DnaJ, and GrpE regulate concentration of α32

Figure 4. Degradation of α32 is defective in the dnaK756, grpE280, and dnaJ259 mutants. Stability of α32 in isogenic wild-type (●), dnaK756 (○), grpE280 (■), and dnaJ259 (□) strains. Cultures were labeled with [35S]methionine for 1 min either at 30°C (A) or at 3 min after shift from 30 to 42°C (B), and the fraction of α32 remaining at various times after addition of unlabeled methionine was determined by immunoprecipitation as described in Materials and methods. Approximate half-lives for α32 in the different strains are indicated.

whether either of these processes was altered in the mutants following temperature upshift. In contrast to the results at 30°C, altered regulation of both synthesis and stability of α32 contribute to delayed shutoff after shift to 42°C.

Each of the mutant strains affect the synthesis of α32 in much the same way that they alter the synthesis of heat shock proteins during the shutoff of the heat shock response. In wild-type strains, α32 synthesis is induced, peaks at 3–4 min after temperature upshift, and drops to a lower rate by 6 min after upshift. In the mutant strains the synthesis of α32 is induced normally but remains elevated up to 15 min after shift from 30 to 42°C (Fig. 5). Measurement of α32 stability during the shutoff phase of the heat shock response indicates that the prolonged heat shock response of the dnaK756 and grpE280 mutants is also due to an increase in α32 stability in these strains (Fig. 4B). In wild-type strains α32 is stable for the first 4 min following shift from 30 to 42°C and then returns to the rapid rate of degradation seen prior to temperature shift [t1/2 = 1 min]. In contrast, the dnaK756 and grpE280 mutants continue to degrade α32 slowly [t1/2 = 5–10 min] during the shutoff phase of the heat shock response. Note that the dnaJ259 mutant has no defect in α32 degradation at 42°C; this strain exhibited the same rapid rate of α32 degradation as did the wild type during the shutoff phase of the heat shock response (Fig. 4B). Because a dnaJ deletion strain is also proficient in degradation of α32 at 42°C, lack of a proteolysis phenotype cannot be explained by suggesting that the dnaJ259 allele is partially functional at 42°C (data not shown). In summary, the altered regulation of the heat shock proteins observed in the mutants after shift from 30 to 42°C results from an increase in both the synthesis and stability of α32 in the dnaK756 and grpE280 strains. Altered regulation in the dnaJ259 mutant results solely from an increase in the rate of α32 synthesis, perhaps accounting for the fact that dnaJ mutants show less induction of α32 during the heat shock response.

Figure 5. Repression of α32 synthesis during shutoff of the heat shock response is defective in the dnaK756, grpE280, and dnaJ259 mutants. Cultures of wild-type (●), dnaK756 (○), grpE280 (■), and dnaJ259 (□) strains were pulse-labeled with [35S]methionine for 1 min either at 30°C or at 3, 6, 10, and 15 min after shift to 42°C. The synthesis of α32 relative to total protein synthesis was determined by immunoprecipitation, as described in Materials and methods, and corrected for a 2.5-fold increase in total protein synthesis following temperature upshift to derive the change in absolute synthesis. The values represent the average of at least four experiments normalized to the maximum rate of synthesis within each strain following temperature upshift. The average maximum increase in the rate of α32 synthesis in wild-type strains was similar in all four strains: Wild type increased 11.1-fold while dnaK756, grpE280, and dnaJ259 increased 9.7-, 12.2-, and 9.5-fold, respectively.
heat shock proteins after temperature upshift than dnaK and grpE mutants [Fig. 1].

We have found previously that the transient induction of $\sigma^{32}$ synthesis that occurs following a shift from 30 to 42°C is due to translational regulation. This conclusion was based on two observations: (1) Induction of $\sigma^{32}$ synthesis is transient even when the synthesis of $\sigma^{32}$ is directed by the $\lambda_p$ promoter instead of the normal rpoH promoters [Grossman et al. 1987]; and (2) correct regulation of an rpoH–$\lambda$-lacZ fusion after temperature upshift requires that its expression be governed by rpoH translational signals [Straus et al. 1987]. Previously, we used the $\lambda_p$ fusion to rpoH to demonstrate that the dnaK756 mutant was defective in this translational regulation [Grossman et al. 1987]. We used the same $\lambda_p$ fusion to rpoH to determine whether dnaJ259 and grpE280 alleles also affected translational regulation. Immunoprecipitation analysis indicated that synthesis of $\sigma^{32}$ in the wild-type strain peaked at 10 min following temperature upshift and was then repressed while synthesis of $\sigma^{32}$ in the mutants remained elevated following induction [Fig. 6]. This finding indicates that the alteration in $\sigma^{32}$ synthesis observed in the heat shock gene mutants is probably due to a defect in translational regulation.

Discussion

Characterization of the dnaK, dnaJ, and grpE mutants indicates that the DnaK, DnaJ, and GrpE heat shock proteins negatively regulate heat shock gene expression. The mutants exhibit increased expression of heat shock proteins at 30°C and fail to shut off the heat shock response after shift to high temperature. We have determined the mechanism for these effects. In every case, the key regulatory feature is the overproduction of $\sigma^{32}$ in the mutants relative to the wild-type strain. Each of the mutant strains is defective in degrading $\sigma^{32}$ at 30°C. The resultant accumulation of $\sigma^{32}$ leads to overexpression of heat shock proteins at low temperature. Tilley et al. [1989] have also found that dnaK mutants are defective in degrading $\sigma^{32}$ at 30°C. The mechanism underlying the delayed shut-off of the heat shock response after shift to high temperature is more complex. Each of the mutant strains continues synthesizing $\sigma^{32}$ at a high rate for an extended time after shift to 42°C because they are defective in translational regulation of $\sigma^{32}$ synthesis. In addition, the dnaK756 and grpE280 mutants are also defective in the degrading $\sigma^{32}$ at high temperature. As a consequence, $\sigma^{32}$ levels remain elevated for an extended time after temperature upshift in the mutants leading to a delay in the shut-off of the heat shock response.

Of the major heat shock proteins, DnaK, DnaJ, and GrpE appear to be uniquely involved in regulating the heat shock response because among the mutations tested, only mutations in these genes affect $\sigma^{32}$ synthesis and stability. It is possible that additional, yet-to-be-discovered heat shock proteins will also be involved in regulating $\sigma^{32}$. DnaK, DnaJ, and GrpE have been thought to function together ever since their discovery as host proteins required for $\lambda$ replication. Biochemical analysis indicates that these proteins act in the formation of a productive initiation complex by disassembling an intermediate protein complex at the $\lambda$ origin of replication [Alfano and McMacken 1989; Dodson et al. 1989; Zylicz et al. 1989]. These studies make it unlikely that these heat shock proteins are directly responsible for degrading $\sigma^{32}$ and/or regulating its synthesis. This viewpoint is supported by additional investigations demonstrating that DnaK, DnaJ, and GrpE hinder denatured or partially unfolded proteins such as $\lambda$ repressor [Gaitanaras et al. 1990]. Thus, these three heat shock proteins probably alter the interactions between $\sigma^{32}$ and the proteins directly responsible for mediating its degradation and translational regulation. For example, DnaK, DnaJ, and GrpE could increase the degradation of $\sigma^{32}$ by maintaining it in a partially denatured state to facilitate the function of a protease.

The existence of a negative feedback loop coupling the function of heat shock proteins to the concentration of $\sigma^{32}$ suggests that a homeostatic mechanism is responsible for the regulation of heat shock gene expression. In such a model, the involvement of the DnaK, DnaJ, and GrpE proteins in the negative regulation of $\sigma^{32}$ synthesis and stability permits the cell to monitor whether the amount of these proteins is adequate for cell growth. In the simplest version of this model, the “thermometer” utilized by the cell is actually the free pool of the DnaK, DnaJ, and GrpE proteins themselves. An increase in growth temperature would deplete the “free pool” of DnaK, DnaJ, and GrpE as a consequence of an increased demand for these proteins. Some potential substrates that could transiently decrease the free pool of DnaK, DnaJ, and GrpE are newly synthesized proteins that are
partially or completely unfolded, preexisting proteins that are partially denatured upon temperature upshift, and increased amounts of unstable proteins. As a result of the undersupply of the three heat shock proteins, cells would become defective in σ32 degradation and translational regulation of σ32 expression. This response would be self-limiting because increased levels of heat shock proteins would allow the cell to re-establish negative regulation of σ32.

The regulation of σ32 stability during the heat shock response is explained simply by a homeostatic mechanism, however, an additional postulate is required to explain how this mechanism can account for the transient derepression of σ32 synthesis during the heat shock response. Because the DnaK, DnaJ, and GrpE proteins are not required for regulation of σ32 synthesis at low temperature, titration of heat shock protein function following temperature upshift, by itself, would be insufficient to explain the induction of σ32 synthesis. An additional event must be responsible for making the regulation of σ32 synthesis dependent on heat shock protein function at high temperature. As an example of such a mechanism, a translational repressor of σ32 synthesis might be inherently temperature sensitive, becoming partially denatured upon shift to high temperature. In this case, accumulation of the DnaK, DnaJ, and GrpE heat shock proteins would then provide the functions necessary to repress the repressor and re-establish the repression of σ32 synthesis required for shutoff of the heat shock response.

How likely is it that temperature is sensed by a homeostatic mechanism linking heat shock protein function to regulation? Two basic requirements of this model are that the heat shock proteins be present in limiting concentrations in the cell and that demand for heat shock proteins increases as the temperature rises. The fact that the maximum growth temperature of E. coli is directly correlated with the capacity to synthesize heat shock proteins indicates both that heat shock proteins are limiting and that more of them are required at higher growth temperatures [Yamamori and Yura 1982]. The finding that increasing the amount of heat shock proteins either at 30 or 42°C increases the ability of cells to degrade puromycyl fragments at both temperatures is consistent with the idea that heat shock proteins are present in limiting concentration [Straus et al. 1988]. Notably, addition of puromycin induces heat shock gene expression [Goff and Goldberg 1985], as would be expected if a homeostatic mechanism controls gene expression. Evidence that such a mechanism specifically involves DnaK, DnaJ, and GrpE comes from experiments with λ repressor. Parsell and Sauer [1989] have demonstrated that the heat shock response is induced by overexpression of a mutated amino-terminal fragment of λ repressor that is denatured but stable in E. coli at 30°C. The Gottesman group has found that renaturation of λ repressor is defective in dnaK, dnaJ, and grpE mutant strains both in vivo and in crude extracts and that the in vitro defect is restored by extracts from wild-type cells suggesting that denatured λ repressor is a substrate of DnaK, DnaJ, and GrpE [Gaitanaris et al. 1990]. Taken together, these experiments indicate a direct link between depletion of the free pool of DnaK, DnaJ, and GrpE and induction of the heat shock response.

Reports from several groups suggest that DnaK homologs, the Hsp70 proteins, play a special role in regulating the eukaryotic heat shock response. Early studies showed that when accumulation of functional Hsp70 protein was blocked, the heat shock genes were generally overtranscribed [DiDomenico et al. 1982]. More recent studies directly implicate Hsp70 as a negative regulator of the heat shock response. Underexpression of Hsp70, accomplished by disrupting the yeast SSA1 and SSA2 genes (encoding Hsp70 proteins) results in increased transcription of the heat-inducible SSA3 and SSA4 genes [Boorstein and Craig 1990, Stone and Craig 1990]. Moreover, this increase is mediated by the heat shock element, the same DNA sequence responsible for the increased transcription observed during the heat shock response. Finally, recent studies from the Welch group suggest that while Hsp70 associates only transiently with newly synthesized proteins, it remains permanently associated with amino acid analog-containing proteins or protein fragments generated by puromycin [Beckmann et al. 1990]. Because treatment with amino acid analogs or puromycin induces the synthesis of Hsp70, these experiments indicate that accumulation of an Hsp70 substrate is sufficient to induce the heat shock response. The cellular response to temperature upshift and many of the heat shock proteins themselves are conserved throughout evolution. It would be remarkable if the logic used by cells to sense temperature change had not also been conserved.

Materials and methods

Strains

Strains carrying dnaK756-thr :: Tn10, dnaJ259-thr :: Tn10, grpE280-tyrA :: Tn10, groE530-zid :: Tn10, and groEL140-zid :: Tn10 were from C. Georgopolous, and lon146 :: Tn10 was from S. Gottesman. All of these alleles were transduced into a derivative of strain C600 [galK, leu, lacZ90, lacY, tonA, thi, thr, supE44/F'locP, lacZ :: Tn5], using PIvir, selecting for tetracycline resistance, and scoring for temperature sensitivity, except for lon146 transductants, which were selected for tetracycline resistance and examined for mucoidy. For the experiment in Figure 3, the dnaK756, dnaJ259, and grpE280 alleles were transduced into a derivative of C600 carrying pDC441 and the pcanB8 mutation (to reduce plasmid copy number) using PIvir, by selecting for the kanamycin resistance of a linked Tn10[kan] marker and scoring for temperature sensitivity. Plasmid pDC441 contains sequences from -60 to +77 of the groE promoter cloned into the Smal site of plasmid pK01 such that expression galK is under control of the groE promoter. For the experiment in Figure 6, the dnaK756, dnaJ259, and grpE280 alleles were transduced into strain N5242 [bio, xyl857, Δbam, Δhif1, Δg] carrying pAG37. Plasmid pAG37 [Grossman et al. 1984] places expression of rpoH under control of the Δhif pro- moter. Strain PK101 carries a dnaK deletion, and strain PK102 carries a dnaJ deletion [Kang and Craig 1990]. For the experiments investigating the null phenotype of these strains, MG1655 was used as the isogenic wild type. Strains were grown...
in M9-glucose media supplemented with all amino acids except those used for radiolabeling. Temperature upshifts were accomplished by rapidly transferring flasks from a 30°C shaking water bath to a 42°C shaking water bath.

**Protein synthesis**

The relative synthesis rates of heat shock proteins were determined by pulse-labeling and two-dimensional gel analysis. Cell culture aliquots of 0.5 ml were pulse-labeled with 70 μCi of [³⁵S]methionine for 1 min, and sampled directly into 0.11 ml of 10% trichloroacetic acid (TCA). Samples were analyzed by two-dimensional gel electrophoresis (O'Farrell 1975), following the addition of aliquots of cell lysate from [³⁵S]methionine-labeled cultures, which served as a standard for double-label quantitation of the heat shock proteins. The radioactive protein spots were excised from the two-dimensional gels, solubilized, and counted. The synthesis rates of the DnaK, GroEL, HspG (C62.5), and HspM (F84.1) heat shock proteins and EF-Tu were determined relative to total protein synthesis as described previously (Grossman et al. 1987).

σ^d^ synthesis and stability

The synthesis and stability of σ^d^ was determined by pulse-labeling and immunoprecipitation as described previously (Straus et al. 1987). For measurements of the relative synthesis rate of σ^d^, 1-ml aliquots of cell culture were labeled with 70 μCi of [³⁵S]methionine for 1 min and sampled directly into 0.11 ml of 50% TCA. For measurements of σ^d^ stability, cultures were labeled with 70 μCi/ml [³⁵S]methionine for 1 min and chased with excess unlabeled methionine, and 1-ml aliquots were sampled directly into TCA. σ^d^ was immunoprecipitated from aliquots of each sample representing equal counts per minute. As a correction for losses in the immunoprecipitation analysis, equal amounts of a mutant σ^d^ protein, which migrates more rapidly than the wild-type protein, were added to each sample. The mutant protein was supplied from a lysate of an [³⁵S]methionine-labeled strain that overproduces the mutant protein. Background estimates were made from samples of a wild-type culture that had been pulse-labeled and chased for 20 min during which time >99% of the σ^d^ is degraded. The immunoprecipitates were analyzed on SDS–polyacrylamide gels and the σ^d^ bands were excised and counted.

Galactokinase synthesis

Galactokinase synthesis was determined by pulse-labeling and immunoprecipitation, as described previously (Straus et al. 1987), except that aliquots of mid-log phase cultures were labeled with 20 μCi of [³⁵S]methionine and that the ratio of counts per minute in the standard cells [labeled with [³⁵S]methionine and [³⁵S][lysine] to the experimental cells was 0.25 : 1.

**Acknowledgments**

We thank C. Georgopoulos for providing numerous strains, B. Craig and H. Echols for their excellent suggestions, and National Institutes of Health grant RO1GM-36278 for supporting this work.

The publication costs of this article were defrayed in part by payment of page charges. This article must therefore be hereby marked "advertisement" in accordance with 18 USC section 1734 solely to indicate this fact.

**References**


Kang, P.J. and E.A. Craig. 1990. Identification and characterization of a new *Escherichia coli* gene that is dosage-dependent.
DnaK, DnaJ, and GrpE regulate concentration of σ22


Zhou, Y.N., N. Kusukawa, J. Erickson, C. Gross, and T. Yura. 1988. Isolation and characterization of Escherichia coli mu-
DnaK, DnaJ, and GrpE heat shock proteins negatively regulate heat shock gene expression by controlling the synthesis and stability of sigma 32.

D Straus, W Walter and C A Gross

Genes Dev. 1990, 4:
Access the most recent version at doi:10.1101/gad.4.12a.2202

References
This article cites 39 articles, 21 of which can be accessed free at: http://genesdev.cshlp.org/content/4/12a/2202.full.html#ref-list-1

License

Email Alerting Service
Receive free email alerts when new articles cite this article - sign up in the box at the top right corner of the article or click here.