Nucleotide Triphosphates Inhibit the Degradation of Unfolded Proteins by HsIV Peptidase

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Escherichia coli HsIVU is an ATP-dependent protease consisting of two heat shock proteins, the HslU ATPase and HslV peptidase. In the reconstituted enzyme, HslU stimulates the proteolytic activity of HslV by one to two orders of magnitude, while HslV increases the rate of ATP hydrolysis by HslU several-fold. Here we show that HslV alone can efficiently degrade certain unfolded proteins, such as unfolded lactalbumin and lysozyme prepared by complete reduction of disulfide bonds, but not their native forms. Furthermore, HslV alone cleaved a lactalbumin fragment sandwiched by two thioredoxin molecules, indicating that it can hydrolyze the internal peptide bonds of lactalbumin. Surprisingly, ATP inhibited the degradation of unfolded proteins by HslV. This inhibitory effect of ATP was markedly diminished by substitution of the Arg86 residue located in the apical pore of HslV with Gly, suggesting that interaction of ATP with the Arg residue blocks access of unfolded proteins to the proteolytic chamber of HslV. These results suggest that uncomplexed HslV is inactive under normal conditions, but may can degrade unfolded proteins when the ATP level is low, as it is during carbon starvation.

Keywords: ATP-dependent Protease; ATPase; HslU; HslV; Lactalbumin; Unfolded Proteins.

Introduction

ATP-dependent proteolysis plays an essential role in eliminating proteins with aberrant structures and in controlling the levels of key regulatory proteins (Goldberg, 1992; Gottesman and Maurizi, 1992). In bacteria, these functions are carried out by ATP-dependent proteases, such as Lon (protease La), ClpAP (protease Ti), ClpXP, HslVU (ClpQY), and FtsH (Chung, 1993; 1997; Gottesman, 1996; 2003; Maurizi, 1992; Park et al., 2006). HsIVU is the product of the hslVU operon, which encodes two heat shock proteins, the HslV peptidase and the HslU ATPase (Chaung et al., 1993). HslV forms a dodecamer of two back-to-back stacked hexameric rings (Bochtler et al., 1997) while HslU is a hexameric ring that binds to either one or both ends of HslV to form the HslVU complex (Bochtler et al., 2000; Wang et al., 2001a). In the HslVU complex, the central pores of HslU and HslV are aligned and the proteolytic active sites are sequestered in the internal chamber of HslV, with access to this chamber restricted to small axial pores.

HslV by itself is a weak peptidase that slowly degrades some hydrophobic peptides, such as N-carbobenzoxy-Gly-Gly-Leu-7-amido-4-methylcoumarin (Z-GGL-AMC), and certain unfolded proteins, like casein and insulin B-chain (Rohrwild *et al.*, 1996; Yoo *et al.*, 1996). In the presence of ATP, HslU markedly stimulates the peptide- and protein-degrading activity of HslV (up to about 100-fold) by forming the HslVU complex. Interestingly, ATP γ S, a nonhydrolyzable ATP analog, supports the hydrolysis of the insulin B-chain and Z-GGL-AMC by HslVU even more effectively than ATP. However, ATP γ S and other ATP analogs cannot support the degradation of native, folded protein substrates, such as SulA, an inhibitor of cell division (Seong *et al.*, 1999). Thus, ATP hydrolysis by HslU ap-

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pears to play an essential role in unfolding native protein substrates for subsequent translocation into the inner proteolytic chamber of the HsIV dodecamer. In addition, HsIU alone can function as a molecular chaperone preventing aggregation of SuIA, which has an intrinsic tendency to form aggregates under both *in vitro* and *in vivo* conditions, although ATP hydrolysis appears not to be essential for its chaperone function (Seong *et al.*, 2000).

 α -Lactalbumin (LA), which has four disulfide-bonds and binds a single Ca²⁺ ion, is one of the model proteins studied most extensively to clarify mechanisms of protein stabilization, folding, and unfolding. Under a variety of mild denaturing conditions, such as dissociation of Ca²⁺ or partial reduction of disulfide bonds, LA adopts a partially structured conformation termed a 'molten globule' that is intermediate between native and unfolded states (Acharya *et al.*, 1991; Hiraoka *et al.*, 1980). Fully reduced LA no longer has a native conformation with little or no detectable secondary structure.

In an attempt to define the degree of unfolding required for degradation by HsIVU, we used LA as a model protein in various unfolded states as substrate. In the present study, we found that HsIV can by itself degrade unfolded LA (u-LA), but not the native form or partially unfolded intermediates. More surprisingly, ATP was found to inhibit the activity of HsIV by interacting with the basic apical pore region of the HsIV dodecamer. Possible roles of ATP and uncomplexed HsIV in the degradation of unfolded proteins are discussed.

Materials and Methods

Materials Calcium-depleted bovine α -lactalbumin (LA), chicken egg white lysozyme, and bovine pancreatic RNase A were purchased from Sigma. Z-GGL-AMC was obtained from BACHEM Feinchemikalien AG (Switzerland). A peptide corresponding to the C-terminal ten amino acids of HsIU (NH₂-ADEDLSRFIL-COOH) was synthesized by AnyGen, Korea. Plasmid pBS/HsIV that expresses HsIV only was constructed as described (Yoo *et al.*, 1997a).

HslV and HslU were purified as described previously (Yoo *et al.*, 1997b). Partially and fully reduced forms of LA were prepared as described (Ewbank and Creighton, 1993). Fully reduced forms of lysozyme and RNase A (10 mg/ml) were prepared by treatment with 20 mM DTT in the presence of 8 M urea. Protein concentrations were estimated according to Bradford using bovine serum albumin as standard (Bradford, 1976).

Assay of proteolysis Degradation of proteins was assayed as described previously (Seong *et al.*, 2002). Reaction mixtures (0.1 ml) contained protein substrates (7 nM) and HsIV (50 nM) in 25 mM Tris-HCl (pH 8), 2 mM DTT, and 0.5 mM EDTA in the absence or presence of 1 mM adenine nucleotides. To assay degradation of the reduced forms of proteins, 2 mM DTT was

added to the reaction mixtures. After incubation at 37° C, the reactions were stopped by adding 30 µl of 0.75 M Tris-HCl (pH 6.8) containing 7.5% (w/v) SDS and 10% (v/v) 2-mercaptoethanol. The mixtures were then subjected to Tricine-SDS-PAGE as described (Schagger and Jagow, 1987). After staining the gels with Coomassie blue R-250, densitometry was performed to estimate the amounts of substrate remaining using a LAS-3000 and MultiGauge software (FUJIFILM, Japan), with 5-point standard curve calibration. Peptide hydrolysis was assayed by incubating reaction mixtures (0.1 ml) containing Z-GGL-AMC (0.1 mM) and 20 nM HslV in 25 mM Tris-HCl (pH 8), 1 mM DTT, and 0.5 mM EDTA. Release of AMC was measured by continuous (2-min interval) monitoring of fluorescence using a FLUOstar OPTIMA fluorometer (BMG LABTECH, Germany) (Park *et al.*, 2006).

Purification of TLT A cDNA encoding a bovine LA fragment (amino acids 47–111) flanked by two thioredoxin molecules (TLT) was cloned into pET32a, and transformed into strain BL21(DE3)-*hslVU*. The cells were grown in Luria broth to an OD₆₀₀ of 0.6 and induced by treatment with IPTG. TLT was then purified using a Ni²⁺-NTA column according to the manufacturer's instructions.

Results

HsIV by itself degrades unfolded LA In an attempt to define the degree of unfolding required for degradation by HslVU, we used lactalbumin (LA) in various unfolded states as a model substrate. Surprisingly, HslV alone (i.e., in the absence of HslU and ATP) degraded fully unfolded LA (u-LA) (Fig. 1A, left panel), but not native LA or partially unfolded intermediates (data not shown). To test whether HslV degraded other unfolded proteins, lysozyme and RNase A, which possess four disulfide bonds like LA, were fully unfolded by reduction of all the disulfide bonds. We also tested whether HslV hydrolyzed casein, a natively unfolded protein (Holt and Sawyer, 1988; Syme et al., 2002). HslV degraded u-lysozyme nearly as well as u-LA, but cleaved α-casein at a much slower rate (Fig. 1B, left panel). In contrast, it had no activity against u-RNase A. These result indicate that HslV alone is capable of degrading some unfolded proteins, though at different rates.

The C-terminal tails of HslU play a critical role in the interaction with, and activation of, HslV peptidase (Seong *et al.*, 2002). Moreover, a synthetic tail peptide of 10 amino acids (referred to as the C10-peptide) substituted for HslU in promoting HslV-mediated hydrolysis of casein and the insulin B-chain as well as of Z-GGL-AMC. In addition, X-ray crystallographic studies of the structure of the HslVU complex have shown that the highly conserved HslU C-terminus is inserted at the HslV-HslV sub-unit interface when ATP is bound, but otherwise is buried



Fig. 1. Degradation of various unfolded proteins by HslV. A. HslV (50 nM) was incubated with u-LA (7 μ M) at 37°C in the absence (left panel) or presence of 60 μ M C10-peptide (right panel). After incubation, samples were subjected to SDS-PAGE followed by staining with Coomassie blue R-250. **B.** HslV was incubated as above but with various unfolded proteins (7 μ M) in the absence (left panel) or presence of 60 μ M C10-peptide (right panel). The samples were then subjected to SDS-PAGE followed by staining with Coomassie blue R-250. The protein bands were scanned with a densitometer to estimate the amounts of remaining substrate.

at the HslU-HslU subunit interface (Sousa *et al.*, 2000; Wang *et al.*, 2001b). Therefore, we have suggested that insertion of the C10-peptide into pockets at the HslV-HslV interface causes a widening of the central pore of HslV peptidase that permits access of peptides or unfolded polypeptide substrates into the proteolytic chamber, thus facilitating their degradation (Seong *et al.*, 2002). As expected, incubation of HslV with C10-peptide also markedly increased the rate of hydrolysis of u-LA, u-lysozyme, and α -casein, but not that of u-RNase A (Figs. 1A and 1B, right panels). Thus, it seems clear that some unfolded proteins have access to the inner chamber of the HslV dodecamer and are thus degraded without the aid of the HslU ATPase.

HsIV on its own cleaves internal peptide bonds of LA The diameter of the central pore of HsIV is so small that only a single strand of polypeptide can be threaded through it (Bochtler *et al.*, 1997; Wang *et al.*, 1998). This finding is in accord with our observation that HsIV alone can degrade fully unfolded LA but not nascent or partially unfolded intermediates. To confirm this, we constructed a hybrid protein, with a fragment of LA (amino acids 47– 111) sandwiched by two thioredoxin molecules (Fig. 2A). HsIV alone could also degrade this hybrid protein (Fig.



Fig. 2. Cleavage of internal peptide bonds in the LA fragment by HslV. **A.** A hybrid protein comprising two thioredoxin molecules flanked by the amino acid sequence of LA (47–111) was purified as described in **Materials and Methods**. The sizes of the fusion protein (TLT) and the LA fragment (Δ LA) are shown. **B.** HslV (50 nM) was incubated with TLT (1 μ M) at 37°C for the indicated periods in the absence (left panel) or presence of 60 μ M C10-peptide (right panel). After incubation, the samples were subjected to SDS-PAGE followed by staining with Coomassie blue R-250. Note that the incubations were carried out for shorter times when the C10-peptide was present.

2B, left panel) and its proteolysis was markedly stimulated by C10-peptide (right panel). However, thioredoxin was not degraded by HslV (data not shown).

ATP inhibits HslV-mediated degradation of u-LA and Z-GGL-AMC To determine whether HslU in the presence of ATP facilitates the HslV-mediated degradation of u-LA, HslV was incubated with HslU, ATP, or both. In fact HslU in the presence of ATP inhibited the degradation of u-LA by HslV (Fig. 3A) while HslU in the absence of ATP had little or no effect on HslV activity on u-LA. To confirm the inhibitory effect of ATP, HslV was incubated with u-LA in the presence of increasing concentration of ATP. Figure 3B shows that ATP inhibited HslV-mediated degradation of u-LA in a concentration-dependent fashion. Similar inhibitory effects were observed with α -casein and u-lysozyme as substrates (data not shown). We conclude that ATP, not HslU, is responsible for the inhibition of HslV. We next examined whether other adenine nucleotides also inhibited HslV-mediated hydrolysis of u-LA and Z-GGL-AMC. While AMP had little or no effect, ADP did inhibit HslV activity but only about 30% as well as ATP (Fig. 3C). Similar results were obtained for peptide hydrolysis (Fig. 3D). Inorganic pyrophosphate inhibited HslV-mediated degradation of u-LA nearly as well as ADP, whereas inorganic phosphate had little or no effect. In addition, other nucleotide triphosphates inhibited HslV activity to a similar extent as ATP (data not shown). The inhibitory effects of nucleotide triphosphates was not re-



Fig. 3. Effects of adenine nucleotides and HslU on the hydrolysis of u-LA and Z-GGL-AMC by HslV. **A.** HslV (50 nM) was incubated with u-LA (7 μ M) at 37°C for increasing periods in the presence of HslU (10 nM), ATP (1 mM), or both. The letters V and U denotes HslV and HslU, respectively. **B.** HslV (50 nM) was incubated with u-LA (7 μ M) at 37°C for 1 h in the presence of increasing concentrations of ATP. **C.** HslV (50 nM) was incubated with u-LA (7 μ M) at 37°C for increasing periods in the presence of the indicated adenine nucleotides (1 mM). **D.** HslV (20 nM) was incubated with 0.1 mM Z-GGL-AMC and 20 μ M C10-peptide at 37°C for increasing periods in the presence of the indicated adenine nucleotides (1 mM). Peptide hydrolysis was then assayed as described in **Materials and Methods**.

lieved by high concentrations of NaCl (*e.g.*, 0.2 M), suggesting that the inhibition is not due to a simple charge effect.

Effect of the R86G mutation on HslV activity Structural analysis has shown that basic amino acids are clustered in the apical pore region of the HslV dodecamer (Fig. 4A). Moreover, database analysis revealed that the basic pore motif (amino acids 86-90) is highly conserved (Fig. 4B) and that more than 95% of the HslVs in 93 reference sequences have either Arg or Lys at position 86 (Fig. 4C). To see whether the inhibitory effect of ATP on the HslVmediated degradation of u-LA and Z-GGL-AMC is due to the interaction of ATP with Arg86 we replaced the Arg residue of E. coli HslV with Gly by site-directed mutagenesis. The R86G mutation markedly reduced the ability of HslV to degrade u-LA and other unfolded proteins (data not shown). In contrast, it hydrolyzed Z-GGL-AMC nearly as well as wild-type HslV (Fig. 5A), indicating that the mutation does not alter the catalytic activity of HslV. Furthermore, the inhibitory effect of ATP on peptide hy-



Fig. 4. Structure of HsIV and sequence of the apical pore region. **A.** X-ray crystal structure of an HsIV dodecamer (Protein Data Bank code: 1HT1) viewed from the side proximal to HsIU. The central dark region shows the Arg86 residue in the apical pore region of HsIV. **B.** Alignment of the amino acid sequences of the HsIV pore motifs of various micro-organisms. **C.** Prevalence of each amino acid in the HsIV pore motif shown as a percentage. The data were obtained from 93 reference sequences.

drolysis was dramatically reversed by the R86G mutation (Fig. 5B). These results strongly suggest that binding of ATP to the basic apical pore region, in particular to Arg86, prevents access of unfolded proteins and peptides to the proteolytic core of the HslV dodecamer.

Discussion

The present study has demonstrated that the self-compartmentalizing HslV peptidase can degrade certain unfolded proteins, such as fully unfolded LA and lysozyme, in the absence of HslU and ATP. Furthermore, HslV alone degraded an LA fragment sandwiched by two thioredoxin molecules, indicating that the peptidase can cleave internal peptide bonds. HslVU protease is a bacterial prototype of the eukaryotic 26S proteasome consisting of the 20S proteasome and the 19S regulatory complex (De Mot *et al.*, 1999; Seemuller *et al.*, 1995). Crystallographic and biochemical analyses have demonstrated that the 20S proteasome is a gated protease and that opening of the gate is



Fig. 5. Effect of the R86G mutation in HslV on peptide hydrolysis. A. Wild-type HslV (Wt) or its mutant form (R86G) (20 nM), in which Arg86 was replaced by Gly, was incubated at 37° C with 0.1 mM Z-GGL-AMC and 20 μ M C10-peptide. B. The mixtures were also incubated at 37° C for 30 min in the presence of increasing concentrations of ATP.

achieved by binding of the 19S regulatory complex or PA28 (Groll et al., 2000; Whitby et al., 2000). In the latent state of the 20S proteasome, substrate entry into the inner proteolytic chamber is blocked by the gate formed by the N-terminal tails of certain α -subunits that reside in the outer ring. However, in the absence of the 19S regulatory complex the 20S proteasome can degrade u-LA, but not native LA or partially unfolded intermediates (Wenzel and Baumeister, 1995). It has also been demonstrated that the 20S proteasome efficiently degrades the cyclin-dependent kinase inhibitor of p21 and a-synuclein, both of which are known to be "natively disordered" (Liu et al., 2003). Furthermore, the 20S proteasome degraded p21 and α -synuclein, each of which were sandwiched by two molecules of non-degradable GFP domains, and so lacked both their N- and C-termini. Based on these findings, it was suggested that the latent 20S proteasome can degrade some natively disordered proteins at internal peptide bonds and that these substrates themselves promote gating of the proteasome (Liu et al., 2003). Similarly, certain unfolded proteins, like u-LA and u-lysozyme, may trigger opening of the apical pore of the HslV dodecamer to access the proteolytic chamber.

Noteworthy was the finding that ATP inhibited the degradation of unfolded proteins by HslV whether or not HslU was present. This is in marked contrast to our previous findings that other unfolded proteins, like casein and insulin B-chain, were very poorly degraded by HslV alone, but that their degradation was dramatically stimulated by the presence of HslU and ATP (Rohrwild *et al.*, 1996; Seol *et al.*, 1997; Yoo *et al.*, 1996). A possible explanation for these discrepant results could be a difference in the interaction of two groups of unfolded proteins with HslV and HslU. u-LA and u-lysozyme bind to HslV, but may bind weakly if at all to HslU. On the other hand, casein or the insulin B-chain may interact more strongly with HslU than with HslV, and therefore need to be passed through HslU to reach the inner chamber of HslV peptidase where they are rapidly degraded. Taken together our findings indicate that unfolded protein substrates can be degraded by either uncomplexed HslV or the HslVU complex, respectively, depending on whether they interact preferential with HslV or HslU.

In growing cells, the ATP level is kept constant at millimolar levels. Therefore, uncomplexed HslV is likely to remain in the inactivate state. In this respect, ATP may play an important role in preventing unnecessary breakdown of cellular proteins, such as naturally unstructured proteins, by HslV. However, it has been reported that the ATP pool in E. coli changes nearly 10-fold depending on growth rate (Gaal et al., 1997). In particular, it is likely that the levels of ATP and other nucleotide triphosphates decrease significantly during starvation of an energy source. In addition, it is well documented that carbon starvation and reduced energy production lead to marked increases in rates of proteolysis to replenish energy sources (Goldberg and St. John, 1976; St. John and Goldberg, 1978). Thus, it is possible that HslV plays a role in degradation of unfolded proteins and thus in replenishment of free amino acids as energy sources under conditions in which the cellular level of ATP is low, such as during carbon starvation.

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