Structural basis for intersubunit signaling in a protein disaggregating machine

Amadeo B. Biter, Sukyeong Lee, Nuri Sung, and Francis T.F. Tsai

ClpB is a ring-forming, ATP-dependent protein disaggregase that cooperates with the cognate Hsp70 system to recover functional protein from aggregates. How ClpB harnesses the energy of ATP binding and hydrolysis to facilitate the mechanical unfolding of previously aggregated, stress-damaged proteins remains unclear. Here, we present crystal structures of the ClpB D2 domain in the nucleotide-bound and -free states, and the fitted cryoEM structure of the D2 hexamer ring, which provide a structural understanding of the ATP power stroke that drives protein translocation through the ClpB hexamer. We demonstrate that the conformation of the substrate-translocating pore loop is coupled to the nucleotide state of the cis subunit, which is transmitted to the neighboring subunit via a conserved but structurally distinct intersubunit-signaling pathway common to diverse AAA+ machines. Furthermore, we found that an engineered, disulfide cross-linked ClpB hexamer is fully functional biochemically, suggesting that ClpB deoligomerization is not required for protein disaggregation.

Results

Crystal Structure of the ClpB D2 Domain. To visualize the structural basis for the ATP-driven power stroke, we crystallized the isolated D2 domain (residues 542–854) of Thermus thermophilus ClpB (Fig. S1), featuring an E668A mutation in the Walker B motif. This Walker B mutant can bind nucleotide and, when combined with the analogous E271A mutation in the Walker B motif of the D1 domain, renders ClpB ATP-hydrolysis deficient (8, 36). Three different crystal forms were obtained in the presence or absence of nucleotide, representing a total of seven independent structures of the D2 domain (Fig. L4 and Fig. S2 and Table S1). Notably, crystals obtained in the presence of nucleotide diffracted consistently to higher resolution, but were merohedrally twinned (form I). To determine the crystal structure of the nucleotide-bound conformer, we prepared a selenomethionine (SeMet) derivative of a D2 E668A variant that featured three additional point mutations (I683M, L706M, and L770M). Fortuitously, in addition to twinned crystals, this D2 variant also produced a new, untwinned crystal form (form II) in the same drop, which was used for structure determination by molecular replacement. Although the best crystal of the nucleotide-bound D2 domain diffracted to 3.3 Å resolution, the data were challenging to refine with the M-domain being on the interior side in form I and intercalated between subunits in form II. The structure of the fully functional biochemically active D2 E668A nanoparticle suggested that the M-domain is conformationally disordered and not required for protein disaggregation.

ATPase | chaperone | Hsp100 | protein unfoldase

ClpB is an ATP-dependent protein-remodeling machine that has the remarkable ability to rescue stress-damaged proteins from a previously aggregated state. As the major protein disaggregase in cells, bacterial ClpB and its yeast (Hsp104) and plant (Hsp101) homologs are essential for thermotolerance development (1–3), and for cell survival from acute stress conditions (4).

At the molecular level, ClpB is a multidomain protein composed of two tandem Walker-type ATP-binding domains (AAA+ domains), termed D1 and D2, which drive ClpB’s chaperone activity. The D1 domain features the ClpB-specific M-domain, which forms a long coiled-coil (5) and is essential for protein disaggregation (6, 7). Like other type II AAA+ ATPases, ClpB forms a double-ring structure, with six copies of the D1 and D2 domains each making up a homohexamer ring (5, 8). Although ClpB shares similar quaternary structure with ClpA (9), ClpC (10), and the single-ring ClpX (11) and HslU (12, 13) AAA+ ATPases, which function as the protein unfoldase components of energy-dependent proteases, ClpB does not associate with a chambered peptidase to degrade proteins. Instead, ClpB cooperates with the cognate Hsp70 system (DnaK/GrpE) in a species-specific manner (14, 15) to recover functional protein from aggregates (16–18).

The prevailing model suggests that DnaK/GrpE targets the ClpB motor activity to aggregates (19, 20), which is consistent with an upstream role of the DnaK system in protein disaggregation (21–23). Once targeted, ClpB disaggregates protein aggregates by extracting unfolded polypeptides (24) and threading them through the ClpB hexamer ring (21, 25). In support of a direct ClpB–DnaK/GrpE interaction, it was reported that ClpB interacts with DnaK via the ClpB M-domain (15, 26). Notably, replacing the M-domain of bacterial ClpB with that of its yeast homolog Hsp104 switched the specificity of the chaperone system so that ClpB now cooperated with the eukaryotic Hsp70/40 system and vice versa (7, 27). The role of the M-domain in mediating DnaK/GrpE interaction is consistent with the M-domain being on the outside of the ClpB hexamer (5, 8), but incompatible with the previously proposed structure of yeast Hsp104 (28, 29), which placed the M-domains on the interior or intercalated between subunits.

Although we recently showed that ClpB and Hsp104 share a similar 3D structure (30), functional differences exist (19, 31–33). Moreover, it remains unclear how ClpB exerts the ATP power stroke to thread substrates through the ClpB hexamer ring, and how ClpB unfolds substrates that are typically much larger than the ClpB hexamer itself. The latter might involve ClpB deoligomerization of a substrate-bound chaperone complex (15, 25, 34), or perhaps other mechanical activities that remodel aggregates prior to substrate translocation (5, 35).

Here, we combine structural and biochemical studies to provide mechanistic insights into the functional role of ClpB in protein disaggregation. Notably, we demonstrate using a disulfide cross-linked ClpB variant that deoligomerization of the ClpB hexamer is not required for protein disaggregation. Furthermore, we found that the active conformation of the D2 substrate-translocating pore loop is controlled in cis by the nucleotide state of the same D2 domain, and demonstrate the existence of a conserved but structurally distinct intersubunit-signaling pathway found in diverse AAA+ ATPases. Our findings suggest that AAA+ machines may utilize a common mechanism to harness the energy of ATP binding and hydrolysis to fuel their diverse biological activities.


The authors declare no conflict of interest.

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2.35-Å resolution (form I), the apo structure could not be resolved beyond 4 Å despite extensive efforts to improve the diffraction quality of those crystals (Table S1).

As anticipated, the crystal structures of the isolated D2 domain resemble that of the D2 domain in the full-length protein (5) and form a clamp around the nucleotide (Fig. 1A). Surprisingly, despite lacking ATPase activity, all six copies of the nucleotide-bound conformer unequivocally showed bound ADP in simulated annealed omit maps (Fig. 1A), resulting from either spontaneous ATP hydrolysis or presence of ADP in commercial ATP preparations. Indeed, it was previously reported that the ClpB D2 domain binds ADP two orders of magnitude more tightly than ATP (37). The higher affinity for ADP could explain the bound ADP structure of the apo structure (Fig. S2A and B; also, compare Fig. S2C with D).

**Fitted CryoEM Structure of the D2 Hexamer Ring.** To examine the structure of the D2 subunit interface in the ring assembly, we fitted the X-ray structures of the D2-ADP and apo D2 domain into the mass densities that were previously attributed to the D2 ring in the cryoEM reconstructions of full-length ClpB (8). The resulting atomic structure fits are fully compatible with the cryoEM maps (Fig. 1B and Figs. S3 and S4). Our new fits differ from our previous ClpB–AMPPNP model (8) by a 17° rotation along an axis that is almost perpendicular to the sixfold axis of the ClpB hexamer and a 2.5-Å translation from its center of mass for the D2-ADP hexamer, and an 8° rotation and a 2.8-Å translation for the apo D2 hexamer. Notably, in the D2-ADP hexamer structure, the pore loops line the axial channel (Fig. 1B and Fig. S3), which is consistent with their proposed role in substrate translocation (21). The α/β domain of one D2 subunit and the small α-helical domain of its neighbor contribute primarily to the subunit interface, which is similar to the subunit arrangement in the recently determined crystal structure of the ClpC hexamer complex (10) (Fig. S5).

To confirm our hexamer model, we used structure-based mutagenesis to introduce pairs of cysteines at the D2 domain interface in full-length ClpB (Fig. 2A). The rationale behind this approach was to engineer cysteine pairs, which could potentially form a disulfide bond between neighboring subunits to generate a cross-linked ClpB hexamer. It is entirely fortuitous that *Thermus thermophilus* ClpB lacks cysteines in its amino acid sequence, allowing us to introduce unique cysteine pairs at different locations. Three disulfide-forming ClpB variants were engineered, namely: ClpB<sub>R576C/A821C</sub>, ClpB<sub>L581C/R726C</sub>, and ClpB<sub>576C/R810C</sub> as well as a single-cysteine mutant (ClpB<sub>R576C</sub>) and one that featured a mismatched cysteine pair (ClpB<sub>L581C/R726C</sub>) (Fig. 2A).

Upon oxidation, all three disulfide-forming ClpB variants formed discrete cross-linked oligomers ranging from dimers to hexamers (Fig. 2B and C). As anticipated, our single- and mismatched-cysteine mutants remained monomeric or only formed dimers (Fig. 2B). Notably, ClpB<sub>R576C/A821C</sub> formed stable oligomers with the cross-linked ClpB hexamer as the final product (Fig. 2C and D and Fig. S6). Cross-linked oligomers were also observed in the absence of nucleotide (Fig. S7A), although ATP facilitated the formation of cross-linked hexamers, which is consistent with its known role in stabilizing the ClpB oligomer (38, 39). We found that our crosslinked ClpB<sub>R576C/A821C</sub> hexamer, like wild-type ClpB, has ATPase activity, which is further stimulated by θ-casein (Fig. 2E) and, most remarkably, cooperated with DnaK/GroES/GrpE to rescue heat-aggregated protein substrates (Fig. 2F and G). The ability to recover functional protein was not caused by dissociation of the cross-linked hexamer, because ClpB<sub>R576C/A821C</sub> remained cross-linked even after the refolding assay was complete (Fig. S7B).

To ensure that the cross-linking procedure did not alter the structure or function of ClpB, we generated ClpB variants in the ClpB<sub>R576C/A821C</sub> mutant background by changing Leu396, Leu460, or both, to alanine. We previously showed that mutations of Leu396 and Leu460 reduced ClpB's chaperone activity, presumably by destabilizing the position of the M-domain (5). Consistently, mutating Leu396 and Leu460 to alanine impaired the protein disaggregation activity of cross-linked ClpB<sub>R576C/A821C</sub> to a similar extent, which was completely abolished when both leucine mutations were present (Fig. 2H).

Taken together, our findings suggest that the cross-linked ClpB<sub>R576C/A821C</sub> hexamer is structurally and functionally similar to the wild-type ClpB hexamer, and, perhaps more importantly, that ClpB deoligomerization is not required for protein disaggregation.
Arg747 Is a Critical Trans-Acting Element Required for ATP Hydrolysis.

The structure of the ClpB D2 hexamer provides the structural framework to examine how ATP binding and hydrolysis are coupled to conformational changes that drive protein disaggregation. It is widely accepted that the ATPase activities in the ClpB ring complex are regulated by conserved structural elements, which include the cis-acting Walker A and B and sensor 1 and 2 motifs, and the trans-acting arginine-finger (Arg-finger) residue (6, 39–41). It has been suggested that Arg747 is the Arg-finger residue in T. thermophilus ClpB D2 (41, 42). In our fitted D2 hexamer structure, Arg747 is found at the domain interface and contacts the bound nucleotide in the neighboring subunit (Fig. 3A). To confirm whether Arg747 is the trans-acting Arg-finger residue, we mutated Arg747 to alanine in a full-length ClpB mutant background in which the catalytic glutamate (Glu271) of the first Walker B motif had been additionally mutated to prevent ATP turnover by the D1 domain. We found that the R747A mutation nearly abolished the remaining ATPase activity of ClpBE271A (Fig. 3B), whereas mutating a close-by arginine (Arg741) to alanine did not, confirming that Arg747 is the Arg-finger residue in D2 (41), which is required for the ATPase activity in the D2 ring.

Structural Basis for the Allosteric Mechanism of ATP Binding and Hydrolysis in ClpB.

Structural comparison of the nucleotide-bound and -free structures showed that the D2 pore loop is ordered when ADP is bound (Fig. 1A and Fig. S2D and E), but disordered in the absence of nucleotide (Fig. S2A and C), suggesting that the pore loop conformation is coupled to the nucleotide state of the cis subunit.

In our structure, the D2 pore loop is stabilized by a β-hairpin (residues 688–699), which buttresses the D2 pore loop (Fig. 3C). To examine the functional role of this hairpin, we either deleted the hairpin loop (ClpBΔ691–695) or replaced the loop with three glycines (ClpBΔ690–696/ΔGGS). We found that deleting or replacing the loop impaired ClpB’s ATPase and chaperone function to a
similar extent as the Y643A mutation (Fig. 3 D–F), which nearly abolished substrate translocation through the ClpB hexamer (21). Our structure also shows that His693 in the β-hairpin loop forms stacking interaction with Tyr646 of the pore loop (Fig. 3C). Notably, mutating His693 to alanine significantly reduced the ATPase activity to 43% of wild-type level (Fig. 3D), concomitant with a small but measurable impact on protein disaggregation (Fig. 3 E–F).

Closer inspection further revealed that the β-hairpin extends from a network of residues, which are part of a signaling cascade responsible for sensing the nucleotide state of the neighboring subunit in the m-AAA protease (43), a distantly related AAA+ machine. An important element of this intersubunit-signaling (ISS) motif is an aspartic acid residue, which, in the m-AAA protease, prevents a subunit from hydrolyzing ATP if the neighboring subunit is also in the ATP-bound state (43). In ClpB, this residue is Asp685 (Fig. 3C). Notably, mutating Asp685 to alanine not only impaired ClpB’s ability to turn over ATP (Fig. 3D), but also significantly reduced its ability to recover heat-aggregated protein substrates (Fig. 3E and F), indicating that the ISS motif is conserved and has similar function in diverse AAA+ machines.

Discussion

Bacterial ClpB and its eukaryotic homologs are ATP-dependent protein disaggregases, which have the remarkable ability to rescue stress-damaged proteins from a previously aggregated state. Here, we have presented the complete structure of the ClpB D2 domain, featuring an intact D2 pore loop, which couples the ATP power stroke to substrate translocation in cis. Moreover, we demonstrate that deoligomerization of the ClpB hexamer is not required for protein disaggregation, because our engineered, disulfide cross-linked ClpB hexamer is fully functional in recovering active protein from aggregates.

Although AAA pore loops can differ in length and sequence (Fig. S8), they feature a conserved Φ–Xxx–Gly tripeptide motif (21, 44) and share a common function in substrate translocation. In our structure, the conformation of the D2 pore loop is controlled by the cis subunit that senses the nucleotide state of the neighboring subunit through the Arg-finger, Arg747, and transmits this signal to the ISS motif via Asp685 (Fig. 4). From there, the signal traverses to the Walker B motif through helix D9, and to the D2 pore loop via the β-hairpin loop that includes His693. We propose that this signaling network is crucial to sense the nucleotide state in the adjacent subunit and to reset the nucleotide cycle in the ClpB ring following ATP hydrolysis.
The existence of an ISS network that regulates ATP hydrolysis in diverse AAA+ ring complexes is also consistent with a sequential ATP-hydrolysis mechanism proposed for ClpB (45, 46) and Hsp104 (47), with four out of six subunits in the ClpB homohexamer occupied by nucleotides at any one time (46). This model is similar to the staircase mechanism proposed for the T7 gene 4 ring helicase (48), and is consistent with the nucleotide occupancy observed in the crystal structure of an engineered, covalently linked ClpX hexamer (11). In our model (Fig. 4), the unfolded polypeptide is bound to ClpB in the ATP-bound state, which displays the highest substrate-binding affinity (8, 36). ATP hydrolysis triggers substrate translocation along the hexamer axis. Once hydrolyzed, release of ADP results in substrate dissociation from the cis subunit, concomitant with binding of the unfolded polypeptide by the ATP-bound neighboring subunit. The cycle repeats itself until the substrate is fully translocated through the hexamer ring.

Finally, the conservation of the ISS motif in diverse AAA+ ATPases also suggests that similar structural elements may drive the ATP power stroke in other AAA+ machines, and support a common, ATP-fueled mechanism underlying the distinct cellular activities of diverse AAA+ ATPases.

Materials and Methods


The nucleotide-bound D2 complex was crystallized by the CryoEM Fit. The crystal structure of D2-ADP was superimposed onto the D2 domain of the previous ClpB-AMPNP hexamer fit (8). This arrangement resulted in steric clashes between neighboring D2 pore loops. Therefore, the fit of the D2 monomer was adjusted manually, keeping the D2 hexamer ring within the boundary of the cryoEM density (EMD-1244). The apo D2 domain was fitted similarly into the apo ClpB cryoEM density (EMD-1241).

Disulfide Cross-Linking. Engineered cysteine-containing ClpB mutants (1.5 mg/ml in 40 mM Tris-HCl pH 7.5, 100 mM KCl, 10% glycerol were incubated at 55 °C in the presence of 5 mM ATP, 5 mM MgCl2, 10 μM CuCl2, and 10 μM 1,10-phenanthroline for 20 min unless indicated otherwise. Reactions were quenched by adding 20 mM (final) EDTA. Formation of disulfide bonds was followed by nonreducing 3–8% Tris-acetate gradient PAGE.

Analytical Size-Exclusion Chromatography. ClpB and ClpBSeMet/AED2C1 were analyzed on a Superdex 200 10/300 size-exclusion column prequillibrated with 40 mM MOPS-NaOH pH 7.5 and 100 mM KCl.

ATPase Assay. ATPase activities were measured with or without 0.1 mg/ml λ-casein at 50 °C in 40 mM MOPS-NaOH pH 7.5, 150 mM KCl, 10 mM MgCl2, using 0.1–0.5 μM ClpB (monomer), 10 mM ATP, and a coupled ATP-regenerating system consisting of 0.2 μM pyruvate kinase, 0.5 μM lactate dehydrogenase, 0.5 mM phosphoenolpyruvate, and 0.2 mM NADH. ATP-hydrolysis rates were calculated from the oxidation of NADH, which was monitored by absorbance at 340 nm.

Protein Disaggregation Assays. Recombinant Bacillus stearothermophilus α-glucosidase (0.2 μM) was heat-denatured in refolding buffer A (25 mM HEPES–NaOH pH 7.5, 150 mM KCl, 10 mM MgCl2, 5 mM ATP) at 73 °C for 12 min in the presence of 0.2 μM DnaKJ-DaFα complex (trimmer) and 0.1 μM GrpE (monomer). The aggregate was immediately diluted 20-fold into refolding buffer containing 0.3 μM ClpB (monomer), 0.2 μM DnaKJ-DaFα complex (trimmer), and 0.1 μM GrpE (monomer), and incubated at 55 °C for 40 min. To measure the recovered α-glucosidase activity, p-nitrophenyl glucopyranoside was added to a final concentration of 2 mM. The reaction mixture was incubated at 55 °C for 20 min and stopped with 0.4 M (final) sodium carbonate. The amount of p-nitrophenol released was measured by absorbance at 400 nm.

Purified enhanced GFP (EGFP, 3 μM) was heat-denatured in refolding buffer A at 80 °C for 10 min in the presence of 0.2 μM DnaKJ-DaFα complex (trimmer) and 0.1 μM GrpE (monomer). The aggregate was immediately diluted 12-fold into refolding buffer A containing 0.2 μM DnaKJ-DaFα complex (trimmer), and 0.1 μM GrpE (monomer). The amount of refolded EGFP was monitored continuously at 25 °C for 40 min using a L555 fluorescence spectrometer at an excitation wavelength of 488 nm and emission wavelength of 510 nm.

Recombinant firefly luciferase (1 μM) was heat-denatured in refolding buffer B (25 mM MOPS–NaOH pH 7.5, 150 mM KACo, and 10 mM Mg(OAc)2) at 42 °C for 30 min in the presence of 5 mM ATP and 1 μM DnaKJ-DaFα complex (trimmer) and 0.1 μM GrpE (monomer). The aggregate was immediately diluted 12-fold into refolding buffer B containing 1 μM ClpB (monomer), 1 μM DnaKJGrpE (monomer), and an ATP-regenerating system consisting of 5 mM ATP, 20 mM phosphoenolpyruvate, and 0.2 μM pyruvate kinase. Reactions were incubated at 25 °C for 2 h, and the amount of recovered enzymatic activity was measured at 560 nm after addition of 0.1 μM luciferin.

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Supporting Information

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Fig. S1. Sequence alignment of the analogous D2 domains of ClpA, ClpB, and ClpC from *Thermus thermophilus* (Tt), *Escherichia coli* (Ec), and *Bacillus subtilis* (Bs). Conserved residues are highlighted in green and similar residues in cyan. Secondary structure elements are illustrated and labeled. The asterisk marks the Arg-finger residue, Arg747, in *T. thermophilus* ClpB.

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Translocating pore loop

Walker A

Walker B

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Substrate-
Fig. S2. (A) Crystal structure of the apo D2 domain (PDB entry code: 4FCT). The location of the D2 pore loop, which is disordered in the apo structure, is indicated by a dashed line. (B) Superposition of the apo (magenta, PDB entry code: 4FCT) and ADP-bound (gold, PDB entry code: 4FCV_A) D2 monomers. Structures were superimposed through the Cα atoms of the α/β domain (rmsd of 0.69 ± 0.03 Å). The relative orientation of the large α/β and small α-helical domain is different between the ADP-bound and apo states, and involves a 12.6° rotation and 3.8 Å translation of the α-helical domain in the pair shown. (C) Section of the simulated-annealed composite omit map in the crystal structure of the apo D2 domain (PDB entry code: 4FCT). The map is contoured at 1.0σ level. (D) Section of the simulated-annealed omit map in the crystal structure of the ADP-bound D2 domain (form II, PDB entry code: 4FCV_A) with the D2 pore loop (pink) omitted and the dataset truncated to 4 Å resolution. The map is contoured at the 1.0σ level. (E) Superposition of the four ordered (magenta, PDB entry code: 4FCV_A; dark cyan, PDB entry code: 4FCV_B; gray, PDB entry code: 4FCV_C; brown, PDB entry code: 4FCW_A) and (F) the two partially ordered pore loop conformations (green, PDB entry code: 4FCW_B; blue, PDB entry code: 4FCW_C) seen in the six independent representations of the X-ray structure of the D2-ADP domain. The four ordered loops have rmsd of 0.92 ± 0.28 Å over 14 Cα atoms.
Fig. S3. Stereo diagram of top-down, side, and cutaway side views of the fitted cryoEM structure of the D2-ADP hexamer ring. Fit was obtained by docking the X-ray structure of the isolated D2-ADP domain into the cryoEM reconstruction of the full-length ClpB8227A:6666A–ATP hexamer (EMDB-1244) (1). Neighboring subunits are colored differently for clarity. ADP is depicted as stick model.

Fig. S4. Stereo diagram of top-down, side, and cutaway side view of the fitted cryoEM structure of the apo D2 hexamer ring. Fit was obtained by docking the X-ray structure of the isolated apo D2 domain into the cryoEM reconstruction of the full-length apo ClpB hexamer (EMDB-1241) (1). Neighboring subunits are colored differently for clarity.

Fig. S5. Juxtaposition of the homohexamer structures of (A) the ClpB D2-ADP ring (this work) and (B) the ClpC D2 ring, which was derived from the X-ray structure of the MecA–ClpC hexamer complex (PDB entry code: 3PX1) (1). To improve the diffraction quality of the crystals, a total of four loops were deleted (two in D2) in ClpC. Hence, although the arrangement of D2 subunits is similar in ClpB and ClpC, the ClpB D2 hexamer ring appears more compact. Neighboring subunits are colored in different hues for clarity. The symbols indicate the sixfold and twofold symmetry axes present in the structures.


Fig. S6. Size-exclusion chromatograms of wild-type (wt) ClpB, crosslinked (xl) ClpB*R576C/A821C, and ClpB*R576C/A821C*. Non-crosslinked ClpB*R576C/A821C mostly exists as a smaller oligomer (tetramer), although some hexamers are also present.

Fig. S7. (A) Apo ClpB*R576C/A821C forms high-molecular weight, cross-linked oligomers, albeit less efficiently than in the presence of nucleotide. Time course of the catalyzed cross-linking reaction of ClpB*R576C/A821C in the absence of ATP. M, marker; T, disulfide cross-linked ClpB*R576C/A821C hexamer after 10 min in the presence of ATP. (B) The cross-linked ClpB*R576C/A821C hexamer remains cross-linked during the protein disaggregation reaction. Samples of the α-glucosidase disaggregation reactions before (+) and after (*) refolding by DnaKJ/GrpE only (KJE), and in the presence of the wild-type ClpB bichaperone system (KJE + WT), or cross-linked ClpB*R576C/A821C plus DnaKJ/GrpE (KJE + XL). M, molecular weight markers.
Fig. S8. Structure-based survey of the substrate-translocating pore loop (pink) and flanking secondary structure elements (cyan, β-sheet; gray, α-helix) in diverse AAA+ ATPases. *Thermus thermophilus* (Tt) ClpB–D2 (this work), *Escherichia coli* (Ec) ClpA–D2 (PDB entry code: 1R6B) (1), EcClpX (PDB entry code: 3HWS) (2), *Escherichia coli* (Ec) HslU (PDB entry code: 1E94) (3), *Mus musculus* (Mm) p97–D2 (PDB entry code: 3CF0) (4), *Thermotoga maritima* (Tm) FtsH (PDB entry code: 3KDS) (5), *Aquifex aeolicus* (Aa) NtrC1 (PDB entry code: 3M0E) (6), TtClpB–D1 (PDB entry code: 1QVR) (7), EcClpA–D1 (PDB entry code: 1R6B) (1), Mmp97–D1 (PDB entry code: 1E32) (8).


### Table S1. Data collection and refinement statistics

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<td>15.9(8.3)</td>
<td>18.8(4.3)</td>
<td>10.6(5.1)</td>
<td>18.5(5.9)</td>
</tr>
<tr>
<td>Completeness (%)</td>
<td>99.2(99.8)</td>
<td>99.7(99.9)</td>
<td>99.0(98.8)</td>
<td>98.0(99.4)</td>
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<tr>
<td>Redundancy</td>
<td>6.9(6.2)</td>
<td>6.2(4.8)</td>
<td>7.4(7.6)</td>
<td>6.2(4.4)</td>
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<tr>
<td>Refinement</td>
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<tr>
<td>Total no. of reflections</td>
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<td>300,635</td>
<td>90,324</td>
<td>16,702</td>
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<td>23,566</td>
<td>46,053</td>
<td>12,206</td>
<td>2,694</td>
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<tr>
<td>Rwork/Rfree (%</td>
<td>28.2/29.9</td>
<td>23.5/29.1</td>
<td>23.3/26.0</td>
<td>29.1/29.2</td>
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<tr>
<td>No. of atoms</td>
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<tr>
<td>Protein</td>
<td>7,392</td>
<td>7,368</td>
<td>7,446</td>
<td>2,348</td>
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<td>Water</td>
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<tr>
<td>ADP</td>
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<td>81</td>
<td>81</td>
<td>0</td>
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<tr>
<td>B-factors</td>
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<td>66.5</td>
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<td>42.0</td>
<td>73.8</td>
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<td>RMSD</td>
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<tr>
<td>Bond lengths</td>
<td>0.013 Å</td>
<td>0.009 Å</td>
<td>0.009 Å</td>
<td>0.006 Å</td>
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<td>Bond angles</td>
<td>1.51°</td>
<td>1.27°</td>
<td>1.19°</td>
<td>0.93°</td>
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<tr>
<td>Ramachandran plot</td>
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<tr>
<td>Favored (%)</td>
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<td>92.4</td>
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<td>85.6</td>
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<td>Allowed (%)</td>
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<td>6.7</td>
<td>11.2</td>
<td>12.8</td>
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<td>Generous (%)</td>
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<td>Disallowed (%)</td>
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<td>4FD2</td>
<td>4FCW</td>
<td>4FCV</td>
<td>4FCT</td>
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</table>

Values in parentheses are for the highest-resolution shell. \( R_{sym} = \sum_{hkl} |I(hkl)\|/|\sum_{hkl} I(hkl)|\), where \(|I(hkl)|\) is the mean of the symmetry-equivalent reflections of \(I(hkl)\). The \( R_{free}\) was calculated with 5% of randomly selected unique reflections for form I and form II, and 10% for apo.