Protein Structure and Folding: 
Mechanism of an ATP-independent Protein Disaggregase: II. DISTINCT MOLECULAR INTERACTIONS DRIVE MULTIPLE STEPS DURING AGGREGATE DISASSEMBLY

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Mechanism of an ATP-independent Protein Disaggregase

II. DISTINCT MOLECULAR INTERACTIONS DRIVE MULTIPLE STEPS DURING AGGREGATE DISASSEMBLY

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Background: A novel chaperone, cpSRP43, disassembles a family of membrane protein aggregates.

Results: cpSRP43-mediated disaggregation requires two steps, recognition and remodeling, each with distinct molecular requirements.

Conclusion: cpSRP43 uses distinct substrate binding interactions to recognize and then remodel and disrupt the protein aggregate.

Significance: Mechanism of this novel ATP-independent disaggregate guides the understanding of analogous systems and design efforts to target protein aggregates of interest.

The ability of molecular chaperones to overcome the misfolding and aggregation of proteins is essential for the maintenance of proper protein homeostasis in all cells. Thus far, the best studied disaggregation systems are the Clp/Hsp100 family of “ATPases associated with various cellular activities” (AAA+) ATPases, which use mechanical forces powered by ATP hydrolysis to remodel protein aggregates. An alternative system to disassemble large protein aggregates is provided by the 38-kDa subunit of the chloroplast signal recognition particle (cpSRP43), which uses binding energy with its substrate proteins to drive disaggregation. The mechanism of this novel chaperone remains unclear. Here, molecular genetics and structure-activity analyses show that the action of cpSRP43 can be dissected into two steps with distinct molecular requirements: (i) initial recognition, during which cpSRP43 binds specifically to a recognition motif displayed on the surface of the aggregate; and (ii) aggregate remodeling, during which highly adaptable binding interactions of cpSRP43 with hydrophobic transmembrane domains of the substrate protein compete with the packing interactions within the aggregate. This establishes a useful framework to understand the molecular mechanism by which binding interactions from a molecular chaperone can be used to overcome protein aggregates in the absence of external energy input from ATP.

Protein homeostasis is vital to all cells and requires the proper production, folding, localization, assembly, and degradation of all cellular proteins (1). Crucial to the maintenance of protein homeostasis is an elaborate network of “molecular chaperones” (2–4), which prevents the misfolding and aggregation of proteins by protecting exposed hydrophobic residues in non-native states or unstructured regions and, in some cases, actively promotes protein folding (2). However under stress conditions, the folding capacity of the chaperone network could be exceeded or impaired, leading to protein aggregation. A special set of chaperone machineries, the “disaggregases,” plays a crucial role in rescuing these detrimental processes. The best studied disaggregases belong to the Hsp100 family: Hsp104 in yeast and ClpB in bacteria (5). Both are members of the “ATPases associated with various cellular activities” (AAA+) superfamily that assemble into hexameric ring structures (5). These disaggregases use repetitive ATPase cycles and, in collaboration with their co-chaperones, remodel large protein aggregates via translocation of the substrate protein through their central pores (6–8).

Despite the fascinating activity displayed by ClpB/Hsp104, their homologues have not been found beyond bacteria and yeast cells. Nevertheless, multiple lines of evidence indicate that maintenance of protein homeostasis in mammalian cells is critically dependent on cellular programs to overcome the deleterious effects of protein aggregation (9). Recently, it was demonstrated that ATP-independent actions of the mammalian Hsp110 and small heat shock proteins can engage and facilitate the remodeling of protein aggregates in collaboration with Hsp70 and Hsp40 homologues (10, 11). These observations suggest that cells, especially higher eukaryotic cells, have evolved alternative strategies and mechanisms to rescue protein aggregates.

Recently, we described a novel disaggregate system that operates independently of ATP: the 38-kDa subunit of the chloroplast signal recognition particle (cpSRP43) disassembles a family of membrane protein aggregates in the absence of external energy input from ATP. This ability to overcome the misfolding and aggregation of proteins is essential for the maintenance of proper protein homeostasis in all cells. Thus far, the best studied disaggregation systems are the Clp/Hsp100 family of “ATPases associated with various cellular activities” (AAA+). Both are members of the “ATPases associated with various cellular activities” (AAA+) superfamily that assemble into hexameric ring structures. These disaggregases use repetitive ATPase cycles, in collaboration with their co-chaperones, to remodel large protein aggregates via translocation of the substrate protein through their central pores.
An ATP-independent Disaggregase Mechanism

The abbreviations used are: cpSRP, chloroplast signal recognition particle; cpSRP43, a subunit of cpSRP; cpSRP from the chloroplast stroma to the thylakoid membrane (13). The most abundant member of this family, LHCP, comprises up to 50% of the protein content in the thylakoid membrane and is likely the most abundant membrane protein on earth (13, 14). LHCP proteins contain three hydrophobic transmembrane (TM) helices, making them highly prone to aggregation as they traverse aqueous compartments in the cell (14, 15). Recently, we and others showed that the cpSRP43 subunit of cpSRP acts as an effective molecular chaperone for the LHCP proteins (12, 16). Intriguingly, cpSRP43 also efficiently reverses the aggregation of LHCP proteins without the requirements for ATP hydrolysis or co-chaperones (12, 16).

cpSRP43 provides a valuable example of a novel category of ATP-independent chaperones/disaggregases that operates with energy derived solely from binding interactions with its substrate proteins. Understanding its mechanism of action will provide valuable insights into alternative principles and approaches that can be used to overcome protein aggregation problems. An increasing number of examples speak to the generality of this phenomenon. Mitochondria import stimulation factor (MSF) reverses the aggregation of mitochondrial precursor proteins and restores their import in an ATP-independent mode (17, 18). Small heat shock proteins play crucial roles in remodeling protein aggregates and facilitate their resolubilization by Hsp70/100 (10, 11). Cyclophilins reactivate the aggregates of adenosine kinase (19). ATP-independent disaggregase activities have also been reported in nematode and mammalian tissue (9, 20, 21). However, the mechanism by which protein aggregates can be disassembled based solely on the substrate binding energy of a chaperone remains elusive.

Many questions arise in addressing these mechanisms. First, what are the precise binding interactions between cpSRP43 and its substrate proteins? Previous work has demonstrated a specific interaction of cpSRP43 with a highly conserved 18-amino acid loop, L18, between TM2 and TM3 of LHCP proteins (22, 23). However, the ability of cpSRP43 to prevent LHCP proteins from aggregation implies that it must also protect the hydrophobic TMs of the substrate protein. Consistent with this notion, the binding affinity between cpSRP43 and full-length LHCP is at least an order of magnitude higher than that for the L18 peptide (12, 24). Thus additional interactions most likely exist between LHCP and cpSRP43, but the nature of these interactions remains to be determined. Second, how does cpSRP43 use these binding interactions to effect the reversal of protein aggregation? Previous kinetic analyses revealed that disaggregation is a cooperative process during which multiple cpSRP43 molecules recognize and actively remodel the LHCP aggregate (12). However, how the recognition and remodeling of the protein aggregate were accomplished by cpSRP43 has been elusive.

By combining molecular genetics with kinetic and thermodynamic analysis, here we present evidence that the interaction of cpSRP43 with its substrate proteins is composed of two components: sequence-specific recognition of the L18 motif and highly promiscuous interactions with hydrophobic TMs. These interactions enable distinct steps in the cpSRP43-mediated disaggregation of LHCP proteins: initial recognition and subsequent remodeling and disruption of the aggregate. The balance of these binding interactions with the energetics of packing interactions within the aggregate dictates the efficiency of the disaggregation reaction.

EXPERIMENTAL PROCEDURES

Materials—To construct the LHCP TM mutants (see Table 1), a pair of unique restriction sites was introduced into the expression plasmid encoding LHCP before and after the sequences encoding TM1, TM2, or TM3. The sequences coding for the TMs were replaced with PCR fragments encoding alternative TMs using the corresponding restriction sites. TM deletion mutants and Lhcb5 cysteine mutants were constructed using the QuickChange mutagenesis procedure (Stratagene). cpSRP43, LHCP, and its variants were purified as described (12).

Determination of Binding Affinity between cpSRP43 and Soluble Protein Substrates—Two independent methods were used to determine the apparent dissociation constant $K_d^{app}$ for cpSRP43-substrate binding. (i) The first method was prevention of LHCP aggregation by cpSRP43, monitored by light scattering at 360 nm after a 10-min incubation of the substrate protein with varying concentrations of cpSRP43 (12). The light scattering is linearly proportional to the concentration of LHCP except at very low concentrations (Ref. 12 and the accompanying manuscript (34)). The percentage of soluble substrates (% soluble) was analyzed as a function of cpSRP43 concentration. The data were fit to Equation 1,

$$%\text{soluble} = 100 \times \frac{[L] + [43] + K_d^{app} - \sqrt{[L] + [43] + K_d^{app}^2 - 4 \times [L][43]}}{2 \times [L]}$$

(Eq. 1)

in which [L] is the LHCP protein concentration and [43] is the cpSRP43 concentration. (ii) The second method was fluorescence anisotropy, as described previously (12). Briefly, all anisotropy measurements were conducted at room temperature using a Fluorolog 3-22 spectrofluorometer (HORIBA Jobin Yvon). Fluorescence-labeled LHCP or its variants (100 nm) were diluted into buffer containing different concentrations of cpSRP43. The samples were excited at 450 nm, and the fluorescence anisotropy was recorded at 524 nm. The data were fit to Equation 2,

$$A_{obsd} = A_0 + \Delta A \times \frac{[L] + [43] + K_d - \sqrt{[L] + [43] + K_d^2 - 4 \times [L][43]}}{2 \times [L]}$$

(Eq. 2)

in which $A_{obsd}$ is the observed anisotropy value, $A_0$ is the anisotropy value without cpSRP43, $\Delta A$ is the total change in anis-
sotropy, and \( K_d \) is the equilibrium dissociation constant. The \( K_d \) values measured by these two methods produced consistent results for the substrates tested (see Fig. 2A).

**Thermodynamic and Kinetic Analyses of cpSRP43-mediated Disaggregation**—Disaggregation reactions were performed as described previously (12), with the exception that aggregate formation was allowed to proceed for 1 min before the addition of cpSRP43. The observed light scattering intensity was normalized to that prior to the addition of cpSRP43. The disaggregation time courses were fit to an exponential function, Equation 3,

\[
A = A_0 + \Delta A e^{-k_{obs}t}
\]  
(Eq. 3)

in which \( A \) is the observed light scattering, \( A_0 \) is the amount of light scattering at \( t \to \infty \), \( \Delta A \) is the extent of light scattering change, and \( k_{obs} \) is the observed rate constant. The fractions disaggregated (\( K \)) were calculated as \( (\Delta A/(\Delta A + A_0)) \). The cpSRP43 concentration dependences of the value of \( K \) were fit to Equation 4,

\[
K = K_{max} \times \frac{[43]^{h}}{(K_m)^{h} + [43]^{h}}
\]  
(Eq. 4)

in which \( K_{max} \) is the extent of disaggregation at saturating cpSRP43 concentration, \( K_m \) is the concentration of cpSRP43 that enables 50% solubilization of the aggregates, and \( h \) is the Hill coefficient.

Kinetic analysis was performed and analyzed as described previously (12) to obtain the forward rate disaggregation rate constant, \( k_f \) from the observed rate constants (\( k_{obs} \)) and the extent of disaggregation (\( K \)). The cpSRP43 concentration dependence of the \( K_f \) values was fit to Equation 5,

\[
k = k_0 \times \frac{(K_m)^{h}}{(K_m)^{h} + [43]^{h}} + k_{max} \times \frac{[43]^{h}}{(K_m)^{h} + [43]^{h}}
\]  
(Eq. 5)

in which \( k_0 \) is the rate of spontaneous LHCP disaggregation in the absence of the chaperone, \( K_m \) is the concentration of cpSRP43 required to achieve half-maximal disaggregation rate, \( h \) is the Hill coefficient, and \( k_{max} \) is the disaggregation rate constant at saturating cpSRP43 concentration.

For some of the irreversible mutants (see Table 4, red), both the reaction equilibrium and the kinetics did not show detectable cooperative concentration dependences. Therefore, the data were fit to Michaelis-Menten equations (Equations 6 and 7).

\[
K = K_{max} \times \frac{[43]}{(K_m)^{h} + [43]}
\]  
(Eq. 6)

\[
k = k_{max} \times \frac{[43]}{(K_m)^{h} + [43]}
\]  
(Eq. 7)

Although direct evidence remains to be obtained, the following strongly suggests that the formation of an initial recognition complex between cpSRP43 and LHCP aggregates is fast when compared with subsequent remodeling and disassembly of the aggregate. First, in all the binding experiments, the cpSRP43-LHCP interaction is complete within the timescale of manual mixing (\( \pm 15 \) s), much faster than the overall disaggregation rates. Second, given an affinity of \( \sim 2 \) \( \mu \)m for the cpSRP43-L18 motif interaction (12, 24) and the typical range of macromolecular association rate constants \( (10^6-10^8 \) M \( \text{-1 s}^{-1}) \), the dissociation rate constant of the recognition complex would be in the range of \( 2-200 \) s \( ^{-1} \), much faster than the overall disaggregation reaction. Together, these observations suggest that the remodeling and disassembly of the aggregate is the rate-limiting step in the disaggregation reaction. Therefore, the cpSRP43 concentration required to achieve half of the maximal rate of disaggregation, \( K_{m/2} \), provides an empirical measure for the average binding affinity of cpSRP43 to the aggregate.

**Determination of the Energies of Aggregate Formation**—The energetics of packing interactions that drive aggregate formation were probed with a sedimentation assay. Briefly, preformed LHCP aggregates (10 \( \mu \)M) were resolubilized by various concentrations of guanidinium hydrochloride or urea for 30 min at 25 °C. The mixtures were centrifuged at 18,000 \( \times \) g for 30 min, and soluble (S) and pellet (P) fractions were visualized by SDS-PAGE. The intensities of the Coomassie Blue-stained bands for the pellet and soluble fractions were quantified using ImageJ (25). The data were fit to a two-state model (Equation 8) analogous to that for protein folding (26),

\[
F(\text{soluble}) = \frac{1}{1 + e^{-m(U_{50} + [\text{urea}]/T)}}
\]  
(Eq. 8)

in which fraction soluble is calculated as \( (S/(S+P)) \), \( R \) is the gas constant, \( T \) is temperature, \( U_{50} \) is the urea concentration to achieve 50% solubilization, and \( m \) is a constant of proportionality.

**Mathematical Analyses**—Linear regression analysis was performed using Mathematica to identify a weighted linear combination of \( U_{50} \) and \( \text{In}K_d \) values that best reproduces the \( \text{In}K_d \) values in Table 4. This was carried out by identifying the global minimum for the scoring function (Equation 9)

\[
f(\alpha, \beta, \gamma) = \sum \text{In}K_d - (\alpha U_{50} + \beta \text{In}K_d + \gamma)^2
\]  
(Eq. 9)

**LHCP Scanning Mutagenesis**—All molecular biology manipulations, including site-directed mutagenesis, transformation, and plating, were performed on a Tecan Freedom EVO liquid-handling robot. Constructs were sequence-verified and re-arrayed into master plates. These master plates served to inoculate 10-ml volumes of Instant TB autoinduction media in 24-well plates. After overnight expression at 37 °C, the hexahistidine-tagged proteins were purified under denaturing conditions. Cysteine scanning of the L18 motif was carried out using QuickChange mutagenesis (Stratagene). Single cysteine mutants of LHCP were analyzed using the light scattering assay as described under “Determination of Binding Affinity between cpSRP43 and Soluble Protein Substrates.”

**Plate-based Aggregation Prevention and Disaggeregase Activity Assay**—All LHCP variants were normalized to 45 \( \mu \)M and then diluted to 5 \( \mu \)M in 384-well plates by the liquid-handling robot. In the aggregation prevention assay, either no cpSRP43 or an equimolar amount (5 \( \mu \)M) or a 1:3 molar ratio (15 \( \mu \)M) of traditionally purified cpSRP43 was already present in each reac-

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5 A. Nisthal and S. L. Mayo, manuscript in preparation.
tion well. The reaction was followed by absorbance at 360 nm and allowed to proceed for at least 20 min. When measuring disaggregase activity, the cpSRP43 concentration was raised to a 1:6 molar ratio (30 μM) and added ~1 min after diluting the LHCP protein into aqueous buffer. Again, the reaction was followed for 20 min by measuring the absorbance at 360 nm. For both types of assays, the first time point was measured ~5 min after mixing, and the final time point was used for data analysis. The percentage of chaperone activity is defined as

$$\frac{\text{no chaperone}_{A_{360}} - \text{equimolar chaperone}_{A_{360}}}{\text{no chaperone}_{A_{360}}} \times 100$$

(Eq. 10)

where the equimolar chaperone$_{A_{360}}$ value can be substituted for the $A_{360}$ values of other chaperone conditions. Relative chaperone activity is then calculated by normalizing the percentage of chaperone activity to the WT LHCP value for each assay plate.

RESULTS

Bipartite Interactions of cpSRP43 with Soluble LHCP—To identify binding interactions of cpSRP43 with LHCP that are crucial to its chaperone and disaggregase activities, we performed exhaustive alanine-scanning mutagenesis in LHCP and assayed the chaperone activity using automated protocols on a Tecan Freedom EVO liquid-handling robot. Residues in the conserved L18 sequence between TM2 and TM3 of LHCP were further mutated to glycine and lysine. We tested the mutational effects on the interaction of cpSRP43 with LHCP by measuring the ability of cpSRP43 to (i) bind and thus prevent the aggregation of LHCP (Fig. 1, A, B, D, and E); and (ii) reverse existing LHCP aggregates (Fig. 1, C and F). Single mutations of every residue in an FDPLGL motif in L18 had large deleterious effects on both activities (Fig. 1, D–F), indicating that this motif plays a crucial role in the ability of cpSRP43 to bind and chaperone LHCP. In contrast, mutations in the remainder of the LHCP had modest to marginal effects (Fig. 1, A–C). An independent cysteine mutagenesis scan of the L18 sequence yielded the same results (Fig. 1, G and H). These results extend previous studies (12, 24), and together, they show that cpSRP43 makes highly sequence-specific interactions with the XDPLGX motif in the L18 sequence.

The absence of significant defects from point mutations of the remainder of LHCP (Fig. 1, A–C) suggests that the interactions of cpSRP43 with the TMs of LHCP are likely promiscuous. To provide independent evidence for this notion and to further probe the nature of the interaction of cpSRP43 with the TMs of the substrate protein, we constructed LHCP variants in which the individual TMs are deleted or swapped. In addition, the TMs in LHCP were replaced with those from unrelated membrane proteins, including the tail-anchored proteins SERP1, Sec61, and cytochrome $b_6$ (see Table 1 for nomenclature and composition of LHCP TM mutants used in this study). If the interaction of cpSRP43 with the TMs is sequence-specific, these mutations should significantly reduce the ability of cpSRP43 to bind and chaperone LHCP (12). On the other hand, if these interactions arise from generic hydrophobic interac-

![FIGURE 1. cpSRP43 makes highly sequence-specific interactions with the FDPLGL motif in the L18 sequence. A–F, alanine-scanning mutagenesis of the entire LHCP (A–C), and alanine- glycine-, and lysine-scanning mutagenesis within the L18 sequence of LHCP (D–F). The aggregation prevention activity of cpSRP43 was assayed at 1:1 (A and D) and 1:3 (B and E) molar ratios of LHCP to cpSRP43. aa, amino acids; mut, mutant. In C and F, the disaggregase activity was measured at 1:6 molar ratio of LHCP to cpSRP43. All assays were performed in 384-well plates using a Tecan Freedom EVO liquid-handling robot, as described under “Experimental Procedures.” G and H, single-cysteine substitutions at individual residues in L18 were tested for their ability to prevent the aggregation of LHCP (G) and to resolubilize existing LHCP aggregates (H). In G, a 1:1 ratio of cpSRP43 and LHCP was used. In H, a 5:1 ratio of cpSRP43 relative to LHCP (in aggregates) was used.](http://www.jbc.org/doi/abs/10.1074/jbc.R113.491978)
aural steps of this reaction. The disaggregation reaction mediated by cpSRP43 can be studied under single turnover conditions (26), minimizing complications from multiple turnover and facilitating interpretation of data. Both the kinetics and the equilibrium of this reaction exhibit saturable cooperative concentration dependences (Fig. 3, A and B) (12), strongly suggesting that the reaction involves at least two steps: (i) a higher-order step dependent on cpSRP43 concentration, presumably the assembly of a recognition complex between cpSRP43 and the aggregate (Fig. 3C, step 1) followed by (ii) a unimolecular step independent of chaperone concentration, presumably involving the remodeling and disruption of the aggregate to generate resolubilized cpSRP43–LHC complexes (Fig. 3C, step 2). Important parameters can be extracted from these data to empirically report on the energetics of these steps (Fig. 3 and Table 2). Assuming that the initial recognition step is fast when compared with the subsequent remodeling (see “Experimental Procedures”), the cpSRP43 concentration required to achieve half of the maximal disaggregation rate provides an empirical measure for the average binding affinity of cpSRP43 to the LHC aggregate (Fig. 3, A and C, and Table 2, $K_{\text{app}}$). The Hill co-efficient, $h$, denotes the minimum number of cpSRP43 molecules that cooperatively act together to disrupt the aggregate (Fig. 3C and Table 2). The maximal rate of disaggregation at saturating chaperone concentration, $k_{\text{max}}$, measures the energetic barrier for remodeling and disrupting the aggregate once the initial recognition complex is formed (Fig. 3, A and C, and Table 2). In equilibrium measurements, the fraction of LHC proteins resolubilized at saturating cpSRP43 concentrations, $K_{\text{app}}$, reports on the extent to which the interactions between LHC and cpSRP43 overcome the forces that stabilize the aggregate. Finally, at a subsaturating cpSRP43 concentration, the observed kinetics and equilibrium of LHC resolubilization ($k_{\text{app}}$ and $K_{\text{app}}$, respectively) measure the overall barrier to reach the transition state and the final cpSRP43–LHC complex, respectively.

To provide independent evidence that the disaggregation reaction can be experimentally dissected into distinct steps and to probe the molecular determinants that underlie each step, we characterized mutant cpSRP43 or LHC proteins that exhibit different defects in the disaggregation reaction. Below, we present evidence for two distinct classes of mutants that uncouple the initial recognition of the aggregate from its subsequent remodeling and solubilization and for the distinct molecular determinants that underlie these steps.

### Interaction with the L18 Motif Is Essential for Initial Recognition of the Aggregate

In the accompanying manuscript (Nguyen et al. (34)), the results of both electron paramagnetic resonance (EPR) and chemical modification experiments showed that in LHC aggregates, the hydrophobic TMs are buried in the interior, whereas the L18 motif is displayed on the exterior. These results suggest an attractive model in which cpSRP43 could recognize the L18 motif presented on the surface of the aggregate, initiating its action as a disaggregase. If this were the case, mutant LHC or cpSRP43s that specifically disrupt the L18-cpSRP43 interaction would impair the initial recognition of the aggregate, exhibiting defects in disaggregation at low chaperone concentrations. As binding is a higher-order process, the defects of these mutants could be overcome

### A Quantitative Framework to Analyze cpSRP43-mediated Disassembly of LHC Aggregates

To understand how the substrate binding interactions of cpSRP43 are used to drive the disassembly of LHC aggregates, it is crucial to establish a quantitative framework that describes the energetics of the individual steps of this reaction. We quantitatively measured the binding interactions of the TM mutants with cpSRP43 using two independent approaches: (i) the ability of cpSRP43 to bind and thus prevent the aggregation of substrate proteins, which provides a convenient measure for the apparent binding affinity ($K_{\text{app}}$) between this chaperone and the soluble LHC; and (ii) equilibrium titrations based on changes in the fluorescence anisotropy of fluorescein-labeled LHC upon its binding to cpSRP43 (12). The values of $K_{\text{app}}$ obtained from the two assays were comparable with one another (12) (Fig. 2A).

All the LHC TM mutants tested could be efficiently bound and protected from aggregation by cpSRP43 (Fig. 2), with efficiencies that differ no more than 5-fold from wild-type LHC. Some mutants, such as ΔTM3, SERP2, and Sec2, bound cpSRP43 with even higher affinity than wild-type LHC and are hence more readily protected by this chaperone (Fig. 2, A–E, green). Collectively, all the TM replacement mutants exhibit moderate to high affinities for cpSRP43, which are 10–100-fold higher than that of the isolated L18 peptide (24) (Fig. 2F). This strongly suggests that the hydrophobic TMs contribute additional binding interactions with cpSRP43. Further, these interactions are fairly generic and highly adaptable, in contrast to the strictly sequence-specific interactions of the L18 motif. Finally, these results show that cpSRP43 can protect a variety of aggregation-prone proteins, as long as the L18 motif is present to provide specific recognition.

### TABLE 1

<table>
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when a sufficiently high chaperone concentration is used to drive the initial binding. To test this hypothesis, we examined how mutations in the L18 motif of LHC or in the L18-binding sites of cpSRP43 affect the efficiency of disaggregation.

We identified two mutations in the L18 motif of Lhcb5 (a close homologue of LHCP), H160C and L170C, that weaken substrate binding with cpSRP43. Equilibrium binding assays showed that wild-type Lhcb5 binds tightly to cpSRP43, with a $K_{\text{d}}^{\text{app}}$ value of 10 nM, whereas mutants H160C and L170C exhibited weakened binding, with $K_{\text{d}}^{\text{app}}$ values of 30 nM and 1.1 μM, respectively (Fig. 4A and Table 3). Reciprocally, mutation of Arg-161 in cpSRP43 (R161A), which provides an important hydrogen bond partner with L18, significantly reduced the binding affinity of cpSRP43 to LHCP ($K_{\text{d}}^{\text{app}}$ = 1.2 μM, when compared with 138 nM with wild-type cpSRP43 (12)).

Consistent with defects in recognition of the LHCP aggregate, mutant cpSRP43-R161A exhibited severe defects in the reversal of LHCP aggregates at low chaperone concentrations (Fig. 4B, magenta versus black; Table 3, $K_{\text{d}}^{\text{app}}$ and $k_{\text{app}}$). However, when the concentration of the mutant chaperone was raised to compensate for the binding defect, cpSRP43-R161A could reverse LHCP aggregation. At saturating chaperone concentrations, close to 50% solubilization of the aggregate could be attained (Fig. 4B, magenta and Table 3). Analogously, the aggregates formed by the Lhcb5 mutants, H160C and L170C, exhibited defects in the disaggregation reaction that can be rescued by higher cpSRP43 concentrations (Fig. 4C, D, and Table 3). At saturating chaperone concentrations, the equilibrium and kinetics of disaggregation with the mutant aggregates are within 2-fold of those of wild-type Lhcb5 (Fig. 4C, D, and Table 3).
and Table 3). Finally, all three mutants exhibited much higher values of \( K_m \) in the disaggregation reaction when compared with the wild-type protein (Fig. 4 and Table 3), which correlated with their reductions in substrate binding affinity (12) (Fig. 4A).

Together, these results showed that L18 binding is a key requirement for the initial recognition of the aggregate by cpSRP43; further, this recognition event can be uncoupled from the subsequent concentration-independent step(s) in the disaggregation reaction.

A Class of LHCP TM Mutants Specifically Blocks the Disaggregation Process—Because the LHCP TM mutants contain intact L18 motifs, they provide a collection of substrates to probe for additional molecular requirements that underlie the disaggregase activity of cpSRP43. Surprisingly, although all the TM mutants can be efficiently bound and prevented from aggregation by cpSRP43 (Fig. 2), they exhibit striking differences in the thermodynamics and kinetics of the disaggregation reaction (Fig. 5 and Table 4).

The aggregates formed by some of the TM mutants, notably those of ATM3, SERP2, Sec2, and Cypb2, showed disaggregation kinetics and efficiencies that are comparable with or even higher than that of wild-type LHCP (Fig. 5 and Table 4, green). Notably, the aggregates formed by a group of mutants, especially 1-1-3, TM3, SERP2, and Cypb2, were virtually irreversible even when saturation in disaggregation rate constants had been reached at high cpSRP43 concentrations (Fig. 5 and Table 4, red). To a lesser extent, mutants 1-2-1 and TM1 also exhibited significant reductions in the disaggregation rates even when saturation was reached at high cpSRP43 concentrations (Fig. 5, E–H, blue, and Table 4). In the aggregate formed by all these mutants, the L18 motif is highly accessible and solvent-exposed (Fig. 6, A–C); this and the observation that saturation in disaggregation kinetics can be reached with these mutants indicate that their defects could not be accounted for by the inability of cpSRP43 to recognize the aggregate. In addition to substantial reductions in maximal disaggregation rates, the disaggregation reaction of these mutants lost cooperative dependence on cpSRP43, further supporting a specific defect in the ability of cpSRP43 to remodel and resolubilize the aggregate. Together, these results provide strong evidence for the presence of an additional remodeling step in the disaggregase mechanism of cpSRP43, whose molecular requirements are distinct from the initial recognition step.

The Irreversible LHCP TMMutants Form Ultrastable Aggregates—Unlike the L18-binding mutants, the irreversible LHCP TM mutants bind reasonably well to cpSRP43. What caused their defects in disaggregation? The results of chemical modification and EPR experiments showed that the TM segments are buried inside the aggregate and engage in strong interactions (see accompanying manuscript (34)). We hypothesized that the internal packing interactions within the aggregate...
gates are altered in these TM mutants, which could present higher barriers for cpSRP43 to remodel and disrupt the aggregate. To test this hypothesis, we probed the energetics of the packing interactions that stabilize the aggregate by quantitatively analyzing its solubility in chemical denaturants. Using the sedimentation assay, we showed that both guanidinium hydrochloride and urea could effectively solubilize the LHCP aggregate in a concentration-dependent manner (Fig. 7A). Quantification of the amount of solubilized LHCP as a function of urea concentration gave an aggregate solubilization curve analogous to protein unfolding curves (26) (Fig. 7, B–E). Based on a two-state model, quantitative analyses of these data yielded information about the energetics of transfer of LHCP from urea to water (ΔG°) and the urea concentration required to achieve 50% solubilization (Table 4, U50; see “Experimental Procedures”). These parameters provide quantitative empirical measures of the energetics of the internal packing interactions that drive aggregate formation.

The aggregates formed by the LHCP TM mutants exhibited a wide range of stabilities, with U50 values ranging from 2.5 to 5.7 M (Fig. 7, B–E, and Table 4). Notably, the four irreversible mutants whose aggregates could not be efficiently resolubilized exhibited the highest U50 values (4.7–5.7 M; Fig. 7 and Table 4, red). In contrast, some of the mutant aggregates that are more readily resolubilized by cpSRP43, such as Lhcb5-H160C (blue), and Lhcb5-L170C (red) by wild-type cpSRP43.

**TABLE 3**

<table>
<thead>
<tr>
<th>Construct</th>
<th>Kdapp</th>
<th>(Km)</th>
<th>Kmax</th>
<th>kcat/Km</th>
<th>kmax</th>
<th>kcat/Km</th>
</tr>
</thead>
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<tr>
<td>cpSRP43 R161A</td>
<td>1200</td>
<td>&gt;50</td>
<td>&gt;0.40</td>
<td>0.02</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Lhcb5</td>
<td>10</td>
<td>8.8</td>
<td>1.06</td>
<td>0.51</td>
<td>0.029</td>
<td>0.0042</td>
</tr>
<tr>
<td>Lhcb5 H160C</td>
<td>30</td>
<td>64</td>
<td>0.85</td>
<td>0.12</td>
<td>0.025</td>
<td>0.0021</td>
</tr>
<tr>
<td>Lhcb5 L170C</td>
<td>1100</td>
<td>&gt;90</td>
<td>&gt;0.35</td>
<td>0.03</td>
<td>&gt;0.014</td>
<td>N.D.</td>
</tr>
</tbody>
</table>

*Previously determined by fluorescence anisotropy (12).

* Denotes the values at the highest cpSRP43 concentration used.

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**FIGURE 4.** L18-binding mutants uncouple initial recognition of the aggregate from its subsequent solubilization. A, binding of cpSRP43 to wild-type Lhcb5 and L18 mutants H160C and H170C. The data were fit to Equation 1 (see “Experimental Procedures”) and gave Kdapp values of 10 nM for wild-type Lhcb5 (black), 30 nM for Lhcb5-H160C (blue), and 1.1 μM for Lhcb5-L170C (red). B, concentration dependences for the equilibrium of disaggregation of LHCP by wild-type cpSRP43 (black) or mutant cpSRP43(R161A) (magenta). C and D, chaperone concentration dependences for the equilibrium (C) and kinetics (D) of disaggregation of Lhcb5 (black), Lhcb5-H160C (blue), and Lhcb5-L170C (red) by wild-type cpSRP43.
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FIGURE 5. LHCP TM mutants exhibit a wide range of disaggregation efficiencies. A–H, representative concentration dependences of the equilibrium (A, C, E, and G) and kinetics (B, D, F, and H) for disassembly of the aggregates formed by the LHCP TM mutants. The data for wild-type LHCP (black) were shown as a reference of comparison in all four sets. The data in A, C, E, and G were fit to Equation 4 (black, blue, and green) or Equation 6 (red) to obtain $K_{\text{max}}$ values and extract $K_{\text{app}}$ values at 4 $\mu$M cpSRP43. The data in B, D, F, and H were fit to Equation 5 (black, blue, and green) or Equation 7 (red) to obtain $k_{\text{max}}$, $K_m$, and $h$ values and to extract $k_{\text{app}}$ values at 4 $\mu$M cpSRP43. All the thermodynamic and kinetic parameters were reported in Table 4.

uated how the maximal rate of disaggregation ($k_{\text{max}}$) correlates with the two energetic parameters that were varied in this set of mutants: (i) the binding affinity between cpSRP43 and the solubilized LHCP ($K_{\text{app}}$), which ultimately drives the disaggregation reaction; and (ii) the energetics of packing interactions that drive aggregate formation ($U_{\text{pack}}$), which must be overcome by
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TABLE 4
Summary of the thermodynamic and kinetic parameters of the LHCP TM mutants

<table>
<thead>
<tr>
<th>Substrate</th>
<th>$K_{\text{max}}$ (M)</th>
<th>$K_\text{app}^{4p}$ (M)</th>
<th>$\langle K_m \rangle$ (µM)</th>
<th>h</th>
<th>$k_{\text{max}}$ (s$^{-1}$)</th>
<th>$k_\text{app}^{4p}$ (s$^{-1}$)</th>
<th>$K_d^{\text{app}}$ (nM)</th>
<th>$U_{50}$ (M)</th>
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</thead>
<tbody>
<tr>
<td>WT</td>
<td>0.98±0.02</td>
<td>0.70±0.05</td>
<td>8.8±4.1</td>
<td>2.9±0.5</td>
<td>0.049±0.005</td>
<td>0.0068±0.001</td>
<td>111±3</td>
<td>3.8±0.2</td>
</tr>
<tr>
<td>1-3-3</td>
<td>1.01±0.04</td>
<td>0.72±0.08</td>
<td>11.7±1.1</td>
<td>1.8±0.1</td>
<td>0.068±0.006</td>
<td>0.0090±0.001</td>
<td>144±34</td>
<td>3.7±0.1</td>
</tr>
<tr>
<td>SERP3</td>
<td>1.19±0.10</td>
<td>0.33±0.03</td>
<td>12.2±1.9</td>
<td>2.6±0.4</td>
<td>0.025±0.005</td>
<td>0.0021±0.0003</td>
<td>207±51</td>
<td>4.0±0.1</td>
</tr>
<tr>
<td>1-2-1</td>
<td>0.95±0.01</td>
<td>0.08±0.01</td>
<td>15.6±0.3</td>
<td>4.6±0.4</td>
<td>0.0099±0.002</td>
<td>0.0011±0.0001</td>
<td>234±16</td>
<td>4.4±0.3</td>
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<td>ΔTM1</td>
<td>0.97±0.02</td>
<td>0.13±0.04</td>
<td>42.5±15</td>
<td>1.6±0.5</td>
<td>0.053±0.021</td>
<td>0.0017±0.0002</td>
<td>413±76</td>
<td>4.7±0.0</td>
</tr>
<tr>
<td>SERP2</td>
<td>1.05±0.03</td>
<td>0.96±0.04</td>
<td>8.5±1.3</td>
<td>2.3±0.2</td>
<td>0.25±0.04</td>
<td>0.042±0.009</td>
<td>9±5</td>
<td>3.5±0.1</td>
</tr>
<tr>
<td>ΔTM3</td>
<td>1.01±0.02</td>
<td>0.95±0.03</td>
<td>5.7±0.7</td>
<td>2.4±0.3</td>
<td>0.11±0.013</td>
<td>0.033±0.009</td>
<td>26±12</td>
<td>2.5±0.1</td>
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<tr>
<td>Sec2</td>
<td>1.09±0.02</td>
<td>0.81±0.01</td>
<td>6.6±0.5</td>
<td>2.9±0.3</td>
<td>0.088±0.008</td>
<td>0.017±0.005</td>
<td>36±17</td>
<td>3.3±0.1</td>
</tr>
<tr>
<td>Cyb2</td>
<td>1.12±0.11</td>
<td>0.66±0.04</td>
<td>6.8±0.4</td>
<td>3.6±0.5</td>
<td>0.056±0.004</td>
<td>0.0086±0.001</td>
<td>51±24</td>
<td>3.6±0.2</td>
</tr>
<tr>
<td>ΔTM2$^a$</td>
<td>0.62</td>
<td>0.04</td>
<td>N.D.</td>
<td>N/A</td>
<td>0.003</td>
<td>0.0004</td>
<td>216±88</td>
<td>5.7±0.1</td>
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<tr>
<td>1-2-2$^a$</td>
<td>N.D.</td>
<td>0.03</td>
<td>N.D.</td>
<td>N/A</td>
<td>0.006</td>
<td>0.0002</td>
<td>489±95</td>
<td>4.7±0.1</td>
</tr>
<tr>
<td>1-3-2$^a$</td>
<td>N.D.</td>
<td>0.03</td>
<td>N.D.</td>
<td>N/A</td>
<td>0.005</td>
<td>0.0003</td>
<td>456±206</td>
<td>4.8±0.1</td>
</tr>
<tr>
<td>1-1-3$^a$</td>
<td>0.56</td>
<td>0.02</td>
<td>N.D.</td>
<td>N/A</td>
<td>0.003</td>
<td>0.0003</td>
<td>490±57</td>
<td>5.7±0.1</td>
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</tbody>
</table>

N.D. = not determined. Values reported are averages from two or more independent experiments ± S.D. # denotes mutants that are fit with Equations 6 and 7 under “Experimental Procedures.” $K_{\text{max}}$ values were estimated for these mutants from the observed disaggregation rate constants at the highest cpSRP43 concentrations, where saturation was reached. Accurate $\langle K_m \rangle$ values could not be determined due to the extremely slow reaction of these mutants at low cpSRP43 concentrations and are hence not reported. N/A = not applicable.

cpSRP43 during disaggregation. A strong correlation was found between the maximal disaggregation rate constant and a weighted combination of the $U_{50}$ and $K_{\text{app}}^{4p}$ values (Fig. 8A, $R^2 = 0.96$), but not with either of the parameters alone. This correlation strongly suggests that once a recognition complex is formed, the competition between the packing interactions that stabilize the aggregate and additional binding interactions that cpSRP43 establishes with the TMsof LHCP dictates the resolubilization of the aggregate. Finally, two of the mutants, ΔTM1 and ΔTM3, exhibit significant deviations from this correlation (Fig. 8A, blue), suggesting preferences in the disaggregation pathway of cpSRP43 that are not accounted for by these two parameters (see “Discussion”).

Analysis of the relationship between the equilibrium and rate constants of disaggregation provides further insights into the nature of the rate-limiting remodeling complex (Fig. 3C, ‡). At a subsaturating cpSRP43 concentration (chosen at 4 µM), the free energy barrier ($\Delta \Delta G \sim ln K_{\text{app}}$) and the activation barrier ($\Delta \Delta G^2 \sim ln k_{\text{app}}$) of the disaggregation reaction showed an excellent linear correlation for the entire set of LHCP mutants (Fig. 8B), giving a slope of $\Phi = 0.73$. Analogous to the $\Phi$-value analysis of protein folding (26), this analysis could be used to infer the nature and structure of the transition state relative to the reaction substrate (the LHCP aggregate) and product (the solubilized cpSRP43-LHCP complex). The observation of a $\Phi$-value that approaches unity implies a fairly late transition state in which a substantial fraction of packing interactions within the LHCP aggregate is disrupted and those interactions with cpSRP43 are formed, albeit not to the same extent as those in the resolubilized cpSRP43-LHCP complex.

DISCUSSION

cpSRP43 provides an example of a novel class of chaperones that can effect the reversal of insoluble protein aggregates based solely on ATP-independent binding interactions with its substrate protein. The simplicity of this system makes it an accessible model system to delineate the molecu
ular mechanisms as well as the capability and limitations of ATP-independent disaggregases. Here, mutational analyses revealed distinct sets of binding interactions that this chaperone establishes with its substrate proteins. Further, molecular genetics combined with thermodynamic and kinetic analyses allowed us to dissect the molecular steps during the cpSRP43-mediated disaggregation reaction and revealed distinct molecular requirements and interactions that underlie each step. These results, combined with previous work (Ref. 12 and the accompanying manuscript (34)), led us to propose a two-step working model for the action of cpSRP43 as a protein disaggregate (Fig. 9A).

**Bipartite Interactions of cpSRP43 with Substrate Protein**

Previous work has established a specific interaction of cpSRP43 with L18, a conserved and relatively hydrophilic segment between TM2 and TM3 of the LHC family of proteins (22–24). The mutagenesis results here further demonstrated that this interaction is localized to the most conserved FDPLGL motif within L18, emphasizing the highly specific nature of this recognition. This is consistent with crystallographic observations in which DPLG was found to form a “turn” that wraps around Tyr-204 of cpSRP43, whereas the side chains in the remainder of the L18 peptide were not well resolved (24). Nevertheless, the ability of cpSRP43 to protect LHC proteins from aggregation implies that additional interactions must exist between this chaperone and the hydrophobic TMs on its substrate proteins. Although the precise motif(s) that mediate these additional interactions remain to be determined, the results here demonstrate that the interactions of cpSRP43 with the TMs are highly promiscuous, enabling it to bind and chaperone a variety of substrate variants in which the TMs were removed or replaced. Some of the substrate variants could be bound and chaperoned by cpSRP43 even more effectively than wild-type LHCP. Together, these results establish two important components of the binding interaction of cpSRP43 with substrate protein: highly specific recognition of the FDPLGL motif in the L18 segment and general hydrophobic interactions with the TMs of the substrate protein that are highly adaptable.

As discussed below, these two sets of binding interactions contribute to distinct stages in the action of cpSRP43 as a disaggregate.

**Different Binding Interactions Drive Distinct Stages of the Disaggregate Activity of cpSRP43**

**Recognition of the Aggregate**—To initiate the disaggregation reaction, cpSRP43 must first recognize and engage the LHC aggregates (Fig. 9A, step 1). Although additional interactions cannot be excluded, an attractive mechanism to drive this initial recognition is the binding of cpSRP43 to the L18 motif, which is displayed on the solvent-accessible exterior of the LHC aggregate (Fig. 9A, step 1). In support of this model, mutant proteins that disrupt the interaction of cpSRP43 with the L18 motif exhibit defects in disaggregation at low cpSRP43 concentrations and require much higher chaperone concentrations to reach saturation (Fig. 9B, ΔΔG). Consistent with a specific defect of these mutants in a binding step, their defects could be overcome by increasing the chaperone concentration such that the maximal rate and efficiency of the disaggregation reaction with these mutants are within 2-fold of that of the wild-type protein (Fig. 9B).

Intriguingly, the values of $K_m$ and $K_{D_H}$, which provide a proxy for the binding of cpSRP43 to the LHCP aggregate, are considerably weaker than the binding of cpSRP43 to the L18 peptide (24) and, with the exception of ΔTM1, vary from ~6 to 16 μM across different TM mutants (Table 4, white and green). On the one hand, this variation is much smaller than the up to 50-fold changes in the binding affinity between cpSRP43 and the soluble substrate protein (Table 4, $K_p$), supporting the notion that interaction with the L18 motif is a major driving force for the
initial recognition step and is less sensitive to variations in the TMs of LHCP. On the other hand, such variation, although modest, could not be explained by the simplest model in which the recognition step is equivalent to interaction of cpSRP43 with an isolated L18 peptide. It is possible that the L18 motif is presented in different configurations on the aggregates formed by the different TM mutants, which could cause the observed variations. Consistent with this possibility, EPR experiments showed that in the aggregate, spin probes in the DPLG motif exhibit much lower mobility than the remainder of the L18 sequence (see accompanying manuscript (34)), suggesting that this motif might contact the remainder of the aggregate and needs to undergo a rearrangement to interact with cpSRP43. Additional structural or sequence elements presented on the aggregate surface could also be recognized by cpSRP43. Consistent with this possibility, cpSRP43 cross-links to residues at the N terminus of TM3 (27); this segment is also exposed on the surface of the LHC aggregate (see accompanying manuscript (34)) and available for recognition by cpSRP43.

**Remodeling and Disruption of the Aggregate**—The class of irreversible TM mutants (Table 4, red), which exhibits severe defects in maximal disaggregation rate constants ($k_{\text{max}}$), provides strong evidence for a distinct remodeling step in the disaggregation reaction (Fig. 9A, step 2) that has different molecular and energetic requirements than the initial recognition step. As cpSRP43 effectively prevents the aggregation of these mutant LHCPs, the defects of these mutants in disaggregation are most likely kinetic, rather than thermodynamic in origin. Further, the observation that all the irreversible mutants form significantly more stable aggregates than wild-type LHCP (Fig. 9C, $k_{\text{max}}$) strongly suggests that the packing interactions within the aggregate present a major barrier for disaggregation (Fig. 9C, $k_{\text{max}}$; Fig. 8A) and that these packing interactions need to be substantially disrupted in the rate-limiting remodeling complex (Fig. 9A, species in brackets).

Additional insights into the remodeling step are provided by analyses of the entire series of TM mutants, which display a wide range of binding interactions with cpSRP43, packing interactions within the aggregate, and kinetics of disaggregation. In this series of mutants, the best predictor for disaggregation kinetics is provided by a combination of two energetic parameters: the packing interactions within the aggregate ($U_50$) compensated by the available binding interactions between cpSRP43 and soluble LHCP ($K_{\text{Dapp}}$; Fig. 8A). This correlation is
striking and implies that the transition state (or rate-limiting remodeling complex) for the disaggregation reaction involves substantial global disruption of the aggregate. Further, these disruptions must be compensated by the establishment of additional binding interactions between cpSRP43 and the TMs of the dislodged LHCP molecules. A notable example of the latter is SERP2, which forms an aggregate with a $U_{so}$ value comparable with wild-type LHCP, 1-3-3 or Cyb2, but displays the fastest disaggregation kinetics as cpSRP43 establishes the strongest binding interactions with this mutant. Together, these results strongly support the model that, once cpSRP43 recognizes and "latches" onto the LHCP aggregate, the competition between its binding interactions with the TMs of LHCP and the packing interactions that stabilize the aggregate dictates the efficiency of the disaggregation process.

Φ value analysis, which compares the extent to which mutations affect the barrier to reach the transition state versus that to the resolubilized cpSRP43-LHCP complex, provide additional insights into the nature of the rate-limiting remodeling complex. The Φ value of 0.73 observed here rules out early (Φ < 0) transition states and suggests a fairly late structure for the rate-limiting remodeling intermediate (26), in which a substantial portion of the packing interactions within the LHCP aggregate is disrupted and significant binding interactions with cpSRP43 have been established. A slightly alternative model, which takes into account potential heterogeneity in the action of cpSRP43, is that cpSRP43 disrupts the packing interactions at certain parts of the aggregate more extensively than at others, giving rise to a Φ value less than unity. As formation of the aggregate is a highly cooperative process (see accompanying manuscript (34)), it is conceivable that extensive disruption at multiple parts of the LHC aggregate could lead to the collapse of the
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network of packing interactions that drive aggregate formation (Fig. 9A), thus leading to its solubilization.

Perspective

The analyses here established two key requirements for how a chaperone can use binding interactions to reverse a protein aggregate. First, the chaperone must efficiently recognize and latch onto the target aggregate, through interactions with structural or sequence motifs displayed on the exterior of the aggregate. Second, the chaperone must effectively compete with and replace the internal packing interactions of the aggregate, by interacting with and protecting the segments of the substrate protein buried in the aggregate interior. Although the specifics of each system differ, these principles may be general to other ATP-independent chaperones that participate in or facilitate protein disaggregation processes.

Nevertheless, these do not represent all the features in the cpSRP43-mediated disaggregation reaction. Most notably, mutants ΔTM1 and ΔTM3 are clear outliers in the correlation analysis (Fig. 8A, blue squares). These deviations imply that factors in addition to the two parameters explored here ($U_{50}$ and $K_p$) contribute to the disaggregation reaction, and suggest preferred pathways in the action of cpSRP43. For example, ΔTM3 forms the loosest aggregate and can bind cpSRP43 tightly, but its rate of disaggregation is significantly slower than that expected from these considerations. Coupled with the observation that N-terminal residues of TM3 are also highly accessible on the aggregate (see accompanying manuscript (34)) and can contact cpSRP43 (27), this raises the intriguing possibility that cpSRP43 preferentially exerts its action on TM3 during the remodeling process to most effectively disrupt the aggregate. In contrast, ΔTM1 forms one of the tightest aggregates and has weaker binding interactions with cpSRP43, yet its maximal rate of disaggregation by cpSRP43 far exceeded what would be expected based on these parameters. This led us to speculate that TM1 is not a preferred site of action of cpSRP43 during the disaggregation process. Finally, despite these deviations, the aggregates formed both by mutants and by "hybrid" substrates containing TMs from unrelated membrane proteins are efficiently reversed by cpSRP43, demonstrating the remarkable adaptability of this chaperone.

It is noteworthy to compare the mechanism of aggregate remodeling by cpSRP43 to that of the ClpB/Hsp104 disaggregases. Many of the insights into the action of ClpB/Hsp104 are based on analogy with the ClpAP/ClpXP proteases (8), which use cycles of ATP binding and hydrolysis to drive repetitive movements of the substrate-binding loops, forcing the polypeptides through a constricted pore in the hexameric assembly and thus unfolding the substrate protein. By analogy, ClpB/Hsp104 could use ATPase cycles to drive translocation of a polypeptide, extracting it out of protein aggregates (28, 29). In this mechanism, each disaggreagase machine can locally sever an aggregate without disrupting the remainder of the aggregate. This is consistent with the observation that local, rather than global, structure and stability near the recognition sites dictate the efficiency of ClpA/ClpX (30, 31) and with the ability of Hsp104 to generate more amyloid fragments and thus promote amyloid propagation (32, 33). Although the precise molecular details remain to be elucidated, our results suggest that cpSRP43 acts globally, rather than locally, on the protein aggregate. The rate-limiting step in the reaction pathway of cpSRP43 involves the generation of a late intermediate in which the packing interactions within the entire aggregate are extensively disrupted and which requires the cooperative action of multiple cpSRP43 molecules. Conceivably, in the absence of external energy input, individual cpSRP43 molecules cannot compete with the packing interactions inside the aggregate and extract a soluble LHC molecule from it. Instead, multiple chaperones collectively disrupt and collapse the entire aggregate (Fig. 9A). The results here provide a valuable framework to probe the capability, effectiveness, and limitations of this alternative ATP-independent chaperone mechanism and to understand the design principles by which binding energy can be used to overcome the problems of protein aggregation.

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REFERENCES


