Insights into the client protein release mechanism of the ATP-independent chaperone Spy

Molecular chaperones play a central role in regulating protein homeostasis, and their active forms often contain intrinsically disordered regions (IDRs). However, how IDRs impact chaperone action remains poorly understood. Here, we discover that the disordered N terminus of the prototype chaperone Spy facilitates client release. With NMR spectroscopy and molecular dynamics simulations, we find that the N terminus can bind transiently to the client-binding cavity of Spy primarily through electrostatic interactions mediated by the N-terminal D26 residue. This intramolecular interaction results in a dynamic competition of the N terminus with the client for binding to Spy, which promotes client discharge. Our results reveal the mechanism by which Spy releases clients independent of energy input, thus enriching the current knowledge on how ATP-independent chaperones release their clients and highlighting the importance of synergy between IDRs and structural domains in regulating protein function.
imbalance in protein homeostasis (“proteostasis”) can lead to protein misfolding and aggregation, which is associated with a variety of human diseases such as peripheral amyloidosis, type II diabetes, cancer, cardiovascular diseases, and many neurodegenerative disorders. To maintain proteostasis, molecular chaperone proteins, which are fundamental to protein quality control systems in cells, exert multiple functions including de novo folding of nascent polypeptides, preventing aggregation of folding intermediates, disassembling protein aggregates, and directing the degradation of off-pathway misfolded proteins.

It is estimated that 36.7% of the amino acid sequences in chaperones can be classified as “intrinsically disordered regions” (IDRs, i.e., protein regions lacking fixed tertiary structures). IDRs exist in a heterogeneous ensemble of conformations and play various roles, including protein complex assembly, post-translational modification, signal transduction, substance transportation, and molecular recognition. The feature of molecular recognition presented by IDRs fits well with the promiscuous nature of molecular chaperones, which often need to recognize and bind to diverse kinds of proteins. Researchers have observed interactions between chaperone IDRs and client proteins and have noted the functional importance of IDRs in chaperone actions. For example, the disordered C terminus of DNAK provides a binding motif with weak affinity for the clients, maintaining the ability of DnaK to support bacterial growth under thermal stress. The disordered C terminus of GroEL actively disrupts kinetic trapping, locally misfolded regions in the client proteins to facilitate their refolding. Several “conditionally disordered” chaperones, such as Hsp26, Hsp33, and HdeA, rely on on-off disorder conformational changes to bind and protect various clients from stress conditions.

Despite increased research on the function of IDRs in chaperones, detailed kinetic and structural characterization of IDRs in chaperones is still lacking. A major challenge is that chaperone clients are prone to aggregation, which limits kinetic and structural analysis. In addition, the high flexibility of IDRs hinders the study of IDRs by traditional biochemical and biophysical methods. Nuclear magnetic resonance (NMR), on the other hand, has the unique advantage of providing detailed residue-by-residue information of protein disorder, making it highly suitable for investigating dynamic IDRs in chaperones. However, high-molecular-weight proteins are a challenge to resolve by NMR and thus, it has been difficult to structurally elucidate the roles of IDRs in canonical chaperones, such as Hsp70 and GroEL.

The ATP-independent chaperone Spy harbors long intrinsically disordered N and C termini, and their influence on Spy’s chaperone action remains enigmatic. The dimeric protein Spy has a molecular weight of 32 kDa and is a good model for structurally understanding the mechanism of IDRs in chaperone action by NMR spectroscopy techniques. Spy was initially identified in a genetic screen of chaperones to stabilize a protein called Im7 in the bacterial periplasm, and overexpression of Spy increased the steady-state level of Im7 up to 700-fold. Im7 is a model protein for protein-folding studies, and mutants of Im7 trapped in fully or partially unfolded states have been constructed and validated as highly soluble. Therefore, we used this model chaperone–client pair as well as a few other well-characterized Spy client proteins to address the fundamental questions of whether and how dynamic IDRs affect chaperone action from a kinetic and structural perspective.

Here, we show that the disordered N terminus of Spy facilitates client protein release. Data from mutagenesis studies, kinetic analyses, NMR spectroscopy, and molecular dynamics (MD) simulations together support that the N terminus can bind transiently to the concave, client-binding surface of Spy, primarily through electrostatic interactions between the N-terminal D26 residue and five positively charged residues on the concave surface. In addition, the binding sites of the N terminus on the concave surface of Spy partially overlap with the binding sites of client proteins. As a result, the N terminus can facilitate client protein release by competitively binding to Spy’s concave surface. Furthermore, the client release-promoting effect of the N terminus depends on its conformational restriction by adjacent regions in the full-length protein, since the peptide with the same sequence as the N terminus but free in solution cannot achieve this effect. Our work expands the existing working mechanism of Spy and reveals tantalizing new features of intrinsic disorder in chaperone action.

Results

The N terminus of Spy contributes to client release. Both the N and C termini of Spy (residues 1–28 and 125–138, respectively) are predicted to be highly disordered and are not visible in the crystal structure of Spy (Fig. 1a and Supplementary Fig. 1a). To investigate the effects of disorder on Spy’s chaperone activity, we deleted one or both termini of Spy and obtained the variants Spy29–138, Spy30–138, Spy31–124, and Spy29–124. We also started from Spy31–138, a previously characterized truncation mutant that retains the ability to stabilize Im7 in vivo and systematically shortened its termini, three residues at a time, to obtain a series of terminally truncated Spy variants (Fig. 1b).

We observed little change in Spy’s overall secondary structure after termini removal or truncation (Supplementary Fig. 1b). We then determined the ability of these variants to prevent the aggregation of dithiothreitol-reduced α-lactalbumin (α-LA). We found that deletion of the entire C terminus (Spy1–124) did not affect the anti-aggregation activity of Spy, but the absence of the entire N terminus (Spy29–138, Spy29–124) increased the activity of Spy by at least 1.7-fold (Fig. 1c). Additionally, removal of up to 23 residues from the N terminus resulted in moderately decreased activity, but further shortening by 26 residues resulted in an abrupt increase in Spy’s activity. In contrast, C-terminal deletion had a lower effect on Spy’s chaperone activity and there was no apparent correlation between the change in activity and the extent of C-terminal truncation (Fig. 1c). These results thus suggest that the N-terminal residues, rather than C-terminal residues, have a decisive influence on the chaperone activity of Spy.

We previously isolated activity-enhancing “Super Spy” variants through a genetic selection that coupled Spy’s ability to stabilize an unstable mutant of Im7 with antibiotic resistance of *E. coli*. These “Super Spy” variants all sacrificed their own stability for flexibility, thereby enhancing chaperone activity. To investigate whether a similar strategy is adopted by the terminally truncated Spy variants constructed in this study, we measured their unfolding Gibbs free energy (ΔG^‡) but found no obvious correlation between the stability and activity of these variants (Supplementary Fig. 1c), suggesting that changes in overall flexibility could not explain the activity differences of these variants.

To understand how terminal truncation affects Spy activity, we determined the association and dissociation kinetics of the Spy variants with the partially folded client, Im7 H40W L53A I54A (Im7AAW), by biolayer interferometry. We found significant differences in the dissociation constant (K_d) and the dissociation rate constant (k_off) between the terminal-truncated variants. Variants that showed higher anti-aggregation activities toward α-LA with the removal of 26 or 28 residues of the N terminus also displayed tighter binding to Im7AAW and slower release of Im7AAW (Fig. 1d, e and Supplementary Fig. 1d–i). In contrast, the association rate constants (k_on) of most variants were comparable to that of the Spy wild type (Fig. 1f). These results suggest that the
long and disordered N terminus of Spy may not be directly involved in the initial binding process of the client, but may contribute to the client release in a length- and composition-dependent manner.

To test whether the facilitation of client release by Spy’s N-terminus also applies to other clients and, more importantly, to clients with different folding states, we measured the binding and release kinetics of Spy wild type and Spy29-138 toward two unfolded clients, namely carboxymethylated α-LA and Im7 L18A L19A L37A (Im7Δ3), and one folded client, Im7 wild type. We found that for all clients, Spy29-138 exhibited slower client-release rates, slightly decreased association rates, and overall tighter affinity compared to those of the Spy wild type (Supplementary Fig. 2). These results thus highlight the generality of Spy’s N terminus in facilitating client release.

**D26 is the key residue in facilitating client release.** The removal of 20 or 23 residues from Spy’s N terminus gave rise to a ~2-fold increase in substrate release rates, while removal of 26 or 28 residues resulted in a sudden decrease in client release rates (Fig. 1e), suggesting that the three residues (H24, Q25, and D26) may play a critical role in facilitating client release. We previously isolated an activity-enhancing point mutation of Spy that substituted Q25 with an arginine residue20. Not coincidentally, the isolated an activity-enhancing point mutation of Spy that substituted Q25 with an arginine residue20. Not coincidentally, the isolated a charged residue at position 25 appears to counteract the effect of the three residues (H24, Q25, and D26) in facilitating client release. We previously isolated an activity-enhancing point mutation of Spy that substituted Q25 with an arginine residue20. Not coincidentally, the isolated an activity-enhancing point mutation of Spy that substituted Q25 with an arginine residue20. Not coincidentally, the isolated a charged residue at position 25 appears to counteract the effect of the three residues (H24, Q25, and D26) in facilitating client release. We previously isolated an activity-enhancing point mutation of Spy that substituted Q25 with an arginine residue20. Not coincidentally, the isolated a charged residue at position 25 appears to counteract the effect of the three residues (H24, Q25, and D26) in facilitating client release.
Fig. 2 The D26 residue located on Spy’s disordered N terminus facilitates client release. a Surface charge distribution of Spy and model client proteins.

The concave, client-binding surface of Spy (PDB: 3039) is predominantly positively charged (blue), while the surfaces of client proteins Im7 (1AYI), α-LA (1F6R), and ADH (4W6Z) are generally negatively charged (red). The electrostatic surface potential was calculated in PyMOL with the Adaptive Poisson-Boltzmann Solver tools 2.1 plugin. A color scale for the charge distribution ranging from -5 to S was applied. b Chaperone activities of different Spy variants in preventing the aggregation of DTT-reduced α-LA relative to the activity of Spy wild type. Kinetic parameters characterizing the interactions between Spy variants and Im7, including the dissociation constants $K_d$ (c), dissociation rate constants $k_{on}$ (d), and association rate constants $k_{off}$ (e). f The anti-aggregation activity of Spy29-124, in the presence and absence of various concentrations of Nt-pep. Representative curves from three independent measurements are shown. In the absence of Spy29-124, massive aggregation of α-LA resulted in the decline of the light scattering signal at 320 nm after 200 min (black curve). g Quantification for remaining anti-aggregation activities of Spy29-124 in the presence of various concentrations of Nt-pep, Nt-pep D26K, and Nt-pep D26R. For b–e, g individual data points (circles) and mean ± SD (n = 3 independent experiments) are shown. Statistical analysis was performed with Graphpad Prism 9.1.0 using one-way ANOVA with Tukey’s multiple comparisons test. α-LA agg. Prev means α-LA aggregation Prevention. Source data are provided as a Source Data file.

(Fig. 1e), suggesting that residues 1–23 are also able to counteract the effects of H24, Q25, and D26. Supporting this idea, we found five positively charged residues (K12, H16, H17, K18, and K20) located in the N terminal region prior to residues 24–26.

The interaction between Spy and its client proteins is mediated by both hydrophobic and electrostatic forces. Spy is a cradle-shaped molecule with an overall positively charged, concave surface for binding to clients, whereas many clients, such as Im7, α-LA, and ADH, are overall negatively charged (Fig. 2a). Since D26 is the only negatively charged residue among the 24th–26th residues, we hypothesize that D26 is the residue that contributes the most to client release. Possible speculation for the mechanism by which the N terminus facilitates client release is that the N terminus containing the negatively charged D26 competes with the client for binding to the concave surface of Spy, which may destabilize the Spy-client complex and facilitate the detachment of the client from Spy’s surface.

To test this hypothesis, we first replaced D26 with a negatively charged residue, glutamate (D26E), a neutral residue, alanine (D26A), or positively charged residues, lysine or arginine (D26K or D26R), and then examined the chaperone activity and substrate binding kinetics of the four variants. We found that D26E showed nearly the same activity and binding kinetics as Spy wild type, whereas variants D26A, D26K, and D26R exhibited enhanced activities compared with Spy wild type (Fig. 2b). More importantly, D26A, D26K, and D26R decreased $k_{on}$ and $K_d$ to levels comparable to the termini-free Spy29-124 variant without significant changes in the $k_{off}$ rates (Fig. 2c–e). These results suggest that the presence of a negative charge at the 26th position facilitates the release of Spy clients but does not interfere with the initial client-binding process. Electrostatic interactions depend on the distance between two objects. Thus, we reckoned that the release-promoting effect of D26 might be attenuated by inserting a linker between residue D26 and M27, which increases the distance between D26 and Spy’s concave surface. Indeed, insertion of the (GGGS)$_2$ linker reduced the $k_{off}$ rate and led to an increase in Spy’s chaperone activity (Supplementary Fig. 3a–c).

To obtain evidence that the D26-containing N terminus of Spy can compete with clients for binding to Spy’s concave surface, we monitored the aggregation kinetics of α-LA in the presence of Spy29-124 and the peptide corresponding to Spy residues 1–28 (hereinafter referred to as Nt-pep). Nt-pep itself had no effect on the aggregation kinetics of α-LA (Supplementary Fig. 3d). In contrast, the addition of Nt-pep in the presence of Spy29-124 reduced the ability of Spy to prevent α-LA aggregation in a dose-dependent manner (Fig. 2f), indicating that Nt-pep had an inhibitory effect on Spy’s activity. The same effect was not observed when increasing concentrations of the C-terminal peptide (corresponding to Spy residues 125–138, Ct-pep) was added to Spy (Supplementary Fig. 3e). Additionally, Nt-pep containing the D26K or D26R mutation showed a weaker inhibitory effect (Fig. 2g), underscoring the role of D26 in mediating the interaction between Nt-pep and Spy. Based on these results, we suggest that D26 may have direct electrostatic...
contact with the concave surface of Spy, leading to competition between the N terminus and negatively charged clients, thus affecting the release of clients from Spy.

The N terminus approaches Spy’s concave surface. To obtain structural insight into the above hypothesis, we performed MD simulations. We found that the N terminus of Spy readily swings into the positively charged, client-binding cavity of Spy, regardless of the starting conformation, and in the presence or absence of C terminus in the MD simulations. In contrast, substituting residue D26 with a positively charged arginine resulted in limited access of the N terminus to the concave surface (Supplementary Movies 1 and 2, Supplementary Fig. 4, and Supplementary Data 1).

To experimentally test our MD simulation results and to elucidate the detailed intramolecular contacts between the N terminus and Spy’s structured region, we performed intramolecular paramagnetic relaxation enhancement (PRE) experiments on 15N-labeled Spy1-124, where the paramagnetic spin label methanethiosulfonate (MTSL) was attached to a cysteine inserted adjacent to D26 or one residue away, i.e., immediately before Q25, D26, M27, or M28 (Fig. 3a and Supplementary Fig. 5a). The PRE phenomenon arises from magnetic dipolar interactions between unpaired electrons at the paramagnetic center and nearby nuclei (typically, backbone amide protons), which can provide long-range distance information up to 35 Å.

Following the backbone resonance assignments of Spy1-124 (Fig. 3b), we found that all the MTSL spin labels around residue D26 had strong effects on many residues of Spy1-124, with the spin label attached to the cysteine inserted between residues Q25 and D26 (hereinafter referred to as CI) having the strongest effect, resulting in the highest PRE intensity ratios (Fig. 3c and Supplementary Fig. 5b–d). In contrast, attaching the MTSL label to the cysteine substitution of M15 or G19, which is more than six residues away from D26, led to small PRE intensity ratios (Supplementary Fig. 5e, f). These results indicate that residues near D26 make extensive transient contacts with the structured region of Spy.

Furthermore, based on the PRE effect at CI, we noticed that all of the positively charged residues with side chains located on the concave surface of Spy (blue bars in Fig. 3c) have a PRE intensity ratio higher than 2, except for residue R43, whose PRE intensity ratio could not be calculated due to peak overlap. In contrast, only two of the eight positively charged residues with side chains pointing to the convex surface (black bars in Fig. 3c) have a comparable PRE signal. Therefore, these results support that the N terminus is predominantly approaching the concave surface of Spy. We then mutated residues Q25 and D26 on Spy1-124 CI to arginine and lysine, respectively (Fig. 3a), to assess the influence of the negative charge at position 26 on the accessibility of the N terminus to the concave surface of Spy. The PRE effects of MTSL attached at position CI of the mutant protein were dramatically reduced (Fig. 3c). This result well supports our speculation based on biochemical data (Fig. 2) that the negative charge of D26 is the key factor mediating intramolecular contacts between Spy’s N terminus and its concave surface.

To investigate whether the other two N-terminal negatively charged residues D2 and D10 also form electrostatic contacts with the concave surface of Spy, we introduced the MTSL spin label on a cysteine substitution at T5 of 15N-labeled Spy1-124 (Fig. 3a) and determined the intramolecular PRE effect. We found that the spin label attached to T5 affected many residues on Spy1-124, with positively charged residues on Spy’s concave surface being more affected than those on the convex side (Fig. 3c). Similarly, substituting D2 and D10 with positively charged lysine residues decreased the PRE intensity ratios (Fig. 3c). However, we also note that the PRE effect of the spin label attached to T5 was much weaker than the signal obtained when the spin label was placed at position CI. In addition, the D2K D10K mutation had a negligible influence on the PRE effect of the spin label attached to CI, but Q25R D26K dramatically decreased the PRE intensity ratios of the spin label attached to T5 (Fig. 3c). Together, these results suggest that although D2 and D10 can also approach the concave surface of Spy, their influence is much smaller than that of D26, the key residue that dictates the long-range electrostatic interactions between the N terminus and the structured region of Spy.

D26 interacts with key residues on Spy’s concave surface. We next set out to identify positively charged residues that might interact electrostatically with D26, focusing on all the arginine or lysine residues located on the central cavity of Spy (Fig. 4a). We reasoned that if we mutated the residue interacting with D26 to aspartate, we could reverse the electrostatic attraction between D26 and this position to repulsion, thus reducing the PRE effect of the spin label of CI on the cavity region. Therefore, we substituted these residues in 15N-labeled Spy1-124 CI and examined the PRE intensity ratios of the resulting five single point mutants (R43D, R50D, H65D, R89D, and H96D) and three double mutants (K54D R55D, R61D R62D, and K121D R122D) (Fig. 4b). In addition, the K75D was chosen as a negative control because the side chain of this residue points to the convex side of Spy, and its peak intensity is nearly not affected by the MTSL spin label attached to CI (Fig. 3c).

For eight of the nine variants, their TROSY spectra remained nearly unchanged, so most of their backbone resonances could be assigned according to the assignment information of Spy1-124 (Fig. 3b). However, the backbone resonance of Spy1-124 CI H65D could not be confidently assigned due to extensive chemical shift perturbations (CSPs). More importantly, these extensive CSPs suggest a non-trivial change in the overall structure of Spy. Therefore, we excluded H65D from further analysis. For accuracy, we only calculated the PRE intensity ratios of the remaining eight variants for residues showing non-overlapping resonance peaks.

We plotted the correlation of the PRE intensity ratios between the eight variants and Spy1-124 CI, and calculated the root-mean-square deviation (RMSD) values (Fig. 4b). Based on the correlation of PRE signals of Spy residues, we classified these variants into four categories. The data points for Spy1-124 CI R89D tends to be distributed below the diagonal compared with other variants, indicating overall increased distances between the MTSL label and its cavity region. The data points of Spy1-124 CI K54D R55D and Spy1-124 CI R61D R62D show the largest RMSD values (1.98 and 2.23, respectively) and data points deviate from the diagonal in both directions, indicating a considerable variation in the distance distribution between the MTSL label and their structured regions. The RMSD values for Spy1-124 CI R43D, Spy1-124 CI H96D, and Spy1-124 CI K121D R122D are moderately high (1.23–1.74), indicating some variation in the distance distribution. The last category includes Spy1-124 CI R50D and Spy1-124 CI K75D, which show the lowest RMSD values (0.89 and 0.78, respectively), suggesting similarity to Spy1-124 CI. Based on these results, we speculated that R89, K54, R55, R61, and R62 are the most promising residues to interact with D26, while R43, H96, K121, and R122 also possess the potential to be D26-interacting sites. Because we saw considerable variation in the distance distribution of the MTSL label, it is possible that D26 interacts with several positively charged residues on the concave surface. The disruption of the interaction between D26 and one such residue may be partially
compensated by enhancing the interaction between D26 and another such residue, which is in accordance with the inherent flexibility of the N terminus.

To further examine the roles of the above residues in mediating the interaction between the N terminus and the structured region of Spy, we substituted them with aspartate in Spy wild type and Spy29-124, respectively, and determined the binding kinetics of Spy, we substituted them with aspartate in Spy wild type and Spy29-124, respectively, and determined the binding kinetics of SpyK54D, R55D, SpyR61D, R62D, and SpyR89D (Fig. 4c), suggesting that these mutations also impede the ability of the N terminus to facilitate client release, which overweighs the reduction in client retention by these mutants.

We then mutated the D26 residue to lysine in the SpyK54D, R55D, SpyR61D, R62D, and SpyR89D variants to test whether re-establishing electrostatic interactions involving the 26th residue could restore the client release-promoting effect of the N terminus in these mutants. Indeed, we found that the $k_{off}$ rate of the double mutant SpyK54D, R55D, SpyR61D, R62D, and SpyR89D was similar to that of Spy wild type, whereas the release rates of the triple mutants SpyD26K, K54D, R55D, and SpyD26K, R61D, R62D were even higher than the rate of Spy wild type (Fig. 4c). For SpyK54D, SpyR55D, SpyR61D, SpyR89D, and the control protein SpyK75D, the additional D26K mutation decreased the $k_{off}$ rates, similar to the change in SpyD26K, SpyK54D, and the control protein SpyK75D, the additional D26K mutation decreased the $k_{off}$ rates, similar to the change in SpyD26K (Figs. 2d and 4c), indicating that these residues have no significant impact on the client release process mediated by the N terminus.
We found that all the four R89D trajectories showed clear tendency of pushing the MTSL spin label out of the bottom of the cavity (Supplementary Fig. 7 and Supplementary Data 1). Thus, based on these experimental and simulation results, we conclude that R89, K54, R55, R61, and R62 are the primary sites on the concave surface that mediate the interactions with D26 on the N terminus.

The N terminus acts in a competitive mode. We have shown that the Spy residues involved in electrostatic interactions with the N-terminal residue D26 are also involved in client binding of Im7AAW (Fig. 4 and Supplementary Fig. 6). This raises the possibility that the N terminus of Spy and the client share overlapping binding regions. To map and compare the two binding interfaces, we collected different NMR spectra and calculated the CSPs of Spy residues.

To identify the binding sites of client proteins on Spy, we assigned the 2D $^{15}$N, $^1$H-TROSY spectra of Spy$_{29-124}$ in its apo
form and in complex with Im7A3 (Supplementary Fig. 8a, b). We found that complex formation did not alter the secondary structure of Spy29-124 (Supplementary Fig. 8c). The binding site of Im7A3 on Spy29-124 was mapped by CSPs, and the Spy residues showing the strongest CSPs were considered to be part of the binding interface (Fig. 5a, d, g).

To determine the binding site of Spy’s N terminus, we used two approaches. In the first approach, we obtained the CSPs of Spy residues by comparing the 2D [15N, 1H]-TROSY spectra of Spy29-124 and Spy1-124 (Fig. 5b, e). The presence of the N terminus affects most of Spy’s residues in the structured region except for the residues located at the bottom of the concave surface and those pointing toward the convex surface (Fig. 5h). The CSPs are not due to changes in the secondary structure of Spy, since the differences in the 13C\textsubscript{\alpha} and 13C\textsubscript{\beta} secondary chemical shifts between Spy29-124 and Spy1-124 are small (Supplementary Fig. 8d).

In the second approach, we obtained the CSPs of Spy residues by comparing the 2D [15N, 1H]-TROSY spectra of Spy1-124 CI and Spy1-124 CI Q25R D26K (Fig. 5c, f, i). As mentioned earlier, the Q25R D26K mutation weakens the interactions between Spy and its N terminus (Fig. 3c). Based on CSPs, we found that these two different approaches revealed very similar residues as potential binding sites of the N terminus (Fig. 5d–i).

Overall, the interaction mode of Spy with Im7A3 and its N terminus is similar (Fig. 5d–i), well supporting our hypothesis that Spy uses at least partially overlapping interfaces to bind the client and its N terminus. Together with the biochemical and biophysical characterization described above, this result suggests that the N terminus can directly compete with the client for binding to Spy, which is the basis for promoting client release.

**Tethering of the N terminus enables efficient competition.** In our previous activity assay, the inhibition of Spy29-124 activity by the Nt-pep was detected only when the concentration of Nt-pep was at least 14-fold higher of Spy (Fig. 2f). In the native Spy protein, the N terminus is present in a 1:1 ratio with Spy29-124 and does not require an excess to function. Apparently, free Nt-pep is
much less competent in promoting client release compared to the native N terminus of Spy. We reasoned that tethering the N terminus to the structured region of Spy not only increases the effective local concentration of the N terminus but also imposes conformational constraints on its movement, making it easier to reach the central cavity of the concave surface and thus more effective to compete with the client. Nt-pep in solution lacks such conformational constraints, and thus it may adopt heterologous poses when interacting with the structured region of Spy, which is less effective for the competition. Indeed, we found that the interaction pattern of Nt-pep with Spy29-124 is different compared to the fused N terminus (Figs. 5e, f, h, i and 6a, b). The CSPs of Spy29-124 upon binding with 4 mM unlabeled Nt-pep were determined and plotted against the residue number of Spy29-124. The secondary structure of Spy29-124 is indicated on top. Color intensities correlate with CSP values, with darker colors indicating larger values. An overlay of the 2D [15N,1H]-HSQC spectra of 100 μM [15N]-labeled Spy29-124 in the absence (red) and presence of Nt-pep at molar ratios (Nt-pep: Spy29-124) of 10:1 (blue) and 40:1 (cyan). Representative residues with remarkable CSPs including M64, D66, N116, and A117 are labeled and their resonances are highlighted on the right. Source data are provided as a Source Data file.

Discussion
Knowing how chaperones bind and release client proteins is fundamental to a full understanding of chaperone action. Chaperones typically recognize and bind hydrophobic amino acid stretches exposed in nascent polypeptides, protein-folding intermediates, and denatured or misfolded proteins24,25. Electrostatic interactions also contribute substantially in client association26–28. During client protein release, ATP-dependent chaperones, such as GroEL/ES and DnaK/J, regulate the affinity for clients through cycles of nucleotide binding and hydrolysis, as well as co-chaperone association and dissociation, thereby actively releasing client proteins at specific stages of their chaperone working cycles29. It is generally believed that ATP-independent chaperones release their client proteins in a passive manner, depending on conformational changes induced by stressors or decreases in chaperone concentration due to protein turnover or cell growth13,21.

Our investigation of the effect of the disordered termini on Spy’s chaperone action revealed an intrinsic client-release mechanism for ATP-independent chaperones. Specifically, the disordered N terminus of Spy forms electrostatic interactions with the client-binding cavity of Spy, thereby promoting client release in a competitive manner (Fig. 7). Interestingly, these intramolecular interactions do not obviously affect the client association rate (k_on). A plausible explanation for this seemingly counterintuitive observation is that upon binding to the Spy
Client protein interconverts between different folding states by a previously proposed “folding-while-bound” mechanism. The N terminus of Spy can competitively bind to Spy’s concave surface via electrostatic interactions. This process interferes with client retention and thus facilitates client protein release. Released client proteins can rebind to Spy under stress conditions or fold spontaneously when stress conditions diminish. For clarity, only one copy of the N and C termini are shown on the structure of the Spy dimer.

In Supplementary Fig. 2, we observed that the client release-facilitating effect of the N terminus holds for both folded and unfolded Im7 variants. Considering the apparent conformational differences between these two clients, one would assume that they bind to different regions on Spy. However, by analyzing the binding patterns of Im7A3 (unfolded) and Im7WT (folded) on Spy, we found that these clients actually bind a very similar surface on Spy (Supplementary Fig. 10). This observation can be explained by the previously proposed “folding-while-bound” mechanism of Spy, whereby the moderate affinities between Spy and clients allow clients in various folding states to make conformational interconversions upon binding to Spy. Therefore, folded and unfolded clients may end up forming similar conformational ensembles on the Spy surface, leading to the observation of similar binding patterns on Spy. Thus, based on our findings in this study and previous results, we propose that the mechanism of action by which the N-terminus of Spy facilitates client release may be generally applicable to clients in different folding states (Fig. 7). This property of Spy may offer kinetic advantages allowing it to rapidly cycle between various folding species of client proteins when cells encounter stress conditions and enable spontaneously folding of the released clients when stress conditions diminish (Fig. 7).

Chaperones need to release client proteins at the proper time and the correct cellular location. Persistent binding of client proteins will result in collapsed proteostasis. For example, tenacious binding of Hsp70 by unfolded client proteins limits the amount of Hsp70 used to sequester the key transcriptional regulator, Heat shock factor 1 (Hsf1), leading to collapsed Hsf1 activity regulation, which is presumably associated with transcriptional activation of cancer-specific genes in malignant cells. In addition, a previous study reported that Spy variants severely deficient in client release negatively impacted bacterial fitness, leading to prolonged filamentous cell morphology.

To evaluate the physiological impact of N terminus removal, we performed a growth assay and found that overexpression of the Spy29-138 variant in a Δspy strain resulted in a decrease in strain fitness when cells were recovered from 1% butanol stress (Supplementary Fig. 11a–c). Similar growth retardation was observed in cells expressing SpyD26K, but not in cells expressing SpyD26E. It has been shown that the folding rate of clients bound to Spy is much slower compared to their spontaneous folding in solution. Consistent with this, we observed a similar inhibitory effect of Spy on the folding rate of another client, α-LA, and noted that the variant Spy29-138 retarded α-LA folding more severely than the Spy wild type (Supplementary Fig. 11d). Thus,
we speculate that the observed decrease in cell fitness could be related to the prolonged binding of Spy29-124 to clients, which hinders rapid spontaneous refolding of clients upon stresses decay. Approximately 97% (224 out of 231) of the Spy protein from 54 species have a disordered N terminus, and the three negatively charged residues (D2/D2/E2, D10/E10, and D26/E26) on the N terminus are highly conserved (Supplementary Fig. 12). Therefore, the presence of an N terminus carrying D/E residues and its facilitation of client release may represent an evolutionarily beneficial trait.

Recently, a similar mechanism of client release has been reported for the RNA chaperone Hfq, whose negatively charged, disordered C terminus forms intramolecular electrostatic contacts with its arginine-rich, client-binding 5m domain, thereby facilitating the release of annealed RNAs from Hfq[14,15,25]. In addition to intramolecular interactions, intermolecular interactions between chaperones and IDRs of their cognate co-chaperones also facilitate client release. For instance, the disordered N terminus of the nucleotide exchange factor FeS1 can mimic the protein client and bind to the substrate binding domain of Hsp70 to promote client release[26]. Intermolecular interactions are also thought to regulate the chaperone function of small heat shock proteins (sHsps): client binding is partially counteracted by interactions between the disordered C terminus of one sHsp monomer and the client-binding surface of another monomer[27]. Thus, these studies suggest that there may be a generic mechanism that is used by IDRs within chaperones to regulate client binding and release actively.

Interestingly, the peptide derived from the N terminus of Spy (Nt-pep) has much less ability to facilitate client release (Figs. 2f, g and 6 and Supplementary Fig. 9). While this may help explain why Spy homologs have chosen to maintain disordered N termini during evolution rather than develop co-chaperones with N terminus-like properties, it also raises more fundamental questions in chaperone biology that need to be addressed. For example, what are the minimum requirements for a prototypical chaperone that performs holding and folding activities in an ATP-independent manner? How can this knowledge guide the de novo design of artificial chaperones for a variety of cellular tasks? We believe that a deeper characterization of other simple chaperone systems will help to answer these questions.

In summary, our work reveals Spy’s active and competitive client release mechanism, which is an example of an emerging paradigm of autoregulation by client-mimicking peptides. This work expands our current knowledge of the functions of IDRs in molecular chaperones. It also highlights the necessity to explore the divergent roles of IDRs in chaperone action to gain a more comprehensive understanding of complex chaperone mechanisms.

**Methods**

**Mutagenesis and protein purification.** QuikChange site-directed mutagenesis (Agilent, catalog no. 200518) was employed to generate mutations in spy and ceiE7 (the gene encoding Im7). Spy wild type and mutant proteins were purified as below. Briefly, Spy or Spy variants containing the N-terminal His6-SUMO tag were overexpressed in BL21 (DE3) and initially purified by immobilized nickel-nitrilotriacetic acid (Ni-NTA) affinity chromatography. The His6-SUMO tag was then removed by overnight digestion with ubiquitin-like-specific protease 1. The tag-free Spy and Spy variants were further purified by cation-exchange chromatography and size-exclusion chromatography. Im7 and Im7 variants were purified by a similar protocol. Protein concentrations were determined by measuring absorbance at 280 nm, and all proteins were aliquoted into 1.5-ml Eppendorf tubes and stored at −80 °C until usage.

**Isotope labeling.** We obtained [15N]-labeled Spy variants by growing cells in medium containing 15NH4Cl. Additionally, we obtained [15N]-labeled Spy variants by growing cells in medium containing 15NH4Cl and 13C glucose and D2O. Isotopes were purchased from Cambridge Isotope Laboratories, Inc. (Tewksbury, Massachusetts).

**Chaperone activity assay.** Anti-aggregation activities of Spy and Spy variants were determined by detecting their abilities to inhibit the aggregation of DTT-reduced bovine α-LA, type III Sigma-Aldrich. Briefly, 50 μM α-LA was induced to aggregate by adding 20 mM DTT in a buffer (pH 6.9) containing 50 mM sodium phosphate, 100 mM sodium chloride, and 5 mM EDTA. The light scattering signals of α-LA aggregation were recorded at 320 nm in the absence or presence of Spy and Spy variants using a Synergy HTX Multi-Mode Microplate Reader for 10 h at room temperature. To quantify the activities of Spy variants relative to the wild type Spy, a standard curve was generated by plotting the slopes of α-LA aggregation curves (from 180 to 240 min) against Spy wild type: α-LA ratios (from 0.0375 to 2). The standard curve was fitted with Origin 8.6 software using a one-phase exponential decay equation and the relative activities of Spy variants to the wild type from 3 independent experiments were calculated according to the standard curve.

**Biolayer interferometry.** The association and dissociation kinetics of Spy variants toward two fully unfolded clients (carboxymethylated α-LA and Im7αL) were determined. A partially folded client (Im7ωα), and one folded client (Im7 wild type) were measured at an Octet RED96 system (Fortebio, Menlo Park, CA). Each client protein was biotinylated with EZ-link NHS-biotin (Thermo Fisher Scientific, Waltham, Massachusetts) at a 1:1 molar ratio of biotin to protein before loading onto the streptavidin sensors. Then each client-loaded sensor was immersed into a buffer (40 mM HEPES and 150 mM NaCl, pH 7.5) containing different concentrations of Spy and Spy variants (0.25, 0.5, 1.5, and 2 μM) for chaperone-client complex binding. Afterward, each sensor was transferred to the buffer without Spy to dissociate the chaperone-client complexes. Kinetic parameters were obtained by fitting the association and dissociation curves in SigmaPlot 12.0 software according to the following equations, respectively:

\[
y = ax + b e^{-k_y y + y_0}\times t
\]

where \(y\) is the observed wavelength shift, \(y_0\) is the constant to correct for baseline drift, \(a\) is the maximal wavelength shift, \(k_y\) is the observed association rate constant, \(k_d\) is the dissociation rate constant, respectively. The association rate constant (\(k_a\)) is then obtained according to the following equation:

\[
k_a = k_{obs} \times [Spy] + k_{aff}
\]

where \([Spy]\) is the Spy concentration. The dissociation constant \(K_d\) is calculated according to:

\[
K_d = k_{obs} / k_{aff}
\]

**NMR spectroscopy.** The NMR experiments were carried out at 25 °C on an Agilent DD2 600 MHz or 800 MHz spectrometers equipped with cryogenic probes. The U-[1H, 13C, 15N] Spy NMR samples were prepared in NMR buffer containing 40 mM HEPES-NaOH (pH 7.5), 150 mM NaCl, 100cocktail protease inhibitor, 1 mM PMSF, 0.03 % Na3C2O4, 3 mM DTT, and 90% H2O/10 % D2O. Chemical shifts were referenced to an external 4.4-dimethyl-4-silapentane-1-sulfonic acid. The backbone assignments were obtained using the TROSY versions of standard triple resonance experiments, including HNCO, HN(CA)CO, HNCA, HN(CO)CA, HN(COA)CB and HNCA(CO)CB in [15N,1H]-LA and Im7A3, one partially folded client (Im7AAW), and one folded client (Im7 wild type) were titrated with unlabeled Im7 or Im7αL at a 1:1 molar ratio in NMR buffer A. The 2D H-15N TROSY spectra were recorded at 25 °C. The chemical shift was calculated using the following equation:

\[
\Delta \chi = \chi_q - \chi_0 = \left( \chi_q - \chi_{max} \right) / \chi_{max}
\]

**Nt-pep NMR titration experiments.** Titration experiments were performed on the Agilent DD2 800 MHz equipped with a cryogenic probe by mixing U-[1H, 13C, 15N] Spy124, U-[1H, 13C, 15N] Spy29-124 and U-[1H, 13C, 15N] Spy29-124 were titrated with unlabeled Im7 or Im7αL at a 1:1 molar ratio in NMR buffer A. The 2D 1H-15N TROSY spectra were recorded at 25 °C. The chemical shift was calculated using the following equation:

\[
\Delta \chi = \Delta \chi_0 = \left( \chi_q - \chi_0 \right) / \chi_{max}
\]

\[
\Delta \chi_0 = \chi_0 - \chi_{max}
\]

\[
\chi_{max} = \chi_{max} - \chi_{max}
\]

\[
[\text{Spy}] = \left( \chi_q - \chi_0 \right) / \chi_{max}
\]

\[
[\text{Spy}] = \left( \chi_q - \chi_0 \right) / \chi_{max}
\]

\[
[\text{Spy}] = \left( \chi_q - \chi_0 \right) / \chi_{max}
\]

\[
[\text{Spy}] = \left( \chi_q - \chi_0 \right) / \chi_{max}
\]
Spin labeling and PRE experiments. Spin labeling of the cysteine mutants of different Spy variants with MTS was performed according to a published protocol45, which is described as follows. First, upon reducing the cysteines within the Spy mutants, DTT was removed from the samples by buffer exchange into 40 mM HEPES-NaOH (pH 7.5), 150 mM NaCl. The protein samples were then diluted to the concentration of 20 μM and reacted with a 26-fold excess of MTSL. Labeling was performed for 12 h on a rocking shaker at room temperature. The unreacted MTS was removed by ultrafiltration. Labeling efficiency was measured using matrix assisted laser desorption ionization–time of flight mass spectrometry (MALDI-TOF MS) and was above 80% for all samples. The paramagnetic NMR sample was solvated in a truncated octahedral SPC/E water box and neutralized using the ff14SB force field49. The HSLC/TOY hydrogen HSQC experiments were performed at 25 °C on a Bruker Avance 600 MHz spectrometer for both paramagnetic and diamagnetic samples. The PRE effect is quantified by the ratio of each individual peak between the diamagnetic and paramagnetic states of the same protein. Spectra were processed and analyzed with Sparky and NMRRpipe41.

Modeling of disordered N terminus. Since the N terminus is missing in the crystal structure of Spy (PDB: 3O39), we have to rebuild it and attach it to the crystal structure. Four N termini with different conformations were constructed. The first one was generated simply using CABS-dock46. The other three were constructed using Amber1880, according to an in-house developed protocol, as detailed below. First, 5000 random coil conformations were generated using OOPS and SCWRL449 and then linked to both monomers of the crystal structure of Spy. The ff14SB force field49 was used for the structural refinement. Each structural model was solvated in a truncated octahedral SPCE/water box and neutralized using the chloride ions. The solvated system was subjected to energy minimization for 1000 steps. Harmonic constraints applied on the heavy atoms of Spy’s cavity regions (force constant 500 kcal mol−1 Å−2). The optimized system was first heated from 0 to 1000 K and then cooled from 1000 to 0 K under NVT ensemble with Berendsen thermostat. Both heating and cooling were divided into five stages with a temperature increment of 200 K. During this simulated annealing procedure, simulations were performed for 40 ps (heating) or 2 ps (cooling) with harmonic restraints (force constant 500 kcal mol−1 Å−2) applied on all heavy atoms except for the residues of N terminus. Finally, the structure was minimized for 1000 steps under implicit solvent (igb = 5) in the absence of restraints and the energy was reported. In the resulting structure pool of Spy with N termini, we filtered out those with D26 located inside the cavity of Spy to make sure D26 will not by accidentally “stick to” the positively charged residues in the cavity and thus cannot escape from the cavity within the reasonable simulation time. In doing so, an upper plane of Spy’s cavity was determined by four points (the geometry center of the Ca atoms of E110 and K112, and the geometry center of the Ca atoms of E120 and K121, with both centers calculated from each monomer). The top 3 structures with lowest energy were selected for final MD simulations.

MD simulations. A total of four conformations were used as the template to build the Spy variants. These variants were modeled so that each Spy monomer has a cysteine insertion before the D26 residue and one or two mutations to the predicted D26-interacting residues, including K41D R55D, R61D R62D, and R89D, as well as the K75D that serves as a control. To model the PRE spin label, an MTSL molecule was attached to the N-terminal cysteine residue. The resulting proteins were simulated with the ff14SB force field. The force field parameters of the MTSL tag were generated using the CHAMBER module of Amber1880. The proteins were solvated in a truncated octahedral box of SPCE/water molecules such that the boundary of water box was at least 10.5 A away from any atoms of the protein. Sodium and chloride ions were then added to achieve electroneutrality of the system and an ionic strength of 150 mM NaCl. The hydrogen mass reparation scheme was applied to enable the use of a larger integration step (Δt) in order to achieve higher production rate of MD, by using the ParmEd module of AMBER18 to generate modified topology files for subsequent MD simulations. The solvated system was minimized for 1000 steps with harmonic restraints (force constant 500 kcal mol−1 Å−2) on Spy’s structured region, followed by 1000 steps without restraints. The system was then heated for 14 ps from 0 to 310 K, and equilibrated for 1 ns at 310 K. This temperature (rather than the experimental temperature 298 K) was chosen to increase the conformational sampling. The 1 μs production stage of MD simulation was conducted at 310 K using the NPT ensemble. During the simulations, all bonds involving hydrogen atoms were constrained using the SHAKE algorithm implemented in Amber18. The nonbonded cutoff was set to 10 Å. Both the cpptrj module of Amber18 and our in-house Python scripts (using Python 2.7) were used for analyzing the distance distribution. VMD45 was used together with PyMOL (The PyMOL Molecular Graphics System, Version 2.1.1 Schrödinger, LLC.) for visualizing the MD results.

Data availability. The backbone chemical shifts generated in this study have been deposited in the BioMagResBank under the accession code BMRB 51107 (H, 15N and 13C backbone chemical shift assignment of Spy29−124) and BMRB 51108 (H, 15N and 13C backbone chemical shift assignment of Spy29−124 in complex with Im7). The previously published crystal structure of Spy used was obtained from the Protein Data Bank: 3O39. Source data are provided with this paper.

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Author contributions
W.H. and X.L. contributed equally to this work. W.H., B.W., Y.X., and S.Q. conceptualized project. W.H., Y.J., M.L., and I.Z. performed biochemical experiments, analyzed the data, and prepared NMR samples. X.L. and H.X. collected and analyzed the NMR data. X.L. performed MD simulations and analyzed the data. W.H., X.L., J.X., B.W., Y.X., and S.Q. wrote the paper with input from all authors. All authors read and approved the manuscript.

Competing interests
The authors declare no competing interests.

Additional information
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