Covalently Linked HslU Hexamers Support a Probabilistic Mechanism that Links ATP Hydrolysis to Protein Unfolding and Translocation

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running head: Disulfide-linked HslU pseudohexamers

keywords: AAA+ protease; mixed hexameric rings; protein unfolding; protein degradation; HslU
ABSTRACT

The HslUV proteolytic machine consists of HslV, a double-ring self-compartmentalized peptidase, and one or two AAA+ HslU ring hexamers that hydrolyze ATP to power the unfolding of protein substrates and their translocation into the proteolytic chamber of HslV. Here, we use genetic-tethering and disulfide-bonding strategies to construct HslU pseudohexamers containing mixtures of ATPase active and inactive subunits at defined positions in the hexameric ring. Genetic tethering impairs HslV binding and degradation, even for pseudohexamers with six active subunits, but disulfide-linked pseudohexamers do not have these defects, indicating that the peptide tether interferes with HslV interactions. Importantly, pseudohexamers containing different patterns of hydrolytically active and inactive subunits retain the ability to unfold protein substrates and/or collaborate with HslV in their degradation, supporting a model in which ATP hydrolysis and linked mechanical function in the HslU ring operate by a probabilistic mechanism.
Enzymes of the AAA+ ATPase superfamily play roles in proteolysis, protein remodeling and disaggregation, replication, transcription, membrane fusion, vesicle transport, and other cellular processes in all organisms (1-2). These enzymes share conserved sequence and structural motifs and typically function as homohexameric or heterohexameric rings. Fueled by the energy of ATP binding and hydrolysis, AAA+ enzymes act as molecular machines that disassemble, remodel, or denature macromolecule targets. There are three general models for how subunits in AAA+ hexamers hydrolyze ATP and generate the mechanical power strokes required for function: (i) concerted ATP hydrolysis that occurs simultaneously in all subunits (3); (ii) sequential hydrolysis by individual subunits that occurs in an invariant kinetic pattern (4); and (iii) probabilistic hydrolysis in which following a power stroke, any ATP-bound subunit has some chance of hydrolyzing ATP to drive the next power stroke (5).

The HslUV protease consists of one or two AAA+ HslU hexamers and the dodecameric HslV peptidase (Fig. 1A; refs. 6-12). In ATP-dependent reactions, HslU hexamers recognize protein substrates, unfold any native structure that is present, and then translocate the unfolded polypeptide into the luminal chamber of HslV for degradation. In all crystal structures of the *Escherichia coli* or *Haemophilus influenza* HslUV complexes and some structures of HslV alone, the HslU hexamer is highly symmetric and binds six ATP or ADP molecules, as might be expected for a concerted mechanism of hydrolysis (6-12). In other structures of HslU alone, only three or four nucleotides are bound to the hexameric HslU ring (6), and solution experiments show detectable binding of a maximum of 3 to 4 nucleotides and the existence of at least two types of nucleotide-binding sites (13). These results suggest that the six subunits of HslU assume non-equivalent functional roles within the hexamer and are more consistent with sequential or probabilistic models. Here, we test different models by which ATP hydrolysis could power the mechanical functions of *E. coli* HslU by introducing hydrolytically inactive subunits at defined positions in its hexameric ring. Using subunit crosslinking strategies involving genetic tethering or disulfide bonding, we find that HslU pseudohexamers with mixtures of hydrolytically active and inactive subunits retain protein unfolding activity and support HslV degradation. These studies support a probabilistic mechanism in which ATP hydrolysis powers mechanical function in the HslU ring and also reveal new information about interactions between HslU and HslV.

**RESULTS**

**HslU dimers with covalent peptide tethers.** HslU homohexamers containing the Walker-B E257Q mutation are defective in ATP hydrolysis and protein degradation but retain the ability to bind HslV and protein substrates (13). We engineered genes to encode tandem *E. coli* HslU subunits connected by a 20-residue peptide tether (Fig. 1B). One encoded dimer consisted of two wild-type subunits (W-W), another had a wild-type subunit followed by a E257Q subunit (W-E), and a third had a E257Q subunit followed by a wild-type subunit (E-W). These dimers behaved like wild-type HslU during purification, suggesting that they form W-W₃, W-E₃, and E-W₃ pseudohexamers. Indeed, the asymmetric unit of a low-resolution W-E₃ crystal structure contained four hexamers similar to wild-type HslU (Fig. 1C; Table 1), although electron density for the C-terminal 8-10 residues was missing in alternating subunits of several hexamers or was generally poor.
throughout a hexamer, as expected if the tether disrupts normal C-terminal contacts.

W-W₃ hydrolyzed ATP at about twice the rate of the W-E₃ and E-W₃ enzymes (Fig. 2A), suggesting that ATP hydrolysis is largely restricted to the W subunits in these enzymes. We constructed and expressed a W-W-W trimer, but this protein was insoluble. To assay protein unfolding, we constructed an 137A Arc-GFP-st11-ssrA fusion protein with a thrombin cleavage site located between β-strands 5 and 6 (14). 137A Arc is a denatured variant of Arc repressor that targets the substrate to HslU, and the C-terminal st11-ssrA sequence increases the substrate turnover rate ~2-fold (15, 16). Following thrombin cleavage, unfolding of the split substrate by wild-type HslU results in an irreversible loss of GFP fluorescence. In experiments performed at different concentrations of the split substrate, unfolding by wild-type HslU and W-W₃ occurred with steady-state Vₘₐₓ rates that were similar, whereas Vₘₐₓ for unfolding by the W-E₃ and E-W₃ pseudohexamers was about half of the wild-type value (Fig. 2B). Thus, pseudohexamers with alternating ATPase active and inactive subunits retain substantial protein-unfolding activity. However, compared to wild-type HslU, W-W₃ supported HslV degradation very poorly (Fig. 2C) and bound HslV ~20-fold more weakly (Table 2), probably because the peptide tether interferes with contacts between HslU and HslV (see Discussion). Thus, we explored a different method of constructing covalently linked HslU hexamers.

Construction of disulfide-crosslinked HslU pseudohexamers. HslU hexamers consist of rigid-body units formed by the large and small domains of adjacent subunits (17). We used the disulfide-by-design algorithm (18) to identify Glu⁴⁷/Ala³⁴⁹ and Gln³⁹/Thr³⁶¹ as sites for potential disulfide bonds across the rigid-body interfaces of an HslU hexamer. Fig. 3A shows a model of an otherwise Cys-free HslU pseudohexamer in which red subunits contain Cys⁴⁷ and blue subunits contain Cys³⁴⁹, potentially allowing formation of three disulfide-linked dimers. To make WSSW₃ pseudohexamers, the Cys⁴⁷ and Cys³⁴⁹ subunits both had wild-type Walker-B ATPase motifs. To make WSS₂E₃ pseudohexamers, the Cys⁴⁷ subunit had a wild-type Walker-B sequence and the Cys³⁴⁹ subunit contained the Walker-B E₂⁵⁷Q mutation to inactivate ATP hydrolysis. Fig. 3B shows a pseudohexamer in which red subunits contain Cys³⁴⁹, green subunits contain Cys⁴⁷ and Cys³⁶¹, and blue subunits contain Cys³⁹. In this configuration, formation of disulfide-linked trimers is possible. We designed WSS₂WSS₂W trimers, WSS₂E₂W trimers, and WSS₂E₂E trimers by changing which subunits had wild-type or E₂⁵⁷Q Walker-B sites.

Relatively efficient formation of disulfide-linked HslU dimers or trimers was achieved by cytosolic coexpression of appropriate variants in the oxidizing SHuffle strain of E. coli (19). For example, following purification by Ni-NTA affinity chromatography, non-reducing SDS-PAGE showed ~50% formation of WSS₂W and WSS₂E and ~33% formation of WSS₂WSS₂W, WSS₂ESS₂W, and WSS₂E₂₂E (not shown). To further purify disulfide-linked pseudohexamers, we performed ion-exchange chromatography and gel-filtration chromatography once in the presence and once in the absence of urea, which destabilizes unlinked HslU hexamers more than linked hexamers. Following the final chromatography step, the WSS₂W, WSS₂E, WSS₂WSS₂W, WSS₂E₂₂W, and WSS₂E₂₂E variants had purities of >95% (Fig. 3C). The disulfide-linked pseudohexamers bound
HslV slightly more tightly than wild-type HslU (Table 2).

**ATP hydrolysis by disulfide-linked pseudohexamers.** Like the basal ATP-hydrolysis activities of the genetically-tethered pseudohexamers, those of the disulfide-linked enzymes were roughly proportional to the number of hydrolytically active W subunits in each pseudohexamer (Figs. 4A, 4B). Compared to wild-type HslU, however, the W_{SS}W_{3} and W_{SS}W_{SS}W_{2} pseudohexamers were ~3-fold more hydrolytically active (Fig. 4A), possibly as a consequence of small conformational changes stabilized by the disulfide bonds. In the presence of HslV, the ATPase rate of wild-type HslU was stimulated ~3-fold, as previously observed (13, 20, 21), ATP hydrolysis by W_{SS}W_{3} and W_{SS}W_{SS}W_{2} was stimulated ~2-fold, but ATP hydrolysis by W_{SS}E_{3}, W_{SS}E_{SS}W_{2}, or W_{SS}E_{SS}E_{2} was not markedly stimulated (Fig. 4C). Thus, the presence of E subunits suppressed normal HslV stimulation of ATP hydrolysis by HslU pseudohexamers.

**Degradation supported by disulfide-linked pseudohexamers.** Arc repressor, a good substrate for HslUV degradation, is a metastable dimer that unfolds/dissociates with a half-life of ~10 s but refolds in milliseconds to maintain a predominantly native structure (15, 22). We assayed the ability of different disulfide-linked pseudohexamers to support HslV degradation of Arc-cysA, where cysA designates a unique cysteine labeled with an Alexa-488 fluorophore (23), as auto-quenching of the fluorophores in the native protein is relieved upon degradation. The W_{SS}W_{3} and W_{SS}W_{SS}W_{2} pseudohexamers supported HslV degradation of a near-saturating concentration of Arc-cysA at rates comparable to the wild-type HslU hexamer (Fig. 5A). Importantly, W_{SS}E_{SS}W_{3} and W_{SS}E_{3} also supported degradation at 35-45% of the wild-type rate, demonstrating that pseudohexamers with only three or four hydrolytically active subunits also have substantial degradation activity. Degradation supported by W_{SS}E_{SS}E_{2} proceeded very slowly, at ~3% of the wild-type rate.

The degradation defects caused by E subunits in pseudohexamers were more severe for Arc-cp6GFP-st11-ssrA, a stable substrate that wild-type HslU degrades ~5-fold more slowly than Arc-cysA. W_{SS}W_{3} and W_{SS}W_{SS}W_{2} supported HslV degradation of a high concentration of Arc-cp6GFP-st11-ssrA at near wild-type rates, but W_{SS}E_{SS}W_{2} had only ~20% activity, W_{SS}E_{3} had only ~10% activity, and W_{SS}E_{SS}E_{2} was inactive (Fig. 5B). Degradation rates were normalized by dividing by the number of wild-type subunits in HslU or different variants and are plotted in Fig. 5C for the Arc-cysA substrate and Fig. 5D for the Arc-cp6GFP-st11-ssrA substrate. For Arc-cysA, there was a sharp discontinuity between 2 and 3 wild-type subunits. For Arc-cp6GFP-st11-ssrA, by contrast, the major discontinuity was between 4 and 6 subunits.

To determine the energetic efficiency of degradation of Arc-cp6GFP-st11-ssrA, we assayed the rate of ATP-hydrolysis for each disulfide-linked pseudohexamer in the presence of HslV and Arc-cp6GFP-st11-ssrA (Fig. 5E). We then divided the ATPase rate by the degradation rate to determine the average number of ATPs hydrolyzed during degradation of a single substrate (Fig. 5F). Notably, W_{SS}W_{3}, W_{SS}E_{3}, W_{SS}W_{SS}W_{2}, and W_{SS}E_{SS}W_{2} all had similar energetic efficiencies, hydrolyzing ~500 ± 100 ATPs for each substrate degraded. Assuming that power strokes are tightly coupled to ATP hydrolysis, this result suggests that the
**DISCUSSION**

Our studies show that *E. coli* HslU variants containing hydrolytically active and inactive subunits at specific positions in the hexameric AAA+ ring can hydrolyze ATP, unfold proteins, and degrade substrates in collaboration with HslV. As we discuss below, these results support a probabilistic model of ATP hydrolysis and provide insights into the multivalent interactions between HslU and HslV that are required for efficient protein degradation.

Genetic tethering allowed us to express and purify HslU pseudohexamers consisting of a trimer of linked dimers. However, the W-W₃ enzyme binds HslV poorly, suggesting that the tether interferes with HslV binding. Consistently, disulfide-linked pseudohexamers bind HslV well. In crystal structures of HslU hexamers alone, the C-terminal tails dock into a pocket and the α-carboxyl group forms a salt bridge with an arginine in the sensor-2 motif of the same subunit (6, 10). These tail interactions were disrupted in several subunits in our low-resolution structure of W-E₃ pseudohexamers, as expected if the attached tether prevents proper packing of these residues. In the *H. influenza* HslUV complex, the C-terminal tails are detached from HslU and pack into grooves on HslV, with the HslU α-carboxylate forming a salt bridge with a HslV lysine side chain (7, 11, 12). In *E. coli* HslUV structures, by contrast, the tails remain docked into HslU (8, 9). Moreover, deletion of the five C-terminal residues of *E. coli* HslU does not prevent stimulation of HslV peptidase activity or degradation (20). Nevertheless, peptides corresponding to the C-terminal residues of HslU activate peptide cleavage by *E. coli* HslV, and mutations in HslV predicted to disrupt contacts with the C-terminal tails prevent HslU activation (20, 24). Our results support the importance of the C-terminal tails in high-affinity HslV binding and indicate that more than three tails of *E. coli* HslU must interact optimally with HslV to allow tight binding and efficient proteolysis.

The pseudohexamers we studied have basal ATP hydrolysis rates roughly proportional to their total number of hydrolytically-active W subunits. This result supports a model in which the W subunits in these pseudohexamers contribute independently to basal ATPase activity, making concerted or strictly sequential models unlikely. For example, if ATP hydrolysis in a specific subunit required prior hydrolysis in a neighboring subunit, as expected in a strictly sequential model, then a non-linear relationship between W subunits and ATP hydrolysis would be expected. HslV stimulates ATP hydrolysis by W₃E₃ and W₃E₃W₂ but caused little change in hydrolysis by W₃E₃W₂, W₃E₃, or W₃E₃E₂. As the E subunits in these latter enzymes also increase pseudohexamer affinity for HslV, stronger interactions between E subunits and HslV might restrict HslV-induced conformational changes required for higher ATPase activity in neighboring W subunits and thus explain the lack of ATPase stimulation. Alternatively, HslV binding might slow ATP dissociation from inactive E subunits, which becomes rate-limiting for hydrolysis in W subunits.

W₃E₃E₃W₂ and W₃E₃ pseudohexamers use approximately the same number of power strokes as hexamers with six wild-type subunits to unfold and translocate Arc-cp6GFP-st11-ssrA. Thus, the slower degradation activities of the W₃E₃E₃W₂ and W₃E₃ enzymes compared to pseudohexamers with only wild-type subunits are principally a consequence of their slower rates of ATP hydrolysis.
especially if only a subset of subunits is nucleotide bound at any given time.

Our strategies for engineering HslU rings with defined mixtures of active and inactive subunits were motivated by prior studies that applied these subunit-crosslinking methods to different hexameric AAA+ unfoldases and remodeling machines (5, 25-27). Indeed, genetic-tethering experiments originally showed that ClpX also operates by a probabilistic mechanism, as ClpX pseudohexamers containing different combinations of hydrolytically active and inactive subunits unfold and degrade protein substrates in collaboration with the ClpP protease (5). Prior to the current study, however, it was not obvious that HslU and ClpX would operate using similar probabilistic mechanisms of ATP-hydrolysis. First, HslU and ClpX contain unique family specific auxiliary domains. The I-domain of HslU emerges from the top of the AAA+ ring, between the Walker-A and Walker-B ATPase motifs, and regulates ATP hydrolysis, degradation, and autoinhibition (9, 20, 23). The N-domain of ClpX, by contrast, serves as a docking site for some adaptors/substrates but its deletion has little effect on ATP hydrolysis or degradation of many ClpXP substrates (28). Second, in crystal structures, the large and small AAA+ domains of each HslU subunit assume orientations that create a potential nucleotide binding site, whereas the corresponding domains of ClpX adopt structures that allow nucleotide binding in some subunits but prevent binding in other subunits (6-12, 29). Third, HslU hexamers make symmetric interactions with hexameric rings of HslIV, whereas ClpX hexamers make asymmetric interactions with heptameric rings of ClpP (30).

The tethered W-E₃ and E-W₃ HslU pseudohexamers have ~50% of the protein unfolding activity of the parental W-W₃ enzyme, and the disulfide-linked W⁸⁴E₃⁻W₂ and W⁸⁴E₃ enzymes support HslV degradation of an easily degraded Arc-CysA substrate at 35-45% of the parental rates. Thus, protein unfolding and translocation by the AAA+ ring of HslU do not require ATP hydrolysis in adjacent subunits or in subunits immediately across the ring from each other. Again, these results support a model in which probabilistic hydrolysis in the AAA+ ring powers unfolding and translocation. Slower rates of ATP hydrolysis in rings with mixtures of active and inactive subunits correlate with their reduced mechanical activities, as the ATP cost of degradation of the more stable Arc-cp₆GFP-st11-ssrA substrate is similar for the W⁸⁴E₃⁻W₂ and W⁸⁴E₃ pseudohexamers and their disulfide-linked parental enzymes containing six W subunits. Although optimal rates of ATP hydrolysis, unfolding, and degradation require six hydrolytically active HslU subunits, rings with only three or four active subunits use approximately the same number of power strokes to degrade this substrate.

We note that the W⁸⁴E₃⁻⁸⁴E₃ pseudohexamer hydrolyzes ATP at ~40-60% of the W⁸⁴E₃ rate but supports no degradation of Arc-cp₆GFP-st11-ssrA. These results suggest that some subunit-subunit communication is required for efficient unfolding and degradation by W⁸⁴E₃⁻E₃ and/or that a minimum rate of ATP hydrolysis is required to unfold Arc-cp₆GFP-st11-ssrA. Probabilistic models of ATP hydrolysis do not exclude coordination between ring subunits for efficient ATP hydrolysis, substrate binding, or mechanical function. In fact, the rate of ATP hydrolysis and the degree of substrate binding by HslU change in a positively cooperative fashion with ATP
concentration, as expected for functional linkage between different subunits (13). In the AAA+ ring of ClpX, communication between subunits is necessary to allow staged ATP binding to drive conformational changes needed for function (31-33). Moreover, optical-trapping experiments reveal that kinetic bursts of power strokes in the ClpX ring result in random patterns of shorter and longer translocation steps, supporting a probabilistic but coordinated mechanism (34-35). Similar themes of probabilistic hydrolysis but coordinated function are seen in AAA+ unfolding rings assembled from non-identical subunits. In the Yta10/Yta12 m-AAA protease, for example, mutating the Walker-A motif of either the Yta10 or the Yta12 subunits does not affect hydrolysis in the remaining wild-type subunits, but Walker-B mutations in Yta12 trap ATP and prevent robust hydrolysis in adjacent Yta10 subunits (36).

Finally, in the six distinct subunits of the Rpt1-6 AAA+ ring of the 26S proteasome, Walker-B mutations in individual subunits have a wide range of effects on ATP hydrolysis and mechanical activity, requiring a model with some subunit-subunit coordination within the context of an inherently probabilistic mechanism of ATP hydrolysis (37).

EXPERIMENTAL PROCEDURES

Cloning, expression, and protein purification. Mutants were constructed and cloned by standard PCR techniques unless noted. To construct genetically linked HslU dimers, a gene encoding two HslU subunits separated by the 20-residue ASAGGGSEGSEGTTSGAT linker was cloned into the pet11a vector (Novagen). HslU mutants used to make disulfide-crosslinked pseudohexamers were constructed in the cysteine-free C262A/C288S HslU background with or without the E257Q mutation. Cysteine-free HslU supports robust ATP hydrolysis and substrate degradation (38-40). To make disulfide crosslinked dimers, a gene encoding untagged E47C HslU was cloned into the first multiple cloning site (MCS1) of the pCOLADuet-1 (Novagen) vector between the NcoI and BamHI sites and a gene encoding His$_6$ENLYFQS-A349C HslU was cloned into MCS2 between the Ndel and Xhol sites, where His$_6$ is the hexahistidine tag and ENLYFQS is the sequence recognized and cleaved by the TEV protease. E47C HslU had a glycine residue inserted after the initiator methionine as a result of cloning. This pCOLADuet-1 vector was transformed into the of E. coli SHuffle T7 Express strain (New England Biolabs) for expression. To make disulfide crosslinked trimers, a gene encoding untagged A349C HslU was cloned into MCS1 of pCOLADuet-1 between the NcoI and BamHI sites and a gene encoding His$_6$-ENLYFQS-E47CT361C HslU was cloned into MCS2 between NdeI and XhoI sites. This vector was co-transformed with a pet12b (Novagen) vector encoding the untagged Q39C HslU variant into the SHuffle T7 Express strain. Both A349C HslU and Q39C HslU contained an additional glycine after the initiator methionine as a result of cloning. A gene encoding the Arc repressor from phage P22 followed by a cysteine residue and a hexahistidine tag (Arc-cys-His$_6$) was cloned and expressed in a pet21b vector (Novagen). Wild-type HslU and HslV were expressed from pet12b vectors. Arc-cp6GFP-st11-ssrA, and 137A-Arc-cp6GFP-β5/β6-st11-ssrA (a GGTEGSLVPRGSGESGGS sequence between β-stands 5 and 6 allows thrombin cleavage and generation of a split substrate), and Arc-st11-ssrA were expressed from pet21b vectors as described (14, 15, 23).
Genetically linked HslU dimers were expressed and purified as previously described for wild-type HslU (20). Disulfide crosslinked HslU dimers were expressed and purified as follows. *E. coli* SHuffle T7 Express cells carrying the pCOLA-Duet1 vector coding for the appropriate HslU mutants were grown at 30 °C until OD 0.6-0.8, the temperature was shifted to 18 °C, and protein expression was induced with 0.5 mM IPTG for 20 h. Cells were pelleted and resuspended in buffer A (50 mM Tris, pH 7.5, 300 mM NaCl, 20 mM imidazole, 0.5 mM EDTA), and 0.5 tablets of Complete Ultra EDTA-free protease-inhibitor cocktail (Roche) and 1.5 µL benzonase (250 units/µL, Sigma) per liter of the original culture were added. Cells were sonicated, the lysate was cleared by centrifugation, and 0.1% v/v polyethyleneimine (PEI) was added to the supernatant. This precipitate was cleared by centrifugation and the supernatant was loaded onto Ni²⁺-NTA beads equilibrated in buffer B (50 mM Tris, pH 7.5, 300 mM NaCl, 20 mM imidazole). The Ni²⁺-NTA beads were washed extensively with buffer B and protein was eluted with buffer C (50 mM Tris, pH 7.5, 300 mM NaCl, 250 mM imidazole). The eluate was diluted 3-fold with buffer D (50 mM Tris, pH 7.5, 10% (v/v) glycerol, 1 mM EDTA), loaded onto a Mono Q 10/100 GL column (GE Healthcare), and eluted with a linear gradient from 150 to 500 mM NaCl in buffer D (120 mL total). Appropriate Mono Q fractions were pooled, concentrated using an Amicon Ultra-15 centrifugal filter unit, chromatographed on a Superdex 200 16/60 column (GE Healthcare) equilibrated in buffer E (50 mM Tris, pH 7.5, 300 mM NaCl, 10% (v/v) glycerol, 1.5 M urea, 1 mM EDTA) and 1 mL fractions were collected. Fractions with the highest purity of the crosslinked dimer as judged by non-reducing SDS-PAGE were pooled, concentrated, and run on another Superdex 200 16/60 column equilibrated in buffer F (50 mM Tris, pH 7.5, 300 mM NaCl, 10% (v/v) glycerol, 1 mM EDTA). Fractions with the highest purity of crosslinked dimer were pooled and concentrated. The concentration of crosslinked dimer was determined in hexameric equivalents by measuring the absorbance at 280 nm using 148,545 M⁻¹ cm⁻¹ as the molar extinction coefficient. Concentrated protein was divided into small aliquots and was flash frozen at –80 °C.

Disulfide-crosslinked HslU trimers were expressed and purified largely as described for crosslinked dimers. His₆ tags were present on Cys³⁴⁹ subunits for dimers and Cys⁴⁷/Cys³⁶¹ subunits for trimers. After precipitation with PEI, the sample was cleared by centrifugation and the supernatant was loaded onto a 5 mL HisTrap HP column (GE Healthcare) equilibrated in buffer B. The column was washed with 75 mL of buffer B, the sample was eluted with a gradient of 20 to 500 mM imidazole in buffer B (100 mL total), and 2 mL fractions were collected. Fractions were pooled and dialyzed overnight against buffer G (50 mM Tris, pH 7.5, 150 mM NaCl, 10% glycerol (v/v), 1 mM EDTA) at 4 °C. The dialyzed material was loaded onto a Mono Q 10/100 GL column and purification then proceeded by the method described for crosslinked dimers. The concentration of the crosslinked trimer was determined in hexameric equivalents by measuring absorbance at 280 nm using 147,180 M⁻¹ cm⁻¹ as the molar extinction coefficient. Concentrated protein was aliquoted and flash frozen at –80 °C.

A pet21b vector carrying Arc with a C-terminal CHHHHHHH tail (Arc-cys-His₆) was transformed into the *E. coli* X90 (λDE3) slyD::kan hs/U::tet strain and cells were grown to an OD₆₀₀ of 0.6-0.8 at 37 °C. Protein expression was induced by the
addition of 1 mM IPTG and continued for 4 h at room temperature. Cells were resuspended, lysed, and Arc-cys-His₆ was purified by Ni²⁺-NTA affinity chromatography as described for purification of crosslinked HslU dimers. After Ni²⁺-NTA purification, protein was concentrated and injected onto a Superdex 75 16/60 column (GE Healthcare) equilibrated in buffer H (50 mM Tris, pH 7.5, 300 mM NaCl, 10% glycerol (v/v), 1 mM EDTA, 2 mM DTT). Fractions containing the Arc protein were concentrated and flash frozen at –80 °C. Arc-cys-His₆ was labeled with the maleimide-derivative of Alexa-488 (Thermo Fisher Scientific) to generate Arc-cysA (23). His₆-tagged wild type HslU, His₆-tagged HslV, I₃₇₅Arc-cp₆GFP-β₅/β₆-st11-ssrA, Arc-cp₆GFP-st11-ssrA, and Arc-st11-ssrA were expressed and purified as described (23).

**Biochemical assays.** Unless noted, assays were performed at 37 °C in PD buffer (25 mM HEPES, pH 7.5, 5 mM KCl, 10% glycerol (v/v), 20 mM MgCl₂, 0.032% Igepal CA-630). Hydrolysis of 5 mM ATP was measured using an NADH-coupled assay (41) by monitoring the loss of absorbance at 340 nm on a Spectramax M5 plate reader (Molecular Devices). Cleavage of I₃₇₅Arc-cp₆GFP-β₅/β₆-st11-ssrA with thrombin was performed as described (14). Rates of unfolding of different concentrations of thrombin-split I₃₇₅Arc-cp₆GFP-β₅/β₆-st11-ssrA were determined by changes in cp₆GFP fluorescence (excitation 467 nm; emission 511 nm) in assays that contained 0.5 µM HslU or tethered variants and 5 mM ATP. Degradation of 30 µM Arc-cysA (monomer equivalents) was measured by monitoring the decrease in GFP fluorescence on a Spectramax M5 plate reader (excitation 467 nm; emission 511 nm). Degradation reactions contained 5 mM ATP and a regeneration system consisting of 16 mM creatine phosphate and 10 µg/mL creatine kinase. HslV activation assays were performed at 25 °C in the presence of ATP as described (13, 23), and K₁/₂ values were determined by fitting to a hyperbolic equation.

**Crystallography.** The W-E₃ pseudohexamer was crystallized by the hanging-drop method using 100 mM Bis-Tris (pH 5.8), 26% (w/v) PEG 3350, and 260 mM ammonium sulfate as the well solution. Molecular replacement using Phaser (42) was initially used to solve the structure using a 1HQY hexamer (17) as the search model. We then replaced each 1HQY subunit with a 5JI3 subunit (23) to improve geometry and used rigid-body refinement of individual domains, refinement of one B-factor and TLS group per subunit, and very tightly constrained positional refinement with torsional NCS constraints in Phenix (43). Coot (44) was used for model building, and MolProbity (45) was used to assess the geometry of the model.

**Acknowledgements.** Supported by NIH grant AI-16892. T.A.B. is an employee of the Howard Hughes Medical Institute.

**Conflict of interest.** The authors declare no conflict of interest.

**Author contributions.** J.C. performed experiments with HslU subunits linked by genetically encoded tethers. A.R.N. designed the split substrate for unfolding
experiments. S.E.G., R.A.G., and R.T.S. performed crystallographic experiments. V.B. performed all remaining experiments. V.B. and R.T.S. wrote the manuscript. All authors contributed to the design and interpretation of experiments and approved the final manuscript.
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Figure legends

**Figure 1. HslUV structure.** (A) An HslU hexamer (cartoon representation) bound to an HslV dodecamer (surface representation; pdb code 1g3i). The large and small AAA+ domains of HslU and its C-terminal tails are colored blue, cyan, and red, respectively. (B) Tandem HslU subunits connected by a genetically encoded peptide tether. (C) Three W-E₃ hexamers in the asymmetric unit of structure 5TXV are shown in cartoon representation; the fourth hexamer is shown in ribbon representation with electron density from a composite omit map contoured at 1σ.

**Fig. 2. Activity of genetically tethered pseudohexamers.** (A) Rates of hydrolysis of 5 mM ATP by the genetically tethered W-W₃, W-E₃, and E-W₃ pseudohexamers. Values are averages (N ≥ 5) ± SD. (B) Rates of unfolding of different concentrations of thrombin-split I₃₇Δarc-cp6GFP-β5/β6-st11-ssrA by wild-type HslU or genetically-tethered variants. Lines are non-linear least-squares fits to the Michaelis-Menten equation. Kₘ’s values for all enzymes were 1-3 μM but were not well determined because of the small number of low-concentration data points. Average Vₘₐₓ values ± SD calculated from the highest four substrate concentrations were 0.155 ± 0.003 min⁻¹ enz⁻¹ (HslU), 0.143 ± 0.008 min⁻¹ enz⁻¹ (W-W₃), 0.0802 ± 0.007 min⁻¹ enz⁻¹ (E-W₃), and 0.0716 ± 0.006 min⁻¹ enz⁻¹ (W-E₃). Fitted Vₘₐₓ values were 10-15% higher. (C) The kinetics of degradation of Arc-st11-ssrA (10 μM) by HslV (10 μM) and HslU (0.3 μM) or W-W₃ (0.3 μM) at 50 °C was monitored by SDS-PAGE. The first lane contains molecular weight standards.

**Fig. 3. Design and purification of disulfide-crosslinked HslU variants.** (A) Spheres show the positions of Cys⁴⁷ (normally Glu) in red subunits and Cys³⁴⁹ (normally Ala) in blue subunits of an HslU hexamer, suggesting that Cys⁴⁷-Cys³⁴⁹ disulfides would stabilize a pseudohexamer consisting of three linked dimers. (B) Spheres show the positions of Cys³⁴⁹ in red subunits, Cys⁴⁷ and Cys³⁶¹ (normally Thr) in green subunits, and Cys³⁹ (normally Gln) in blue subunits. Disulfide-bond formation in this configuration would stabilize a pseudohexamer consisting of two linked trimers. (C) Non-reducing SDS-PAGE of purified wild-type HslU, purified W₅₆W, purified W₅₆E, purified W₅₆W₅₆W, purified W₅₆E₅₆W, and purified W₅₆E₅₆E. In each case, 0.5 μM of the purified protein (in hexamer equivalents) was loaded on the gel.

**Fig. 4. ATP hydrolysis by disulfide-crosslinked variants.** (A) Basal rates of ATP hydrolysis by disulfide-crosslinked variants and wild-type HslU. Assays contained 0.3 μM HslU or pseudohexamers and 5 mM ATP. Values are averages of at least three replicates ± 1 SD. (B) Basal ATP hydrolysis rates for disulfide-crosslinked pseudohexamers plotted as a function of the number of W subunits. (C) Rates of ATP hydrolysis determined in the presence of 0.9 μM HslV dodecamer (other conditions as in panel A). The numbers above each bar represent the rate in the presence of HslV divided by the rate in the absence of ATP.
Fig. 5. Degradation rates and energetic efficiencies of disulfide-crosslinked variants. (A) Rates of degradation of Arc-cysA (30 µM) assayed by increased fluorescence. (B) Rates of degradation of Arc-cp6GFP-st11-ssrA (20 µM) assayed by decreased fluorescence. (C) Degradation rates for Arc-cysA from panel A were divided by the number of wild-type subunits in the HslU variant and plotted against this number. (D) Degradation rates for Arc-cp6GFP-st11-ssrA from panel B were divided by the number of wild-type subunits in the HslU variant and plotted against this number. (E) Rates of ATP hydrolysis in the presence of 20 µM Arc-cp6GFP-st11-ssrA and 1 µM HslV. (F) Energetic efficiency determined by dividing the ATPase rates by the degradation rates. For panels A-D, experiments were performed using 0.3 µM HslU or variants and 0.9 µM HslV. For panels, E and F, experiments were performed using 0.5 µM HslU or variants and 1 µM HslV. All assays contained 5 mM ATP and were performed at 37 °C. Values in panels A-E are averages (N ≥ 3) ± 1 SD. The error bars in panel F represent propagated errors. In panels C and D, values for the three variants with six wild-type subunits were offset slightly on the x-axis to allow visualization of error bars.
Table 1. Crystallographic statistics.

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<th>Description</th>
<th>Value</th>
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<td>wavelength (Å)</td>
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<td>space group</td>
<td>P 1 2 1</td>
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<td>Cβ deviations</td>
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Values in parenthesis represent the highest resolution shell.
Table 2. HslV binding.

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<th>protein</th>
<th>$K_{1/2}$ (nM)</th>
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<tr>
<td>wild-type HslU</td>
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<tr>
<td>W-W$_3$</td>
<td>1603 ± 278</td>
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<td>W$_{SS}W_3$</td>
<td>59 ± 9</td>
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<tr>
<td>W$<em>{SS}W</em>{SS}W_2$</td>
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<td>W$<em>{SS}E</em>{SS}W_2$</td>
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<td>W$_{SS}E_3$</td>
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<td>W$<em>{SS}E</em>{SS}E_3$</td>
<td>28 ± 19</td>
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Titration assays, monitored by changes in HslV peptidase activity, were performed at 25 °C in the presence of 5 mM ATP.
Figure 1

A

large AAA+ domains
small AAA+ domains
HslU
HslV
C-tails

B

peptide tethered HslU dimer
first HslU subunit
C-tail
large AAA+
small AAA+
second HslU subunit
small AAA+
large AAA+

C

W-E₃ hexamers

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Figure 2

A

ATP hydrolyzed (min⁻¹ enz⁻¹)

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<thead>
<tr>
<th></th>
<th>W-W₃</th>
<th>W-E₃</th>
<th>E-W₃</th>
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<tr>
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<tr>
<td>E-W₃</td>
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</table>

B

Unfolding rate (min⁻¹ enz⁻¹)

[protein substrate] (µM)

C

Degradation time (min)

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<tr>
<td>E-W₃/HslV</td>
<td>E-W₃/HslV</td>
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</tr>
</tbody>
</table>

---

Arc-st11-ssrA
Figure 3

(A) 3 SS-linked dimers form pseudohexamer

(B) 2 SS-linked trimers form pseudohexamer

(C) SDS-PAGE gel showing protein standards and recombinant protein samples.
Figure 4
Figure 5
Covalently Linked HslU Hexamers Support a Probabilistic Mechanism that Links ATP Hydrolysis to Protein Unfolding and Translocation
Vladimir Baytshtok, Jiejen Chen, Steven E. Glynn, Andrew R. Nager, Robert A. Grant, Tania A. Baker and Robert T. Sauer

J. Biol. Chem. published online February 21, 2017

Access the most updated version of this article at doi: 10.1074/jbc.M116.768978

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