

# Degradation of carboxy-terminal-tagged cytoplasmic proteins by the *Escherichia coli* protease HflB (FtsH)

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**Proteins with short nonpolar carboxyl termini are unstable in *Escherichia coli*. This proteolytic pathway is used to dispose of polypeptides synthesized from truncated mRNA molecules. Such proteins are tagged with an 11-amino-acid nonpolar destabilizing tail via a mechanism involving the 10Sa (SsrA) stable RNA and then degraded. We show here that the ATP-dependent zinc protease HflB (FtsH) is involved in the degradation of four unstable derivatives of the amino-terminal domain of the  $\lambda$ C1 repressor: three with nonpolar pentapeptide tails (cI104, cI105, cI108) and one with the SsrA tag (cI-SsrA). cI105 and cI-SsrA are also degraded by the ClpP-dependent proteases. Loss of ClpP can be compensated for by overproducing HflB. In an in vitro system, cI108 and cI-SsrA are degraded by HflB in an energy-dependent reaction, indicating that HflB itself recognizes the carboxyl terminus. These results establish a tail-specific pathway for removing abnormal cytoplasmic proteins via the HflB and Clp proteases.**

[Key Words: 10Sa RNA; AAA ATPase family; intracellular proteolysis; tail-specific proteolysis; Clp protease; proteasome 26S]

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Living cells have elaborate mechanisms such as check points and coupling devices to evaluate their physiological state and environment to maintain a harmonious cell cycle. In addition, cell growth requires efficient systems to clean up erroneous metabolites and macromolecules that would otherwise accumulate in the cytoplasm and ultimately arrest growth. Abnormal proteins are a frequent and cumbersome type of cytoplasmic junk, and ingenious proteolytic systems have evolved to recognize and remove them. Abnormal proteins can arise in various ways: from denaturing treatments, from improper folding of newly synthesized polypeptides (often the case for foreign proteins), and from premature termination of transcription or translation (Gottesman 1996; Miller 1996).

Cytoplasmic proteolysis serves several purposes (Gottesman 1996; Herman and D'Ari 1998). It permits rapid regulatory responses to specific signals, it creates protein turnover in times of starvation, and it removes abnormal proteins. The latter housekeeping role for cytoplasmic

proteolysis is a major cell need, and inability to perform it quickly perturbs growth.

Regulatory proteolysis is generally quite specific, each unstable regulator being the preferential substrate of a particular protease, whereas in housekeeping proteolysis, abnormal proteins are often substrates of several proteases (Maurizi et al. 1985). Such lack of specificity is perhaps not surprising, since virtually any protein can become "abnormal" by denaturation, in which case recognition presumably exploits general criteria, shared by most proteases. However, the specific features of proteins that target them for degradation remain largely unknown.

In a systematic search for protease recognition signals in *Escherichia coli*, Parsell et al. (1990) showed that a stable protein can be destabilized by addition of nonpolar carboxy-terminal pentapeptide tags. A natural tagging mechanism of this sort was described recently in *E. coli*, by which a nonpolar amino acid tail is added to peptides translated from truncated mRNA molecules (Keiler et al. 1996), making nonpolar tails an important signal in housekeeping proteolysis. Carboxy-terminal nonpolar tags do not cause protein unfolding (Parsell et al. 1990), so they are presumably recognized directly by proteolytic systems. The periplasmic protease Tsp (or Prc) spe-

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cifically recognizes these tagged polypeptides both in vitro and, for periplasmic proteins, in vivo (Keiler and Sauer 1996), but of course it does not degrade tagged cytoplasmic proteins in vivo (Silber and Sauer 1994). The proteases responsible for their degradation, which have not been identified previously, are the subject of this report.

We examined the ability of the HflB- and ClpP-dependent proteases to degrade *ci* repressor variants with nonpolar carboxy-terminal tails. HflB is the only essential protease in *E. coli*. It is a membrane-anchored protein with its active site in the cytoplasm (Tomoyasu et al. 1993) and is responsible for degradation of the heat shock sigma factor  $\sigma^{32}$  and of the regulatory proteins *ciI* and *ciII* of phage  $\lambda$  (Herman et al. 1995; Tomoyasu et al. 1995; Herman et al. 1997; Shotland et al. 1997); few other substrates are known (Kihara et al. 1995). ClpP is the proteolytic subunit of two different cytoplasmic proteases, ClpXP and ClpAP, each of which has distinct recognition signals. For example, ClpXP degrades the  $\sigma$  factor involved in the stationary phase, whereas ClpAP degrades the chromosomal addiction module MazE (Aizenman et al. 1996; Schweder et al. 1996).

In the present work, we study four carboxy-terminally tagged proteins. The amino-terminal domain of phage  $\lambda$  *ci* repressor, normally a stable cytoplasmic protein, is rapidly degraded when tagged with either a nonpolar pentapeptide tail or the SsrA natural 11-amino-acid carboxy-terminal tag (Parsell et al. 1990; Keiler et al. 1996). We show that these tagged proteins are substrates of HflB- and, in some cases, also of ClpP-dependent proteases, strongly suggesting that HflB- and ClpP-dependent proteases can select substrates by their carboxy-terminal tails.

## Results

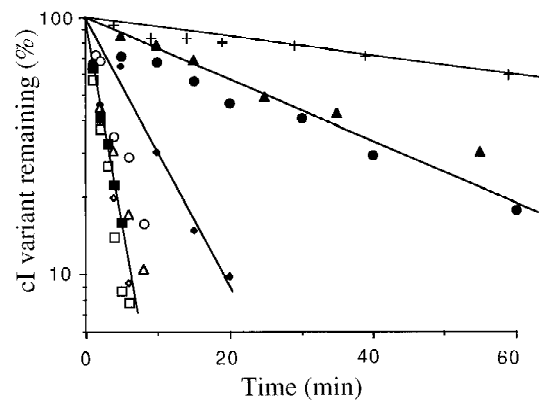
### Tail-specific cytoplasmic degradation in the *hflB1(Ts)* mutant

We first examined the role of HflB in the degradation of three unstable *ci* variants carrying nonpolar pentapeptide tails—*ci104*, *ci105*, and *ci108* (Parsell et al. 1990)—and the *ci* variant carrying the natural nonpolar tail, *ci-SsrA*; the stable variant *ci102* served as control (see Table 1). To determine whether HflB is responsible for the cytoplasmic degradation of these unstable *ci* variants, we carried out pulse chase experiments at 42°C in a ther-

**Table 1.** Carboxy-terminal amino acid sequence of the *ci* variants

<i>ci</i> variant	Carboxyl terminus
<i>ci102</i>	–Arg–Ser–Glu– <u>Tyr</u> –Glu
<i>ci104</i>	–Ile– <u>His</u> –Trp–Val– <u>Thr</u>
<i>ci105</i>	–Trp–Val–Ala–Ala–Ala
<i>ci108</i>	–Ser–Leu–Leu–Trp–Ser
<i>ci-SsrA</i>	–Ala–Ala–Asn–Asp–Glu–Asn– <u>Tyr</u> –Ala–Leu–Ala–Ala

(Boldface type) Nonpolar amino acids; (underline) uncharged polar amino acids; (others) acidic and basic amino acids.



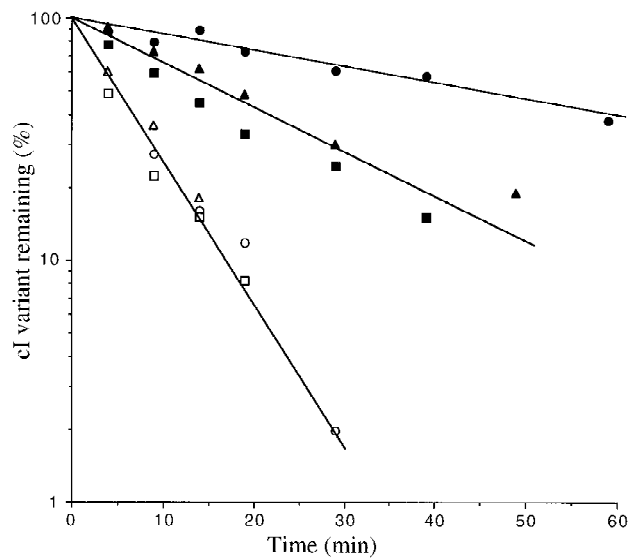
**Figure 1.** Stability of *ci* variants in the *hflB1(Ts)* mutant. The strains used were JM105 (WT) or its *hflB1(Ts)* derivative carrying one of the following plasmids: pDP102 (*ci102*), p104 (*ci104*), p105 (*ci105*), p108 (*ci108*), or p $\lambda$ N-SPT (*ci-SsrA*). (+) Wild-type, *ci102*; ( $\Delta$ ) wild type, *ci108*; ( $\blacktriangle$ ) *hflB1*, *ci108*; ( $\circ$ ) wild type, *ci104*; ( $\bullet$ ) *hflB1*, *ci104*; ( $\diamond$ ) wild type, *ci105*; ( $\blacklozenge$ ) *hflB1*, *ci105*; ( $\square$ ) wild type, *ci-SsrA*; ( $\blacksquare$ ) *hflB1*, *ci-SsrA*. Cultures growing exponentially at 30°C in M63 supplemented with glycerol and ampicillin were shifted to 42°C, induced, pulse labeled, and chased, and the amount of labeled *ci* variant was measured periodically, as described in Materials and Methods.

mosensitive *hflB1(Ts)* mutant carrying a plasmid coding for one or another *ci* variant. In wild-type cells at 42°C, the half-life of all four unstable variants was  $\sim$ 2 min, whereas the stable control, *ci102*, had a half-life of 200 min (Fig. 1). In the *hflB1(Ts)* mutant at 42°C (Fig. 1), two variants, *ci104* and *ci108*, were strongly stabilized (30-min half-lives), one variant, *ci105*, was partially stabilized (7-min half-life), and *ci-SsrA* was not stabilized at all (2-min half-life). These data show that HflB is involved in the degradation of carboxy-terminal-tagged *ci104*, *ci108*, and, to a lesser extent, *ci105*.

### Tail-specific cytoplasmic degradation after HflB depletion

Previous experiments have indicated that the HflB1 mutant protein still has degradative activity toward some of its specific substrates at 42°C (Herman et al. 1997), which could explain the continued instability of *ci105* and *ci-SsrA* in the mutant. Therefore, we examined the stability of these *ci* variants in a different conditional system in which the HflB protein is depleted. The HflB depletion strain carries a plasmid-borne copy of *hflB* under control of the tightly regulated arabinose promoter and a deletion of the chromosomal *hflB* gene. Cells growing in glycerol plus arabinose at 37°C were transferred to glucose medium lacking arabinose to shut off *hflB* expression. After 6 hr in glucose, the half-lives of induced *ci108*, *ci105*, and *ci-SsrA* were 45 min, 18 min, and 16 min, respectively, compared to 6 min in glycerol plus arabinose medium (Fig. 2). These data again show that HflB is involved in the degradation of *ci108* and *ci105* and show further that it is also involved in *ci-SsrA* degradation. However, the fact that *ci105* and *ci-SsrA* are

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**Figure 2.** Stability of cI variants in the  $\Delta hflB3::kan$  mutant. The strains used were JM105 (WT) or its  $\Delta hflB$  derivative, carrying two plasmids, pBHB1 (*paraBAD-hflB*) and p108 (cI108), p105 (cI105), or pAN-SPT (cI-SsrA). (○) Wild-type, cI108; (●)  $\Delta hflB$ , cI108; (△) wild type, cI105; (▲)  $\Delta hflB$ , cI105; (□) wild type, cI-SsrA; (■)  $\Delta hflB$ , cI-SsrA. Cultures were grown to exponential phase at 37°C in M63 supplemented with glycerol, arabinose, ampicillin, and chloramphenicol. HflB protein was depleted by 6 hr incubation in glucose medium lacking arabinose, after which the cultures were pulse labeled and chased and the amount of labeled cI variant was measured periodically, as described in Materials and Methods.

only partially stabilized suggests that these proteins are recognized by another protease as well.

#### Role of ClpP in tail-specific cytoplasmic degradation

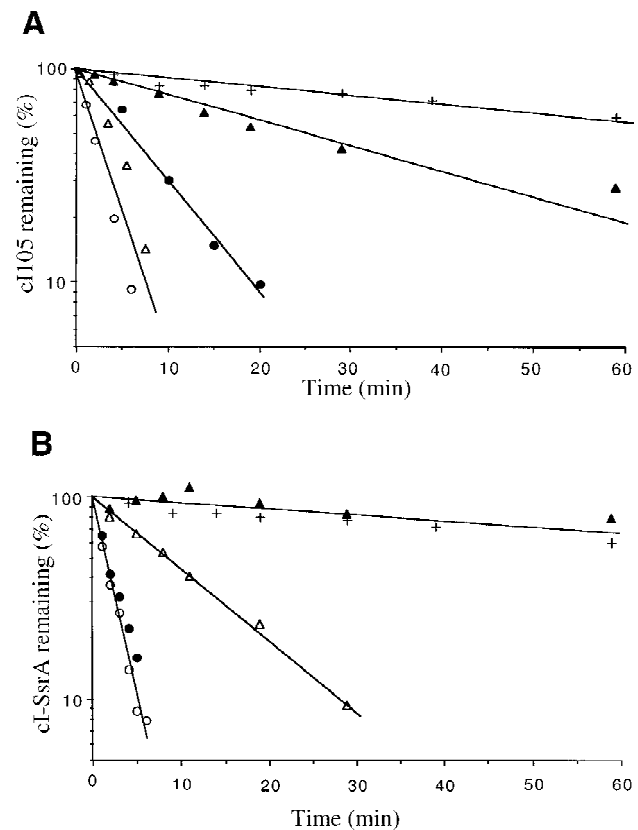
We tested the effect of a *clpP* mutation (which eliminates the ClpXP and ClpAP proteases) in combination with the *hflB1*(Ts) mutation on cI105 and cI-SsrA degradation. Both cI variants were strikingly stabilized in the *hflB1*(Ts) *clpP*::Tn9 double mutant at 42°C, with half-lives of 31 and 120 min for cI105 and cI-SsrA, respectively, compared to ~2 min in wild type (Fig. 3). Strains mutant for each protease separately exhibited less stabilization than that observed in the double mutant. In the single mutants, the cI105 protein (Fig. 3A) was slightly stabilized in *clpP*::Tn9 (3 min) and in *hflB1*(Ts) (6 min), and the cI-SsrA protein (Fig. 3B) was partially stabilized in *clpP*::Tn9 (9 min) but not at all in *hflB1*(Ts) (2 min).

These data show clearly that the degradation of cI105 and cI-SsrA involves both ClpP and HflB. Stabilization of cI-SsrA in a *clpP* mutant has also been observed by Gottesman et al. (1998), who further showed that a related molecule (with a small insert between the cI and SsrA moieties) can be degraded in vitro by both ClpXP and ClpAP. It is interesting to note that cI105 is not totally stabilized in the *hflB1*(Ts) *clpP*::Tn9 double mu-

tant, suggesting that yet a third protease may be involved in its degradation.

#### Overexpression of HflB compensates for the proteolysis defect of the *clpP* mutant

As shown above, cI-SsrA is recognized by the ClpP-dependent proteases and *hflB* protease. To investigate their respective roles, we measured the half-life of cI-SsrA in the *clpP*::Tn9 mutant at 37°C so that we could compare it to that in HflB-depleted cells. In *clpP*::Tn9 cells there was a 5-fold increase in stability (30 vs. 6 min in wild type), compared to the 2.5-fold increase in HflB-depleted cells (16 vs. 6 min in wild type). We have previously shown that the HflB concentration is a limiting factor in the degradation of its substrates  $\sigma^{32}$  and  $\lambda$ CII (Herman et al. 1995, 1997). We therefore asked whether overproduction of HflB could overcome the proteolytic defect of the *clpP*::Tn9 mutant. We introduced into the *clpP*::Tn9 strain a plasmid in which the expression of *hflB* is under control of *plac*. In the resulting strain, the addition of



**Figure 3.** Degradation of cI105 and cI-SsrA by both HflB and Clp. The strains used were JM105 (WT) (○), its *hflB1* derivative (●), its *clpP*::Tn9 derivative (△), and its *hflB1 clpP*::Tn9 derivative (▲). (A) These strains carry plasmid p105 (cI105); (B) they carry plasmid pAN-SPT (cI-SsrA); a wild-type control (+) carrying plasmid pDP102 (cI102), coding for a stable cI variant, is shown in A and B. Cultures were grown, pulse labeled, chased, and analyzed as in the experiments shown in Fig. 1.

Tail-specific degradation of HflB in *E. coli*

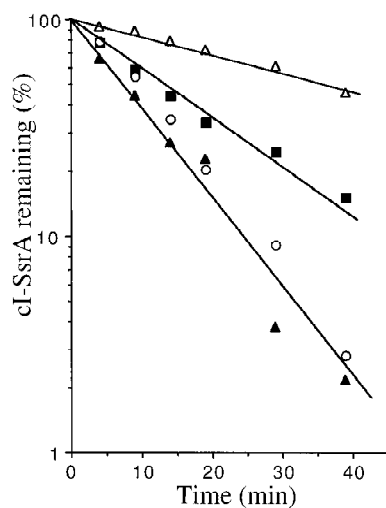
IPTG causes overexpression of both HflB and *cl*-SsrA. Under these conditions, the half-life of *cl*-SsrA was reduced to 7 min, compared to 28 min in the *clpP*::Tn9 control strain in which HflB was not overproduced (Fig. 4). The half-lives reported here are longer than those shown in Fig. 3 because of the lower temperature (see below).

These data show that excess HflB is able to replace the ClpP-dependent proteolytic pathway for *cl*-SsrA degradation. If the role of HflB is direct, the results suggest that HflB can fully recognize the SsrA tag and that the concentration of SsrA-specific proteases determines the half-life of SsrA-tagged proteins.

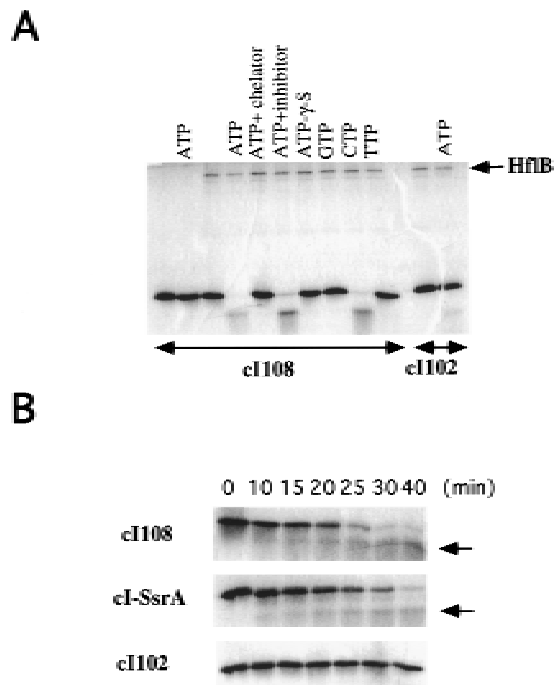
*In vitro* degradation of *cI108* and *cI*-SsrA

To determine whether the action of HflB on unstable variants is direct, we purified HflB-His<sub>6</sub>-Myc from the membrane fractions of an overproducing strain (Kihara et al. 1996). The purification steps included a nickel column followed by a monoQ column. His<sub>6</sub>-*cI108*, His<sub>6</sub>-*cI*-SsrA, and His<sub>6</sub>-*cI102* were purified in a single step, using a nickel affinity column; the preparations were free of ATP-dependent proteases since addition of ATP to the purified *cl* derivatives did not stimulate their degradation.

With these preparations, we investigated the ability of HflB to degrade tail-tagged  $\lambda$  *cl* repressor molecules in vitro. As shown in Figure 5A, purified HflB degraded *cI108* in the presence of ATP or CTP, under protease-limiting conditions (the molar ratio of HflB to *cI108* was



**Figure 4.** Excess HflB compensates for loss of ClpP in the degradation of *cl*-SsrA. The strains used were JM105 (○), its  $\Delta hflB3::kan$  derivative (■), and its *clpP*::Tn9 derivative carrying either the HflB-overproducing plasmid pULB6234 (▲) or the control plasmid pRK7813 (△). All strains carried plasmid pAN-SPT, coding for *cl*-SsrA. Cultures were grown to exponential phase at 37°C in M63 supplemented with glycerol, ampicillin, and chloramphenicol. The cultures were pulse labeled and chased and the amount of labeled *cl* variant was measured periodically, as described in Materials and Methods.



**Figure 5.** Degradation of *cl*-SsrA and *cI108* by HflB in vitro. His-tagged HflB, *cl*-SsrA, *cI108*, and *cI102* proteins were purified as described in Materials and Methods. (A) The indicated reactants were mixed in P buffer for 30 min at 37°C, and the reaction mixture was electrophoresed. Lanes 1–10 contained the unstable variant *cI108*; lanes 11 and 12 contained the stable variant *cI102*; HflB was present everywhere except in the controls shown in lanes 1 and 2. The other reactants in the different lanes were (lane 1) none; (lane 2) ATP; (lane 3) none; (lane 4) ATP; (lane 5) ATP + 1-10 phenanthroline; (lane 6) ATP + pefabloc; (lane 7) ATP- $\gamma$ -S; (lane 8) GTP; (lane 9) CTP; (lane 10) TTP; (lane 11) none; and (lane 12) ATP. (B) Kinetic analysis of reaction mixtures containing HflB, ATP and *cl*-SsrA, *cI108*, or *cI102*. Arrows indicate a major breakdown peptide appearing in the first two mixtures. Concentrations of the various reactants are indicated in Materials and Methods.

1:45). A major product of degradation, about half the protein (5 kD), was visible, suggesting that HflB acts as an endoprotease. In contrast, the stable variant *cI102* was not degraded under the same conditions, in agreement with the in vivo results and suggesting that HflB is specific for *cl* molecules with a nonpolar tail, such as *cI108*. No degradation was observed when ATP was replaced with GTP or TTP. The degradation is HflB-dependent since addition of a zinc chelator inhibited the reaction whereas a serine protease inhibitor did not. The nonhydrolyzable ATP analog ATP- $\gamma$ -S did not allow *cI108* degradation, suggesting that this proteolysis requires ATP hydrolysis and thus is energy dependent. Finally, we compared the kinetics of degradation of *cI108* and *cl*-SsrA; with a 45-fold molar excess of substrate at 37°C, the in vitro half-life of *cI108* was 10 min, whereas that of *cl*-SsrA was ~15 min (Fig. 5B).

These results confirm that the unstable *cl* variants with nonpolar tails are substrates of the protease HflB,

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suggesting that the nonpolar tail is directly recognized by HflB.

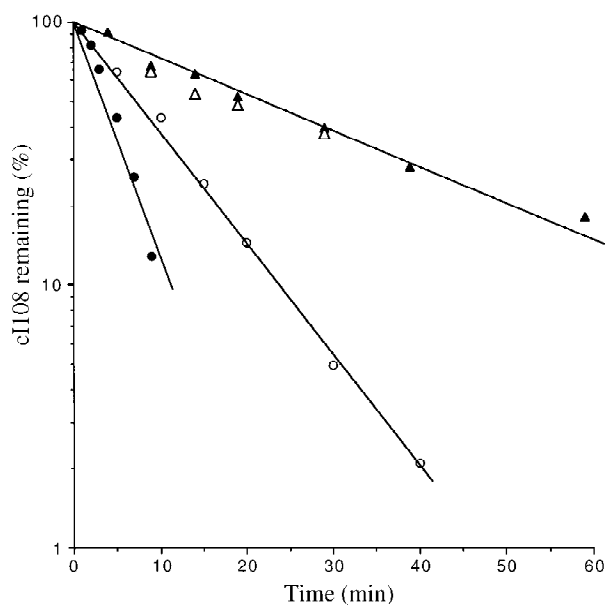
#### Temperature dependence of HflB degradation

In the course of the above work, we noticed that the half-lives of the different *cI* variants varied with the temperature. To characterize this phenomenon quantitatively, we measured the half-life of *cI108* in vivo as a function of temperature. As shown in Figure 6, the half-life of *cI108* was ~20 min at 30°C and 34°C, 6 min at 37°C, 3 min at 40°C and, as already shown, 2 min at 42°C.

These data demonstrate that proteolysis of *cI108* increases with temperature. The effect is larger than the known heat-shock induction of the protease HflB (Herman et al. 1995); this could reflect temperature-dependent changes in substrate conformation or in HflB activity.

#### Discussion

Removing abnormal proteins from the cytoplasm is a vital housekeeping function (Miller 1996). In bacteria, mRNA is highly unstable; thus, truncated messenger molecules may be a significant source of such abnormal proteins. In *E. coli*, peptides arising from such mRNA molecules have recently been shown to be tagged by a process in which the stable 10Sa RNA encoded by *ssrA* is used to add an 11-amino-acid carboxy-terminal sequence with a nonpolar tail and release them from the ribosome;

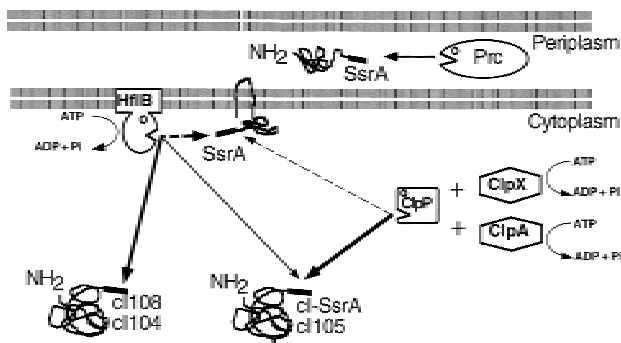


**Figure 6.** Temperature dependence of *cI108* degradation. Strain JM105 carrying plasmid p108, coding for *cI108*, was grown to exponential phase at the indicated temperatures [ (Δ) 30°C; (▲) 34°C; (○) 37°C; (●) 40°C] in M63 supplemented with glycerol and ampicillin, then pulse labeled and chased at that temperature and analyzed as in the experiments shown in Fig. 1.

these SsrA-tagged truncated peptides are targeted for degradation (Keiler et al. 1996), and conservation of the nonpolar tail suggests that this system is common to all bacteria (Williams and Bartel 1996). In a systematic screen for protease recognition signals, it was found that proteins could be destabilized by the addition of similar nonpolar pentapeptide tails without changing their thermal stability (Parsell et al. 1990). The periplasmic protease Tsp (or Prc) recognizes proteins with nonpolar tails in vitro and in vivo as well for periplasmic proteins (Silber et al. 1992; Keiler and Sauer 1996). We show here that cytoplasmic proteins destabilized by such carboxy-terminal tags become substrates of the ATP-dependent protease HflB, an essential, membrane-anchored zinc protease. In some cases the ClpP-dependent proteases are also involved in their degradation. Using purified HflB in vitro, the half-life of proteins tagged with nonpolar tails, in the presence of excess substrate was similar to the in vivo half-life, strongly suggesting that the carboxy-terminal nonpolar sequence is recognized by the protease itself.

All cytoplasmic proteases seem to be polyvalent, with both regulatory and housekeeping functions. Proteolysis involved in regulation is often specific, for example, the heat shock specific sigma factor  $\sigma^{32}$  is degraded primarily by the protease HflB (Herman et al. 1995), whereas the stationary phase sigma factor  $\sigma^S$  is degraded primarily by ClpXP (Schweder et al. 1996). Housekeeping proteolysis, on the contrary, does not usually exhibit such sharp specificity, and abnormal proteins are often substrates of several proteases. For example, puromycin-truncated proteins are degraded in *E. coli* by both Lon and HslUV (Maurizi et al. 1985; Straus et al. 1988; Missiakas et al. 1996). The tagged carboxy-terminal *cI* variants provide an experimental approach to study how proteases can have both specific regulatory functions and nonspecific housekeeping functions. Two of the *cI* variants studied here are degraded exclusively by HflB. Two others exhibit little specificity; they are substrates of HflB, Tsp, and ClpP-dependent proteases (see Fig. 7). Since these variants differ only in their carboxyl terminus, the information in this small tail would seem to be sufficient to confer specificity to a single or several proteases. It is interesting to note that universal tags (*cI105*, *cI-SsrA*) are composed of only nonpolar amino acids (usually buried inside the protein), whereas specific tags are composed of polar and nonpolar amino acids (see Table 1). Using artificial tail-tagged *cI* variants, it should be possible to define the molecular mechanism by which proteases recognize both regulatory and housekeeping substrates.

Among the proteases able to degrade *cI-SsrA* in the cytoplasmic compartment, the ClpXP and ClpAP proteases are collectively more important than HflB, as we observed greater stabilization by loss of ClpP (5-fold) than by loss of *hflB* (2.5-fold). The importance of ClpP-dependent proteases in *cI-SsrA* degradation has also been established by Gottesman et al. (1998). Since HflB is in the membrane, its paramount role may be to degrade abnormal membrane proteins, SsrA-tagged or not, that have their carboxyl termini in the cytoplasm (Fig. 7).



**Figure 7.** Proteolysis of carboxy-terminal-tagged proteins in *E. coli*. The degradation of carboxy-terminal-tagged polypeptides is schematized. The three cytoplasmic proteases, all ATP-dependent, are connected to their substrates by arrows. Thicker arrows indicate higher affinity; broken arrows indicate hypothetical reactions.

This idea is consistent with the fact that HflB plays a further role in membrane quality control by degrading uncomplexed membrane proteins (Kihara et al. 1995). The SsrA tag is thus a signal for protease recognition in all cell compartments—the periplasm (Tsp), the membrane (HflB), and the cytoplasm (ClpXP, ClpAP, and HflB), permitting the rapid elimination of tagged proteins, wherever they may be, as one would expect for a housekeeping function.

HflB, ClpXP, and ClpAP all carry out tail-specific cytoplasmic proteolysis, and all can degrade SsrA-tagged substrates *in vitro* (this work; Gottesman et al. 1998). These proteases recognize nonpolar tails, which presumably provide an anchoring site to which they bind. It has recently been shown that ClpX interacts with the SsrA tail through a PDZ-like domain (Levchenko et al. 1997). PDZ domains, defined in eukaryotes, are modules of ~90 residues that directly bind the carboxy-terminal ends of targeted proteins (Saras and Heldin 1996). Sequence analysis of HflB reveals homology with the PDZ-like domain of Clp in its well conserved “AAA” module (data not shown). The latter, a 220–250-amino-acid sequence, including an ATPase motif, defines a large family of >200 proteins (e.g., regulatory subunits of the 26S proteasome) (Confalonieri and Duguet 1995; Beyer 1997).

HflB, ClpXP, and ClpAP are tail-specific proteases. HslU, which belongs to the Clp family, also possesses a putative tail-specific binding domain (Levchenko et al. 1997). The Lon protease, too, seems to bind to tails (Higashitani et al. 1997; Sen and Rothfield 1998). Tail-specific recognition thus seems to be quite general among *E. coli* cytoplasmic proteases. The resemblance of HflB to a proteasome regulatory subunit suggests that tail-specific recognition may occur in higher organisms as well, and indeed it has been reported in several eukaryotic systems. For example, the S4 proteasome subunit in human cells binds to the carboxy-terminal part of the papillomavirus oncoprotein E7, apparently as part of a mechanism by which E7 delivers the retinoblastoma tumor suppressor protein to the proteasome (Berezutskaya and

Bagchi 1997). Similarly, mammalian ornithine decarboxylase, normally degraded by the 26S proteasome (Murakami et al. 1992), was stabilized when the 37 carboxy-terminal residues were removed (Ghoda et al. 1989), and the degradation of mouse thymidine kinase also depends on its carboxy-terminal residues (Sutterluety et al. 1996). Tail-specific recognition may be a major entry into proteolytic pathways in all organisms.

## Materials and methods

### Bacterial strains

The strains used in this work are all *E. coli* K12 derivatives. There are isogenic strains derived from JM105, which carries a *lac<sup>r</sup>* allele causing Lac repressor overproduction (Sambrook et al. 1989). HflB-His<sub>6</sub>-myc was purified from strain AK1181 (Kihara et al. 1996).

P1 *vir*-mediated transduction was carried out as described by Miller (1992). Plasmid preparations, transformation, cloning, and ligation were carried out as described by Sambrook et al. (1989). The  $\Delta hflB3::kan$  ( $\DeltaftsH3::kan$ ) allele (Akiyama and Shirai 1994) was introduced by transduction to kanamycin resistance in the presence of 70  $\mu$ M L-arabinose into strains carrying plasmid pBHB1. Other protease mutations were introduced by P1 transduction, selecting for an associated antibiotic resistance marker: *hflB1*(Ts) *zgj-203::Tn10*, tetracycline resistance, from strain SL93 (Herman et al. 1993), and *clpP::Tn9*, chloramphenicol resistance, from strain SL246 (Herman et al. 1995).

### Plasmids

Plasmids pDP102, p104, p105, p108 (Parsell et al. 1990), and pAN-SPT (Keiler et al. 1996) carry the different *cl* variant genes under control of the IPTG-inducible *tac* promoter of an ampicillin-resistant pBR322 derivative.

Plasmid pHis-cl108 codes for the cl108 protein fused to a histidine tag at its amino terminus; it was constructed by amplifying plasmid pcl108 with primers CINH2 (5'-GGCGGATCATGAGCACAAAAAGAAACC-3') and CICOHHIII (5'-GCCAAGCTTGCAATTTAACTGTGATAAACT-3'). The resulting product was cut with *Bam*HI and *Hind*III and ligated to the pQE8 (Qiagen) *Bam*HI-*Hind*III backbone fragment. The expected structure of the construct was verified by DNA sequencing.

Plasmids pHis-cl102 and pHis-cl-SsrA overexpress the histidine-tagged cl102 and cl-SsrA, respectively. To construct these plasmids, we replaced the *Nsil*-*Nde*I fragment of pHis-cl108 with the short *Nsil*-*Nde*I fragment of pDP102 or pAN-SPT. The proteolytic susceptibility of each His-tagged variant was tested by pulse chase in a wild-type strain and shown to be identical to that of the untagged variant.

Plasmid pULB6234 (Herman et al. 1993) carries the intact *hflB* gene cloned downstream of the *lac* promoter in a low copy number, tetracycline-resistant plasmid vector, pRK7813 (Jones and Guttererson 1987).

Plasmid pBHB1 (Herman et al. 1997) carries the intact *hflB* gene cloned downstream of the *ara* promoter in a low copy number, chloramphenicol-resistant pACYC184 derivative, pBAD33 (Guzman et al. 1995).

Plasmid pSTD113 (Kihara et al. 1996) carries the *hflB*-his<sub>6</sub>-myc gene fusion under control of the IPTG-inducible *lac* promoter of an ampicillin-resistant pBR322 derivative.

Herman et al.

### Culture conditions

Cells were grown in M63 supplemented with 0.4% glycerol and 1  $\mu\text{g/ml}$  thiamine or in LB broth (Miller 1992). For HflB depletion, cells were grown in M63 glucose (0.4%) plus 19 amino acids (lacking methionine) at 100  $\mu\text{g/ml}$  each. When necessary, antibiotics were added at the following concentrations: 50  $\mu\text{g/ml}$  ampicillin, 15  $\mu\text{g/ml}$  chloramphenicol; 40  $\mu\text{g/ml}$  kanamycin; and 10  $\mu\text{g/ml}$  tetracycline.

### Measurement of *cI* variant stability

All strains carried a *lacI<sup>q</sup>* allele and a plasmid with the *cI* variant gene under control of the *tac* (IPTG-inducible) promoter. The protocol was essentially that described by Parsell et al. (1990). Cells were grown to exponential phase at 30°C in glycerol minimal medium supplemented with antibiotics, then shifted to 42°C ( $t = 36$  min), induced with 1 mM IPTG ( $t = 16$  min), pulse-labeled ( $t = 1$  min) with [<sup>35</sup>S]methionine + [<sup>35</sup>S]cysteine (38.3 TBq/mole, 0.22 MBq/ml, ICN Biomedicals), and chased ( $t = 0$ ) with 100  $\mu\text{g/ml}$  each cold methionine plus cysteine. For pulse-chase experiments done at 37°C, cells were also pregrown at 37°C. Samples were electrophoresed on 15% SDS-polyacrylamide Tris-tricine gels. Radioactivity in the *cI* variant band was revealed with a Molecular Dynamics PhosphorImager and quantified with the ImageQuant 1.33 program (Molecular Dynamics). Half-lives were determined as the slope of the least squares best fit through the points plotted on a semilog graph. All experiments were repeated two to four times; there were no significant variations in the results.

### Protein purification

Purification of HflB-his<sub>6</sub>-myc was essentially as described by Kihara et al. (1996), with the following modifications. Cells of AK1272 carrying pSTD113 were grown at 37°C to midexponential phase in 1 L of TB with ampicillin (50 mg/liter). IPTG (final concentration 1 mM) was added 3 hr before cells were harvested by centrifugation. The DEAE-Sepharose fast flow column was replaced by a monoQ fast flow column.

The His<sub>6</sub>-cI102, His<sub>6</sub>-cI108, and His<sub>6</sub>-cI-SsrA proteins were purified by denaturing nickel NTA chromatography according to the Qiagen protocol.

### In vitro degradation of *cI* variants by HflB-his<sub>6</sub>-myc

Purified *cI* variant preparations (30  $\mu\text{M}$ ) were incubated with HflB (0.67  $\mu\text{M}$ ) in buffer P (50 mM Tris-HCl at pH 7.6, 5 mM MgCl<sub>2</sub>, 25  $\mu\text{M}$  zinc acetate, 1 mM DTT, 0.1% NP-40). The mixture was incubated at 37°C and stopped with 2  $\times$  SDS sample buffer (Sambrook et al. 1989). ATP, CTP, GTP, TTP, and ATP- $\gamma$ -S were added at a final concentration of 5 mM. When indicated, pefabloc or 1-10 phenanthroline was added at a final concentration of 10 mM. Samples were electrophoresed on 15% sodium SDS-polyacrylamide Tris-glycine gels and revealed by Coomassie stain. Gels were digitalized with an Apple scanner.

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