**Structural investigation of a chaperonin in action reveals how nucleotide binding regulates the functional cycle**

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Chaperonins are ubiquitous protein assemblies present in bacteria, eukaryota, and archaea, facilitating the folding of proteins, preventing protein aggregation, and thus participating in maintaining protein homeostasis in the cell. During their functional cycle, they bind unfolded client proteins inside their double ring structure and promote protein folding by closing the ring chamber in an adenosine 5′-triphosphate (ATP)–dependent manner. Although the static structures of fully open and closed forms of chaperonins were solved by x-ray crystallography or electron microscopy, elucidating the mechanisms of such ATP-driven molecular events requires studying the proteins at the structural level under working conditions. We introduce an approach that combines site-specific nuclear magnetic resonance observation of very large proteins, enabled by advanced isotope labeling methods, with an in situ ATP regeneration system. Using this method, we provide functional insight into the 1-MDa large hsp60 chaperonin while processing client proteins and reveal how nucleotide binding, hydrolysis, and release control switching between closed and open states. While the open conformation stabilizes the unfolded state of client proteins, the internalization of the client protein inside the chaperonin cavity speeds up its functional cycle. This approach opens new perspectives to study structures and mechanisms of various ATP-driven biological machineries in the heat of action.

**INTRODUCTION**

Chaperones, proteins that assist folding and stabilization of other proteins, are essential for cellular homeostasis and viability. Chaperonins, a particular class of chaperones also noted hsp60 in the following, form a giant barrel-like structure composed of two back-to-back stacked rings. While group I chaperonins, such as bacterial GroEL, require a co-chaperonin lid (GroES) to cap the ring chamber, group II chaperonins, present in the eukaryotic cytosol and in archaea, are composed of subunits with a built-in helical protrusion which closes the folding chamber. Biochemical studies have provided limited structural information on key steps of the chaperonin functional cycle and mechanism of protein folding (1–4). Hsp60 chaperonin undergoes large-scale adenosine 5′-triphosphate (ATP)–dependent conformational changes to promote folding of the client protein within a large chamber. Structures of open and closed states (5–9) have provided static snapshots of steps along the cycle, but the conformational dynamics of the active ATP-fueled chaperone, as well as the interactions with client proteins being refolded, have remained elusive. Nuclear magnetic resonance (NMR) spectroscopy is the method of choice to study proteins in solution and is, in principle, able to observe conformational changes, interactions, and dynamics under functional conditions. However, the site-specific observation of proteins as large as chaperonin, 1 MDa, remains a significant challenge due to the extensive line broadening of NMR signals in large proteins. Furthermore, retaining the protein in a functional state throughout a structural investigation is complicated by the rapid consumption of ATP, thereby leading to inactivation and inhibition of the ATP-dependent machinery.

To overcome these methodological limitations, we have developed an ATP regeneration system directly inside an NMR tube to keep the ATP concentration constant and to prevent the accumulation of adenosine 5′-diphosphate (ADP), thereby maintaining the chaperonin in a steady working condition powered by ATP hydrolysis. We furthermore elaborated a combined methyl-specific labeling approach to probe the different conformational states of the chaperonin and its client proteins. This combination of methods allows us to probe, on the structural level, the different states involved in the functional cycle of active hsp60 processing client proteins while the chaperonin is being powered by ATP hydrolysis (10, 11). We provide quantitative measures of the thermodynamics and kinetics of the transitions between the open and closed conformations and reveal how the chaperonin affects the folding/unfolding equilibrium of the client protein. Specifically, we find that hsp60 acts as a holdase of unfolded proteins until activation by ATP. ATP binding closes the chaperonin cavity, while the ATP hydrolysis rate controls the timing of the closed state, during which encapsulated proteins can refold for a period of ~17 s. After ATP hydrolysis, the chamber reopens during the ADP residence time (~10 s), which allows release of folded protein and client proteins to be reloaded before the start of a new cycle. Our results reveal how nucleotides and client protein binding regulate the transition between different conformations populated during the hsp60 functional cycle. We anticipate this work to be a starting point to obtain insights into the mechanisms of various chaperones. Furthermore, our approach opens new perspectives for the study of structures and mechanisms of ATP-powered biomolecular machines.

**RESULTS**

**ATP binding closes the chaperonin cavity**

We have chosen to study the conformations, nucleotide binding, and client protein interaction of a group II chaperonin from the
hyperthermophilic *Pyrococcus horikoshii* (12). This hsp60 chaperonin is a homohexadecameric assembly (Fig. 1, A and B), that is, a simplified mechanistic model of its eukaryotic heterohexadecameric counterpart. Furthermore, with an optimal functional temperature above 60°C, where NMR relaxation properties are more favorable, site-specific NMR information can be obtained at the level of individual residues under physiologically relevant conditions. However, studying a 1-MDa protein assembly is challenged by the low sensitivity and resolution associated with rapid NMR spin relaxation (13, 14). By combining selective valine and methionine methyl labeling (15, 16) with methyl–transverse relaxation optimized spectroscopy (TROSY) experiments (17, 18), we have obtained high-quality NMR spectra, allowing us to probe the structure and dynamics at 62 individual sites evenly dispersed throughout the structure of the chaperonin (Fig. 1, B and C, and figs. S1 and S2). In nucleotide-free or ADP-bound states, chaperonins are in an open state allowing the access of client proteins to the cavity (Fig. 1A) (19–22). However, the event that triggers the open-to-closed transition (Fig. 1B) remains a matter of discussion (4, 19, 23–26). Having highly resolved NMR spectra, we probed the conformational states of chaperonin in the presence of different nucleotide ligands. Chemical shift perturbations induced by the binding of ADP and a nonhydrolyzable ATP analog (App-NHp) revealed that M159, located in the vicinity of the nucleotide binding site, exhibits distinct peak positions in the apo and nucleotide-bound states and, thus, directly reports on the occupancy of the binding site (Fig. 1C). Upon binding of App-NHp, however, the signals of M275 and M279 located in the lid are significantly shifted (fig. S3C), revealing a global conformational change to the closed state (4). In contrast, binding of ADP does not affect the signals of these two methionines, confirming that the ADP-bound state, just like the apo state, is in an open conformation (fig. S3B). Negative staining electron microscopy (EM) analysis of the NMR sample (fig. S4) confirmed that App-NHp binding induces the closed conformation. Together, these results show that ATP binding triggers the transition from the open to the closed conformation of the ring, while ADP binding does not affect the closed conformation.

**The open state of chaperonin acts as a holdase of unfolded proteins**

Hen egg white lysozyme (HEWL; 14.5 kDa) and malate synthase G (MSG; 82 kDa) were selected as model proteins to observe the interaction of hsp60 with two client proteins with a significant size difference. At 65°C, a temperature close to the optimal conditions for hsp60 activity, HEWL features two sets of NMR resonances corresponding to the unfolded and folded states, which interconvert on a time scale of 350 ms (fig. S5). The addition of chaperonin shifts only the peaks of the unfolded form of HEWL, showing that hsp60 interacts exclusively with the unfolded form (Fig. 2A and fig. S5). Quantitative analysis of the titration data reveals a dissociation constant of $1.6 \pm 0.4 \mu M$, with a binding stoichiometry of one unfolded HEWL per chaperonin cavity. NMR-based thermal unfolding studies of HEWL revealed that the presence of hsp60 decreases the melting temperature by 5°C (Fig. 2B and fig. S5), confirming the stabilization of the unfolded state by the chaperonin. The exclusive interaction of hsp60 with the unfolded state is furthermore confirmed by the observation that the translational diffusion coefficients of the unfolded states of HEWL and MSG are shifted to the one of the nucleotide-bound state, just like the apo state, is in an open conformation (fig. S3B). Negative staining electron microscopy (EM) analysis of the NMR sample (fig. S4) confirmed that App-NHp binding induces the closed conformation. Together, these results show that ATP binding triggers the transition from the open to the closed conformation of the ring, while ADP binding does not affect the closed conformation.

**Fig. 1.** Methionine-directed methyl NMR allows the identification of the nucleotide binding site and the different chaperonin conformational states. (A) Model of the chaperonin structure in the open conformation obtained by homology modeling from Protein Data Bank (PDB 3IZH). (B) Magnified view of one hsp60 monomer in open (gray) and closed (red) conformations (obtained by homology modeling from PDB 1Q3R). The methyl groups of methionine residues are represented by green spheres, and the nucleotide in the binding site is represented in yellow. (C) 2D $^1H$–$^{13}C$-methyl-TROSY spectrum of a 125 μM U-[$^2H$, $^{15}N$], Met-$^{13}CH_3$ hsp60 sample with the assignment of the methionine residues obtained by combination of a mutation-based approach (fig. S1) with detection of NOEs between methyl groups (fig. S2). The inserts focus on cross peaks of M159 and M279 in the apo state (black) and in the presence of 2 mM ATP analog App-NHp (red, closed state) or ADP (blue, open state).
Client protein accelerates the functional cycle of hsp60

To study the system in action under steady-state conditions, without accumulation of the competitive inhibitor ADP (27), we implemented an enzymatic system directly inside the NMR sample to ensure continuous rapid regeneration of ATP from ADP and phosphoenolpyruvate (PEP; Fig. 3A). This system keeps the ATP concentration constant, prevents ADP accumulation, and keeps the chaperonin catalytically active at a constant rate (Fig. 3, B and C). Moreover, interleaved NMR data collection of this complex reaction mixture allows to simultaneously monitor the state of the chaperonin and client protein, as well as the concentrations of ATP, ADP, and PEP. The distinct sets of methionine NMR resonances report on the open/closed conformational state (M275 and M279) and type of nucleotide bound (M159; Figs. 1 and 3D), while the Ile, Leu, and Val resonances of HEWL report on the equilibrium of folded and unfolded states of the client protein (Fig. 4A and fig. S8). The population of folded HEWL of ca. 50% in the presence of chaperonin alone increased to 75% when powered by ATP and the ATP regeneration system, demonstrating that the active hsp60 is able to enhance the client protein refolding (Fig. 4B). Furthermore, the comparison of ATPase activity of hsp60 with and without HEWL, quantified from the decay of PEP signals, reveals that the presence of client protein enhances the ATPase activity of hsp60 by 40% from 24.7 ± 0.5 to 36.0 ± 1.5 ATP/min per hsp60 (Figs. 3 and 4C and fig. S9). These observations indicate a mutual activation of protein refolding and cycling of the chaperonin between different functional states.

Fig. 2. NMR characterization of the interaction between client protein and hsp60. (A) Sum of chemical shift changes of five HEWL methyl signals (V2, I88, V92, V109, and I124) showing a significant shift and no overlap. The inset shows the NMR signal of valine residues of HEWL recorded at 75°C at a constant concentration (31 μM), and an increasing concentration of perdeuterated hsp60 assembly (0 to 97 μM), a.u., arbitrary units. (B) Evolution of the folded population of the HEWL between 50° and 75°C without (black) and with (red) hsp60 (two equivalents of HEWL per chaperonin). The melting temperatures (Tm) are indicated by vertical dotted lines. (C) Translational diffusion properties of hsp60 alone (blue), unfolded HEWL alone (black), and in the presence of hsp60 (red) characterized by diffusion-ordered NMR spectroscopy. Derived diffusion coefficients at 65°C (hsp60: 5.8 ± 0.3 × 10⁻⁷ cm²/s; unfolded HEWL with hsp60: 4.4 ± 1.2 × 10⁻⁷ cm²/s; unfolded HEWL alone: 30 ± 4 × 10⁻⁷ cm²/s) reveal that only unfolded HEWL is interacting with hsp60. (D) Overlay of the 2D ¹H-¹³C NMR spectra of U-[²H, ¹⁵N], Val-[¹³CH₃]proS, Met-[¹³CH₃] methyl–labeled sample of hsp60 with two equivalents of diamagnetic (black) or paramagnetic labeled HEWL (red). (E) Model of the monomer of hsp60 in the open conformation showing the residues affected by the paramagnetically labeled client proteins. hsp60 sensor loop is colored in green.
Nucleotide binding and hydrolysis control the population of chaperonin conformations

While the chaperonin is being powered by the ATP regeneration system, the NMR reporter of the nucleotide binding state, M159, shows that both the ATP- and ADP-bound states are present in steady state, despite the fact that the concentration of free ADP in solution is negligible (Fig. 4D and fig. S10, A and B). The level of unoccupied binding site is below our detection limit (<3%), showing that the lifetime of nucleotide-free state is negligible compared to the ADP- and ATP-bound states. The reporters of the cavity conformational state (M275 and M279) reveal that there is also a mixture of open and closed states present under these steady-state conditions.
conditions (Fig. 4E and fig. S10, C and D). These independent NMR observables allow the determination of population levels of the ATP-bound (62 ± 3%) and ADP-bound states (38 ± 3%), which quantitatively match the ones of the closed and open conformations, respectively (62 ± 3%, 38 ± 3%; Fig. 4, D and E). These relative population levels of open (ADP-bound) and closed (ATP-bound) states of chaperonin are retained also when client protein is present (fig. S10).

**DISCUSSION**

The chaperonin conformational cycle is driven by ATP; however, its exact role remains a matter of debate (10, 11). A proposed model, in which ATP binding alone is sufficient to close the ring (4, 19, 23, 24), was challenged by studies concluding that it is the ATP hydrolysis that drives the ring closing (3, 25, 26). The quantitative equality of the relative populations of ATP-bound (ADP-bound) states and closed (open) conformations of the chaperonin observed in this study (Fig. 5) implies that ATP binding and chamber closing are intimately linked, and along with the fact that nonhydrolyzable ATP analogs close the hsp60 cavity, these results confirm the view that nucleotide binding, rather than hydrolysis, induces the closure of the ring (Fig. 1B).

This study has also enabled resolution of the cooperativity and conformations of both rings during the adenosine triphosphatase (ATPase) cycle of group II chaperonins. The fact that we observe only two sets of NMR signatures, corresponding to closed/ATP-bound and open/ADP-bound states, leads us to propose that each ring comprising eight subunits acts as a cooperative unit, as proposed earlier (Figs. 3 and 4, D and E, and figs. S9 and S10) (27, 28). This view is corroborated by the absence of mixed conformations within a ring in EM images (fig. S4). Several studies advocated the presence of inter-ring cooperativity in the group II chaperonins, suggesting an alternating mechanism in which one ring is open and the opposite
ring closed (1–3, 29, 30). However, there is also evidence of a non-concerted mechanism, in particular for the eukaryotic chaperonin CCT (chaperonin containing TCP-1) (31, 32). The extreme model with strict anticoopetativity (3)—that is, opening of one ring leading to closing of the adjacent ring—predicts the exclusive presence of half-open/half-closed chaperonin particles, that is, populations consisting of 50% open and 50% closed half-rings at each time point. An alternative model based on partial anticoopetativity includes doubly closed particles in addition to the half-open/half-closed state (33, 34). The population levels obtained directly from our NMR data of the ATP-powered chaperonin in action allow resolution of these conflicting models. The skewed population levels (62.38 ± 3%) necessarily imply the presence of chaperonin particles that have two closed chambers. Taking into account that nucleotide-free states are absent (Fig. 4D), we propose a modified model (2), depicted in Fig. 5, which involves four states along the functional cycle, two of them fully closed and two half-open/half-closed states. The kinetics of ATP processing (Figs. 3 and 4) and the relative populations of states (Fig. 5), detected in our NMR sample, allow determining the average lifetimes of the populated states. With an ATP hydrolysis activity per chaperonin of 36 ATPs/min (Fig. 4C), and assuming that a full cycle consumes 16 ATPs, one catalytic turnover takes 26.7 s. Together with the experimentally observed population levels (Fig. 4, D and E), we can derive that the half-open/half-closed state has a threefold longer lifetime (10.2 s) than the doubly closed state (3.2 s; Fig. 5). The long residence time of the half-open/half-closed state allows the release of the folded proteins and the re-binding of a new unfolded client protein to the chaperonin. In addition, the fact that the ratio of populations in the open and closed conformations equals the ratio of ADP- and ATP-bound states shows that the events following the ATP hydrolysis are subsequently leading to the opening of the chaperonin folding chamber, while the ADP residence time controls the duration of the open state of hsp60 cavity. Together, our results reveal that nucleotide binding, hydrolysis, and release control the conformational transitions between closed and open states and their relative lifetimes. While the open chaperonin conformation stabilizes the unfolded state of client proteins, its binding inside the cavity speeds up the functional cycle of the chaperonin. The implementation of a fully active chaperoning system, being constantly powered by fresh ATP inside the NMR tube, allows studying biomolecular function in action. This work establishes the feasibility to probe these events at the structural level even in systems as large as 1 MDa. We anticipate our approaches to be applicable to a wide range of protein assemblies, including HSPs (heat shock proteins) involved in cancer (35) or in the disaggregations of amyloidogenic proteins (30), as well as other large cellular machines in action such as biological motors or polymerases.

**MATERIALS AND METHODS**

**Production and purification of methyl specifically labeled proteins in *Escherichia coli***

Production and purification of specifically methyl labeled hsp60

*E. coli* BL21-CodonPlus-(DE3)-RIL cells transformed with a pET-11a plasmid encoding the hsp60 from *P. horikoshii* (also known as thermosome) were progressively adapted in three stages over 24 hours to M9/D2O medium containing d-glucose-d5 (2 g/liter; Isotec). As part of our standard labeling protocols, the culture media also contained 15ND4Cl (1 g/liter), but incorporation of 15N in produced samples was not used in this study and these extra 15N spins did not interfere with 13CH3-edited NMR experiments reported here. In the final culture, bacteria were grown at 37°C in M9 medium prepared with 99.85% D2O (Eurisotop). When the optical density (OD) at 600 nm reached 0.6 to 0.7, a solution containing the labeled precursors was added. The precursor solution added for 1 liter of culture medium contained the following: either 100 mg of [βγβ2H5, ε-13C]-l-methionine (Cambridge Isotope Laboratories; CIL) for the production of the U-[1H, 15N], Met-[15CH3] hsp60 sample; or 240 mg of 2-hydroxy-2-[13C]methyl-3-oxo-4,4,4-tri-[1H]butanoate (proS acetolactate-13C, NMR-Bio), 30 mg of l-leucine-d10, and 100 mg of [α,β,γ-2H5, ε13C]-l-methionine (CIL) for the production of the U-[1H, 15N], Met-[13CH3], Val-[13CH3] proS hsp60 sample (15). One hour after the addition of the precursors, hsp60 expression was induced by the addition of isopropyl β-D-1-thiogalactopyranoside (IPTG) to a final concentration of 0.5 mM. The induced culture grew for 4 hours at 37°C before harvesting. Cells were lysed by sonication in buffer containing 25 mM Heps (pH 7.5), 150 mM NaCl, 5 mM MgCl2, and DNase (deoxyribonuclease) (0.01 mg/ml). After the first heat shock step at 80°C, hsp60 was purified using an anion exchange chromatography step (Resource Q, 6 ml, GE Healthcare) followed by a size exclusion chromatography step (HiLoad 16/60 Superdex 200 pg, GE Healthcare). The final yield reached 20 mg/liter of methyl-specific protonated hsp60. The protein was concentrated to final NMR concentration in D2O buffer [20 mM MES (pH 6.5 uncorrected), 100 mM NaCl, 150 mM KCl, and 25 mM MgCl2]. After concentration of the sample, the ATPase activity of hsp60 was tested using a malachite green assay (Sigma-Aldrich), and the oligomerization state of the particle was controlled using an analytical size exclusion column (CE Agilent Bio SEC-5A 500 Å) and negative staining EM.

**Production and purification of hsp60 methionine and valine mutants**

Constructs containing valine-to-alanine or methionine-to-leucine single point mutations were generated by an automated molecular biology platform (RoBioMol, Institut de Biologie Structurale) using an automated polymerase chain reaction–based protocol adapted from the QiikChange site-directed mutagenesis method (36, 37). The library of mutants was expressed in parallel in 25 ml of M9/D2O and labeled on methionine and valine methyl groups following the protocol described above. Purification of hsp60 mutants was done following the protocol described above. The hsp60 mutants were concentrated in D2O buffer [20 mM MES (pH 6.5 uncorrected), 100 mM NaCl, 150 mM KCl, and 25 mM MgCl2] using centrifugal filter devices (Vivaspin, 100,000 MWCO, Vivasience) to a final concentration between 6.25 and 12.5 μM hsp60 (100 to 200 μM monomer).

**Production of U-[1H, 15N], Leu/Val-[13CH3] proS, Ile-[13CH3]H EWL**

*E. coli* BL21-CodonPlus-(DE3) cells transformed with a pET-11a plasmid encoding the EWL (Addgene plasmid #39233) (38) were progressively adapted in three stages over 24 hours to M9/D2O medium containing 15ND4Cl (1 g/liter) and d-glucose-d3 (2 g/liter; Isotec). As part of our standard labeling protocols, the culture media also contained 15ND4Cl (1 g/liter), but incorporation of 15N in produced samples was not used in this study and these extra 15N spins did not interfere with 13CH3-edited NMR experiments reported here. In the final culture, bacteria were grown at 37°C in M9 medium prepared with 99.85% D2O (Eurisotop). When the OD at 600 nm reached 0.6 to 0.7, 2-hydroxy-2-[13C]methyl-3-oxo-4,4,4-tri-[1H]butanoate (proS acetolactate-13C, NMR-Bio) was added at a final concentration of 240 mg/liter 1 hour before induction. Forty minutes later (that is, 20 min before induction), 3,3-[2H2],4-[13C]-2-ketobutyrate...
(NMR-Bio) was added to a final concentration of 60 mg/liter (15).
One hour after the addition of the first precursors, HEWL expression was induced by the addition of IPTG to a final concentration of 2 mM. The induced culture grew for 3 hours at 37°C before harvesting.

HEWL was purified from inclusion bodies (38) and refolded using size exclusion chromatography (39). The final yield reached 5 mg/liter of methyl-specific protonated refolded HEWL. The protein was concentrated using centrifugal filter devices (Vivaspin 15, 5000 MWCO, Vivasicence) in D$_2$O buffer [20 mM MES (pH 6.5 uncorrected), 100 mM NaCl, 150 mM KCl, and 25 mM MgCl$_2$].

**Production of U-$^{2H, 15N}$, Leu/Val-$^{13}$CH$_3$proS, Ile-$^{13}$CH$_3$**

MSG

*E. coli* BL21(DE3) cells were transformed by heat shock with the pET21b bearing the MSG gene. Cells were progressively adapted to E. coli BL21(DE3) cells were transformed by heat shock with the pET21b bearing the MSG gene. Cells were progressively adapted to

90 hours with a 75\textmu M (470 to achieve the highest sensitivity.

The angle of the proton excitation pulses were recorded with an adjusted duration depending on experiments (700 MHz equipped with a 5-mm cryogenically cooled pulsed-field gradient triple-resonance probe. A recycle delay of 1 s was used in all experiments. The diffusion time was set to 100 ms, gradient duration was 2 ms, and recovery delay was set to 0.2 ms, with a total acquisition time of 2 hours for each spectrum. The gradient strength varied from 5 to 42.5 G/cm.

**1$^3$C-edited 2D Exchange Spectroscopy (EXSY) experiments** were acquired to characterize the exchange between folded and unfolded HEWL and to transfer assignment between the two forms. For this purpose, we have modified standard methyl-TROSY experiment (17) by adding a first 90° $^1$H pulse just before proton acquisition to store magnetization along the z axis during the EXSY mixing delay. Then, the magnetization was flipped back in the transverse plane by the addition of a second 90° $^1$H pulse for the detection of $^1$H signal. All the 2D methyl-TROSY-EXSY experiments were acquired on a Bruker Avance III HD spectrometer operating at a $^1$H frequency of 700 MHz equipped with a 5-mm cryogenically cooled pulsed-field gradient triple-resonance probe. A recycle delay of 1 s was used in all experiments. The EXSY mixing times were set to 1, 25, 50, 100, 200, 300, 500, 800, 1000, and 1300 ms, with an average acquisition time of 30 min for each spectrum.

**Assignment of methyl group resonances**

**Assignment of HEWL Ile-$^{\delta_3}$, Val and Leu-proS methyl groups**

Folded HEWL assignment was obtained from the published assignment (36). Unfolded HEWL assignment was transferred from the folded HEWL using a methyl-TROSY-EXSY experiment recorded at 65°C, where both unfolded and folded populations of HEWL can be observed (fig. S5D).

**Assignment of hsp60 Met-$^{\varepsilon}$ and Val-$^{\gamma}$ methyl groups**

Each mutant sample (40 \mu l) was loaded in a 1.7-mm tube, and the NMR experiments were recorded on a Bruker Avance III HD spectrometer operating at a $^1$H frequency of 850 MHz and equipped with a 1.7-mm TCI MicroCryoProbe. The 2D SOFAST-methyl-TROSY NMR experiments were recorded at 75°C with an adjusted duration depending on the final concentration of each mutant (experimental time ranging from 60 to 120 min per sample). For each mutant, spectra (fig. S1) were recorded for the apo state, App-NHp–bound state (1 mM), or ADP-bound state (1 mM). Analysis and comparison of the complete library of mutant spectra allowed the assignment of 33 valines (70%) and 15 methionines (100%). To obtain the missing assignment and confirm the assignment obtained by mutagenesis, a 3D HMQC-NOESY-HMQC experiment was recorded with a 47 \mu M (0.75 mM monomer) U-$^{13}$CH$_3$, Met-$^{13}$CH$_3$, Val-$^{13}$CH$_3$proS sample of hsp60 (fig. S2). Comparison of the NOE cross peaks with the model of the structure of hsp60 allowed to confirm the mutagenesis assignment and to assign 43 valines (92%) and 15 methionines (100%).

**Study of the interaction between client proteins and hsp60**

**Titration of the interaction between HEWL and hsp60**

Spectra were recorded using a different sample for each titration point with a fixed concentration of 31 \mu M U-$^{13}$H, $^{15}$N], Val/Leu-$^{13}$CH$_3$proS, Ile-$^{13}$CH$_3$] (1 mM) HEWL and a variation of the concentration of U-$^{13}$H, $^{15}$N], Met-$^{13}$CH$_3$] hsp60. Each sample (40 \mu l) was loaded in a 1.7-mm tube, and the NMR experiments were recorded on a Bruker Avance III HD spectrometer operating at a $^1$H frequency of 850 MHz.
and equipped with a 1.7-mm TCI MicroCryoProbe. The 2D SOFAST-methyl-TROSY NMR (18) experiments were recorded at 75°C.

**Thermal unfolding of HEWL with and without hsp60**

The reversible thermal (un)folding of HEWL (37) was probed using two samples. HEWL reference sample: 31 μM HEWL U-[1H, 15N], Val/Leu-[13CH3]proS, Ile-[13CH3]81. HEWL with hsp60: 15 μM hsp60 (240 μM monomer concentration) U-[1H, 15N], Met-[13CH3] sample and 31 μM HEWL U-[1H, 15N], Val/Leu-[13CH3]proS, Ile-[13CH3]81 sample (two equivalents of hsp60 per hsp60 particle). Each sample (40 μl) was loaded in a 1.7-mm tube, and the NMR experiments were recorded on a Bruker Avance III HD spectrometer operating at a 1H frequency of 850 MHz and equipped with a 1.7-mm TCI MicroCryoProbe. The 2D SOFAST-methyl-TROSY NMR experiments (18) were recorded from a temperature of 50°C to 75°C with steps of 5°C between 50° and 60°C and steps of 2.5°C between 60° and 75°C. Percentage of folded state was extracted from the averaged ratio between intensity of the signals in the folded and unfolded states for different residues.

**Measurement of the translational diffusion coefficient of the free and hsp60-bound HEWL and MSG**

For HEWL, three samples were used to acquire the 2D [13C, 1H]-DOSY data sets (40) at a temperature of 65°C. Reference sample: 31 μM HEWL U-[1H, 15N], Val/Leu-[13CH3]proS, Ile-[13CH3]81; hsp60 reference sample: 15.5 μM hsp60 (250 μM monomer concentration) U-[1H, 15N], Met-[13CH3]; and hsp60 with HEWL: 15.5 μM hsp60 (250 μM monomer concentration) U-[1H, 15N], Met-[13CH3] and 31 μM HEWL U-[1H, 15N], Val/Leu-[13CH3]proS, Ile-[13CH3]81 sample (two equivalents of HEWL per hsp60 particle).

For MSG, three samples were used to acquire the 2D [13C, 1H]-DOSY data sets. As thermally unfolded MSG is not soluble, the MSG reference spectrum was acquired at 50°C with 31 μM folded MSG U-[1H, 15N], Ile-[13CH3] 81. For the sample containing unfolded MSG bound to hsp60, 15.5 μM hsp60 (250 μM monomer concentration) U-[1H, 15N], Met-[13CH3] was mixed with 31 μM MSG U-[1H, 15N], Ile-[13CH3] 81. MSG was thermally unfolded at 60°C in the presence of hsp60 (two equivalents of MSG per hsp60 particle), and DOSY (Diffusion Ordered SpectroscopY) spectra were acquired at 60°C. The hsp60 reference sample spectrum was acquired at 60°C with a sample containing 15.5 μM hsp60 (250 μM monomer concentration) U-[1H, 15N], Met-[13CH3].

**Paramagnetic labeling of HEWL and MSG for the detection of intermolecular paramagnetic relaxation enhancement (PRE) effects**

Spin labeling of the e-amino groups of the solvent accessible HEWL (Sigma-Aldrich) or MSG lysine residues with OXYL-1-NHS (1-oxy-2,2,5,5-tetramethylpyrrolin-3-carboxylate-N-hydroxysuccimide ester, Toronto Research Chemicals) was carried out following a published protocol (41). Protein was dissolved in the labeling buffer [10 mM Na2CO3 (pH 9.2)] at a concentration of 40 μM. A stock solution was prepared by dissolving 10 mg of OXYL-1-NHS in 100 μl of dimethyl sulfoxide. After addition of a 50-fold molar excess of OXYL-1-NHS over lysine residues to the protein solution, the reaction mixture was incubated for 1 hour at room temperature, followed by 4 hours at 4°C. The excess of spin label was removed by washing the sample with approximately 20 volumes of NMR buffer using centrifugal filter devices (Vivaspin 15, 5000 MWCO, Vivascience). Homogeneity of the lysine labeling was checked by mass spectrometry.

We measured paramagnetic relaxation (PRE) enhancement values, to provide qualitative information on the interactions between HEWL or MSG and hsp60, by comparing peak intensities in 2D SOFAST-methyl-TROSY NMR experiments recorded at 75°C with a 16 μM hsp60 (256 μM monomer concentration) U-[1H] Met-[13CH3], Val-[13CH3]proS in interaction with either 32 μM OXYL-1-NHS–labeled 1H HEWL (Iref)/MSG (Iref) or reference diamagnetic HEWL (Iref)/MSG (Iref).

**NMR study of the hsp60 in action with/without HEWL**

**Sample conditions**

Two samples were used to record the data. Hsp60 reference sample: 26 μM hsp60 (416 μM monomer concentration) U-[1H, 15N], Met-[13CH3]. Hsp60 with HEWL: 26 μM hsp60 (416 μM monomer concentration) U-[1H, 15N], Met-[13CH3] and 52 μM HEWL U-[1H, 15N], Val/Leu-[13CH3]proS, Ile-[13CH3]81 (two equivalents of HEWL per hsp60 particle). All the samples were prepared in D2O buffer containing 100 mM MES (pH 6.5 uncorrected), 100 mM NaCl, 150 mM KCl, and 25 mM MgCl2.

**Composition of the ATP regeneration system**

ATP regeneration system was composed of 3 U of pyruvate kinase from *Bacillus stearothermophilus*, 0.1 mM ribulose-5-phosphate, and 220 mM PEP (Sigma-Aldrich).

**Interleaved NMR experiments**

The experiment was started by the addition of 10 mM ATP to the reaction mix, and the NMR tube was placed in the spectrometer at a temperature of 65°C. After an equilibration time of 5 min, 2D SOFAST-methyl-TROSY NMR spectra were recorded in interleaved mode with 1H 1D spectra.

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/4/9/eaau4196/DC1

Fig. S1. Examples of mutant spectra used to assign individual methionine and valine correlations.

Fig. S2. Cross-validation of the valine and methionine methyl group assignments.

Fig. S3. Interaction of hsp60 with different nucleotides.

Fig. S4. Conformational changes of hsp60 induced by a nonhydrolyzable ATP analog (App-NHp) investigated by EM and NMR.

Fig. S5. NMR characterization of the HEWL thermal unfolding without/with hsp60.

Fig. S6. Translational diffusion properties characterized by diffusion-ordered NMR spectroscopy.

Fig. S7. Determination of HEWL binding site on hsp60 using PRE experiments.

Fig. S8. Refolding of HEWL by ATP-powered chaperonin.

Fig. S9. Rebinding of HEWL by ATP-powered chaperonin.

Fig. S10. Real-time NMR study of hsp60 functional cycle without client protein.

Fig. S11. Population of active hsp60 in ATP/ADP/apo state and closed/open state. References (42–45).

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