Intrinsically disordered interaction network in an RNA chaperone revealed by native mass spectrometry

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RNA-binding proteins contain intrinsically disordered regions whose functions in RNA recognition are poorly understood. The RNA chaperone Hfq is a homohexamer that contains six flexible C-terminal domains (CTDs). The effect of the CTDs on Hfq’s integrity and RNA binding has been challenging to study because of their sequence identity and inherent disorder. We used native mass spectrometry coupled with surface-induced dissociation and molecular dynamics simulations to disentangle the arrangement of the CTDs and their impact on the stability of Escherichia coli Hfq with and without RNA. The results show that the CTDs stabilize the Hfq hexamer through multiple interactions with the core and between CTDs. RNA binding perturbs this network of CTD interactions, destabilizing the Hfq ring. This destabilization is partially compensated by binding of RNAs that contact multiple surfaces of Hfq. By contrast, binding of short RNAs that only contact one or two subunits results in net destabilization of the complex. Together, the results show that a network of intrinsically disordered interactions integrates RNA contacts with the six subunits of Hfq. We propose that this CTD network raises the selectivity of RNA binding.

RNA chaperone | small RNA | intrinsically disordered protein | ion mobility mass spectrometry | surface-induced ion dissociation

Many RNA-binding proteins (RBPs) contain intrinsically disordered regions (IDRs) (1) with overlapping functions that have been difficult to disentangle. For example, IDRs may augment specific RNA recognition, connect different RNA-binding modules, and enable the assembly of liquid condensates, while also serving as targets for posttranslational modification (2–4). The heterogeneous and dynamic structures of IDRs make their interactions especially challenging to quantify, and their functions in most RBPs remain poorly understood.

Hfq is a bacterial Sm protein that binds small noncoding RNA (sRNA) and chaperones sRNA regulation of complementary messenger RNAs (mRNAs) (5) (Fig. 1). Deletion of Hfq results in pleiotropic effects, including maladaptive responses to stress. The cell, if Hfq must distinguish its RNA partners from many similar nucleic acids. Mass spectrometry dissociation patterns, together with molecular dynamics simulations, showed that flexible extensions of each Hfq subunit form a dense network that interconnects the entire hexamer. This network is disrupted by RNA binding, but the lost interactions are compensated by RNA-Hfq subunits. By measuring interactions that are too irregular to be counted by other methods, mass spectrometry shows how flexible protein extensions help chaperones like Hfq recognize their RNA partners in the messy interior of the cell.

Significance

Hfq is a protein hexamer necessary for gene regulation by noncoding RNA in bacteria during infection or under stress. In the cell, Hfq must distinguish its RNA partners from many similar nucleic acids. Mass spectrometry dissociation patterns, together with molecular dynamics simulations, showed that flexible extensions of each Hfq subunit form a dense network that interconnects the entire hexamer. This network is disrupted by RNA binding, but the lost interactions are compensated by RNA-Hfq subunits. By measuring interactions that are too irregular to be counted by other methods, mass spectrometry shows how flexible protein extensions help chaperones like Hfq recognize their RNA partners in the messy interior of the cell.

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Disordered CTDs Stabilize Hfq. To better understand the interactions of the flexible CTDs, we first used nMS-SID on wild-type (WT) E. coli Hfq (102 aa per subunit) and a truncated Hfq lacking the CTDs (HfqΔCTD; 65 aa per subunit). Precursor ions (hexamer) and their product ions were resolved by m/z and by drift time (SI Appendix, Fig. S1 B and C). We analyzed the dissociation products (pentamer, tetramer, trimer, dimer, and monomer) obtained at increasing collision energies (CE) with the surface (Fig. 2 A). The resulting energy-resolved mass spectra (ERMS) showed that the two proteins have different stabilities and fragmentation patterns (Fig. 2 B and C). HfqΔCTD reached 20% fragmentation (80% hexamer remaining) at ~220 eV, compared to ~390 eV for WT Hfq (Fig. 2 B and C, black lines). This large difference demonstrated that the WT protein is much more stable than the truncated version, in agreement with previous reports (14).

The fraction of HfqΔCTD hexamer sharply decreased with modest increases in collision energy, with 90% fragmentation of the hexamer at CE ~350 eV. In contrast, dissociation of WT Hfq increased gradually over a wide range of CE, and ~20% of the hexamer remained intact even at CE = 1,000 eV; each CTD acts locally and independently, or if the six CTDs act together to accommodate or displace an incoming RNA (Fig. 1). Moreover, the energetic contributions of individual CTDs to RNA binding have been almost impossible to quantify.

We addressed these challenges by using native mass spectrometry (nMS) coupled with surface-induced dissociation (SID) (Fig. 2 A and SI Appendix, Fig. S1). In nMS, the protein complex is exchanged into a volatile electrolyte, allowing transfer of the intact native complex to the gas phase (24). After ionization, collision of the precursor ion with a surface (nMS-SID) dissociates the complex into product ions that provide information about the stabilities of the noncovalent interfaces within the complex and their molecular organization (25). This method has been used to characterize the stability, structure, and assembly pathways of many protein complexes, including RBPs and membrane proteins (26–28). Although nMS does not reveal atomic detail, it is uniquely capable of resolving mixtures of complexes by mass and shape. Yet, despite its promise for discovery, nMS-SID studies of large biomolecular complexes typically require customized instrumentation (SI Appendix, Fig. S1 A).

Here, by using nMS-SID and all-atom MD simulations, we show that the six disordered CTDs of apo Hfq form extensive interactions that connect and stabilize the entire hexamer. When RNA binds any subunit of Hfq, these stabilizing interactions are disrupted throughout the hexamer. Taken together, our results show how disordered regions can integrate RNA–protein interactions across a multisubunit chaperone.
Hfq fragmentation was less steep than for HfqΔCTD even when considering the broader collision energy range involved (SI Appendix, Fig. S2). This gradual response to higher CE suggested that the CTDs prevent dissociation of the complex across a range of energies. One explanation is that the CTDs form intersubunit interactions that reorganize upon activation. Additionally, the CTDs may be organized differently in each Hfq hexamer, causing dissociation over a continuum of CE.

Disordered CTDs Impact the Connectivity of Hfq. The dissociation patterns for HfqΔCTD and Hfq generated fragments in different ratios, indicating a different degree of connectivity between subunits in the two proteins (Fig. 2 B–E). We compared the dissociation products of the two proteins at energies that deplete the hexamer equally. HfqΔCTD dissociated into twice as many trimers (−30%) as other fragments (−15 to 20%) regardless of the total amount of hexamer dissociated (Fig. 2D).

For WT Hfq, the distribution of dissociation products was markedly different from HfqΔCTD (Fig. 2E), indicating that the CTDs contribute to the subunit interfaces, as also reported earlier (14, 29). For example, the percentage of trimers decreased while the percentage of monomers increased as more hexamer was fragmented. This suggests that some hexamers dissociate at energies high enough to produce secondary fragmentation.

Inspection of ion mobility arrival times revealed that as the collision energy increases the initial WT hexamer converts into two complexes that migrate more slowly in the drift chamber, suggesting partial extension or restructuring of the protein hexamer (Fig. 2 F and G). We observed one additional conformation of HfqΔCTD upon activation (Fig. 2F), suggesting that one of the extended Hfq complexes comes from restructuring of the core beta sheet. The second extended form was only observed for WT Hfq and likely arises from extension of the CTDs (Fig. 2G). Altogether, the nMS results support a model in which the CTDs are bound to the core and each other, stabilizing the entire WT hexamer. At increasing CE, the CTDs disentangle, exposing the core and eliminating the stabilizing intersubunit connections. As a result, the fragile hexamer ruptures into smaller complexes.

MD Simulations Reveal a Network of CTD Interactions on Hfq. To gain more insight into the organization of the disordered CTDs on Hfq, we performed multiple all-atom MD simulations on the WT protein. The simulations were started from 10 previous low-energy Rosetta models of the CTDs (models 1 to 10) (11) and an extended conformation started with four different initial velocities (models e1 to e4). Trajectories ranged from 110 ns to 1500 ns long, with total simulation time for all models = 8.9 μs (SI Appendix, Table S2). For all simulations, the radius of gyration (Rg) dropped after ~100 ns and remained approximately constant thereafter (SI Appendix, Fig. S3). The obtained Rg agreed with the experimentally determined nMS collisional cross-section (SI Appendix, Table S1).

The simulated CTDs adopted a variety of conformations (Fig. 3A and SI Appendix, Fig. S4), in agreement with NMR experiments showing they are disordered (12, 30). These interactions continued to evolve after the first 100 ns, with some CTDs entering periods of mobility and others remaining less mobile (SI Appendix, Fig. S5). CTD movement was observed in all simulations, although no two CTDs adopted the same conformation (SI Appendix, Fig. S6). Thus, the diverse CTD structures confer heterogeneity to individual Hfq hexamers, as implied by the ERMS results (Fig. 2C).
To evaluate the interchain interactions, we counted the number of subunit cores or the number of CTDs contacted by each CTD for all models (SI Appendix, Figs. S7 and S8). Most CTDs interacted with the core of the same subunit and adjacent subunits, but some CTDs engaged as many as five other subunits (SI Appendix, Figs. S7 and S8). These long-range intersubunit interactions were common (SI Appendix, Fig. S8). As illustrated for the three longest (1.5 μs) trajectories in Fig. 3 B and C, an average of two CTDs per hexamer interacted with three or four subunit cores over a 100-ns interval (Fig. 3B and SI Appendix, Figs. S7 and S8). This distribution was similar at the beginning and end of the simulations but shifted to larger values when averaged over the entire 1,400-ns simulation (Fig. 3C, purple and SI Appendix, Figs. S7 and S8). Although the MD simulations cannot sample the full range of structures accessible to the Hfq CTDs, their disorder is apparent even on the 1-μs timescale and among 14 different starting points. The many interchain contacts in the simulations support the conclusion that the CTDs form a mobile network of interactions that connect the entire Hfq hexamer. The effect of this connectivity is illustrated in SI Appendix, Fig. S9 showing how changes in the interactions of one CTD affect the movements of other CTDs.

**RNA Binding Stabilizes Hfq, CTD but Destabilizes WT Hfq.**

Because nucleic acids compete with the CTDs for binding to Hfq’s rim (11), RNA binding has the potential to alter the organization of the CTDs around Hfq. As a result, the stability and fragmentation pattern of the HfqRNA complexes are expected to provide information on how the bound RNA perturbs the connections between Hfq subunits.

To determine whether RNA binding perturbs the structure of Hfq, we designed a series of short RNAs that mimic the Hfq binding motifs in natural sRNAs and mRNA targets of Hfq (Fig. 4 A). The designed RNAs interact with different surfaces of the Hfq hexamer (SI Appendix, Fig. S10). rA18 and rA18G bind two or six subunits on the distal face of Hfq (31). rUC and rAU5G mimic the sRNA 3’-end and contact five or six subunits around the proximal inner pore (32–34). rCU2C2 and rim-SL, which contains rCU2C2 plus a stable stem loop (SL), mimic RNA motifs that interact with the arginine patch on the rim (10). Interestingly, this loss was also substantial for RNAs binding the distal face, even though only the first few residues of the CTDs are close to this surface (10).

Next, we studied how RNA binding affected the dissociation pathways of Hfq and HfqΔCTD, by determining which subcomplexes retained the RNA after hexamer dissociation (SI Appendix, Fig. S13A). Comparison of the dissociation pathways revealed similar fragmentation of RNA complexes with Hfq and with HfqΔCTD (SI Appendix, Fig. S13 B and C). This result suggested that RNA binding stabilizes the core of Hfq and that perturbation of the CTDs likely comes from their displacement from the core and not from direct interactions with the RNA. CTD displacement was greatest for the largest RNA (RybB sRNA), as the ERMS for Hfq and HfqΔCTD almost overlapped (Fig. 4F). The ERMS of shorter RNA complexes were closer to the ERMS of the free proteins, indicating that binding required less CTD displacement (Fig. 4B and D).

**Progressive RNA Binding Displaces the CTDs from Hfq.**

The design of our RNAs allowed us to investigate the interplay between Hfq, the disordered CTDs, and RNA–protein interactions as an sRNA progressively binds to the protein (Fig. 5 A and B). For this, we compared the stabilities and dissociation pathways of WT Hfq and HfqΔCTD in the absence and presence of RNAs mimicking a stepwise binding process (Fig. 5B). On the one hand, we found that as the RNA interacts with more surfaces of Hfq, the RNA conferred stability to the protein core (Fig. 5C, HfqΔCTD). However, RNA binding was accompanied by a loss of favorable CTD interactions (Fig. 5C, HfqΔCTD).
CTD = Hfq\textbullet RNA-Hfq\textbullet CTD\textbullet RNA; see also Fig. 4G that reduced stability overall (Fig. 5 C, Hfq). Progressive RNA binding also shifted the dissociation products to increasingly larger fragments (Fig. 5 D). This dissociation pattern was the same for Hfq and Hfq\textbullet CTD, indicating similar RNA interactions with the core in both proteins. Thus, stable interactions between RNA and the core are established as the CTDs are displaced.

**Discussion**

In this work, we employed nMS-SID to analyze Hfq complexes of defined stoichiometry, stability, and shape. The results demonstrate that the CTDs connect the Hfq subunits, stabilizing the hexamer (Fig. 2 C and G and SI Appendix, Fig. S13A). MD simulations using an improved force field for IDPs (37) suggest an explanation for this stabilization by revealing that each CTD can dynamically interact with several other CTDs and folded domains (Fig. 3 and SI Appendix, Figs. S4, S5, and S8).

Although RNA binding disrupts the network of CTD interactions, RNA also stabilizes the folded core of Hfq. RNAs that form multiple favorable contacts with Hfq offset the loss of stabilization by the CTDs, while RNAs that do not form favorable contacts are displaced by the CTD interactions, explaining how the CTDs help Hfq discriminate among different RNAs (11, 22).

Based on our results, we propose that RNAs stably bind Hfq through a stepwise process (Fig. 5 E). An RNA may first take...
advantage of configurations in which an Hfq region is exposed, or a CTD is loosely folded. The initial interaction with a segment of the RNA perturbs nearby CTDs. Because the CTDs form multiple contacts with the core and each other (Fig. 3 B and C and SI Appendix, Figs. S7 and S8), perturbations propagate to the rest of the hexamer (SI Appendix, Fig. S9), resulting in more CTD displacement and further RNA binding. If binding is favorable, the hexamer is stabilized. However, if binding is not favorable, the hexamer is destabilized, making it more favorable for the CTDs to regain their interactions with the rest of the protein. This search for stability could explain the removal of weakly bound RNAs when the CTDs are present (11).

The current results agree with previous binding experiments reporting a higher prevalence of nonspecific RNA binding to HfqΔCTD and lower populations of kinetically stable complexes (22). Additionally, even a flexible network of CTDs could facilitate sRNA competition and the search for mRNA targets, as binding of a second RNA will also perturb the CTDs, stimulating the disassembly of noncognate ternary complexes. Finally, the varied CTD conformations impart asymmetry to the Hfq homohexamer that may also contribute to the selection of RNA ligands.

Our nMS-SID results showed that the CTDs make the Hfq hexamer more resistant to dissociation by collision (Fig. 2C). This observation agrees with previous collision-induced dissociation MS experiments showing that *E. coli* Hfq is more stable than Hfq from *Vibrio cholerae*, which has a 22-residue CTD (38). Vincent et al. (38) attributed this stabilization to packing of the CTDs along the intersubunit interfaces. Based on the possibility that the CTDs unfold before the subunits dissociate (Fig. 2 G and SI Appendix, Fig. 51) and MD results showing the CTD interact with multiple subunits (Fig. 3 B and C and SI Appendix, Figs. S4 and S5A), we propose that the Hfq hexamer is additionally stabilized by these intersubunit interactions that are facilitated by longer CTDs. Future experiments with longer or shorter CTDs will help elucidate how variable CTD contacts influence Hfq’s stability.

Network connectivity can explain why the binding of even short RNAs reduced the CTD’s stabilizing contribution (Figs. 4 D–G and 5C): Short RNAs such as rU6 still contact all six Hfq subunits and likely reduce the number of allowed CTD configurations, as suggested by the change in the shape of the ERMS plot (Fig. 4D). However, short RNAs may perturb fewer CTDs, or allow reestablishment of CTD contacts lost after RNA binding, explaining why they reduce the CTD contribution less than long RNAs. Since RNA-bound HfqΔCTD and Hfq fragmented similarly (Fig. 5D and SI Appendix, Fig. S13), stable RNA binding seems to involve contacts with the core and not the CTDs, as previously proposed (13, 30, 39). Finally, destabilization of Hfq but not HfqΔCTD when rA18 RNA is bound (Fig. 4C) supports a reported functional link between the distal RNA binding face and R66 at the start of the CTD (23).

Hfq is a model protein with disordered regions that act synergistically to communicate perturbations among its subunits. This feature is enabled by an architecture of identical monomers, each providing an identical disordered region. The single-stranded DNA binding (SSB) protein, a homotetramer involved in DNA repair, replication, and recombination (40), also contains disordered CTDs with parallels to Hfq. They impart stability to the SSB tetramer (41) but are displaced upon partial DNA engagement, modulating binding and SSB oligomerization (42). Importantly, partial deletion of SSB’s CTDs results in impaired activity, indicating a critical role for multiple CTD interactions in cellular function (43). Histone tails are also thought to fold heterogeneously around the DNA (44, 45) and to be disrupted by chaperone binding or posttranslational modification (46). It would be interesting to know if other RNA-binding proteins use interconnect IdPs to integrate the molecular interactions within RNA–protein complexes.

**Materials and Methods**

**RNA Preparation.** RybB RNA was transcribed using phage T7 RNA polymerase followed by 8% polyacrylamide gel purification (8 μM urea). The remaining short RNAs used in the study were purchased from IDT with high-performance liquid chromatography purification. See SI Appendix, Table S3 for RNA sequences.

**Protein Expression and Purification.** HfqΔCTD and Hfq were purified as described before (21).

**Sample Preparation for nMS.** Protein (HfqΔCTD and Hfq) and RNA (rA18, rim SL-U6, and RybB) samples were dialyzed overnight into 500 mM ammonium acetate, pH 6.8 (99.99%, MilliporeSigma), with eight buffer exchanges (3.5 kDa mass cutoff microdialysis; Pierce). This ionic strength prevented precipitation of protein–RNA complexes at the concentrations (10 μM hexamer) required for ion mobility MS. The remaining RNA samples were used as supplied by the manufacturer and did not require additional desalting. Protein–RNA complexes were prepared by mixing 1:1 RNA and HfqΔCTD or Hfq to a final concentration of...
10 μM each; 400 mM ammonium acetate (final concentration) plus triethylammonium acetate (1 M, MilliporeSigma; 100 mM final concentration for charge reduction) were subsequently added to the samples.

MS. All samples were introduced into the mass spectrometer using nanoelectrospray emitters that were prepared in-house using a Sutter P-97 micropipette puller. All spectra in this work were acquired on a Waters Synapt G2 HDMS instrument (Waters Corporation) modified with an SID device between a shortened trap stacked ring ion guide and an ion mobility cell, as described previously (47). SID lenses can be tuned either to transmit ions for MS or to direct the ions onto the surface for collision. Typical voltage settings and instrument parameters used here for transmission mode and SID can be found in SI Appendix, Tables S4 and S5). ERMs were produced by acquiring data from tandem MS experiments with SID voltage potentials ranging from 15 and 140 V. Each experiment was repeated in technical triplicate. Additional information is provided in SI Appendix.

Analysis of MS data. Ion mobility was used to separate product ions and selection rules for each SID product were made using Waters Corporation Driftscope v1.3 (48). CE were calculated as $E(eV) = zV$, where $z$ is the charge state of the precursor ions and $V$ is the SID voltage. ERMs were corrected by $m_{SID\,CTD}/m_{CTD}$ and $m_{protein}/m_{protein\,RNA}$ (see also SI Appendix, Eqs. S1–S3). Additional information is provided in SI Appendix.

MD Simulations. All simulations were performed with the MD program OpenMM (49) and CHARMM36m force-field (50). Simulations were started from the Protein Data Bank ID 1HK9, which included residues 7 to 68 (29). The start-

different initial velocities. All protein structures were embedded in a water box and neutralized with 150 mM NaCl. Additional simulation and setup details are provided in SI Appendix. Following a 100-ns equilibration, the models were run for an additional 1,400 ns (models 1, 3, and 5), 400 ns (models 2 and 4), 100 ns (models 6 to 10), 1,000 ns (e1 and e4), or 10 ns (e2 and e3; SI Appendix, Table S2). To gain information about the short-term structures and their evolution, we analyzed contacts between CTDs and cores of various subu-

nits during various time intervals of the simulation (SI Appendix, Figs. S7 and S8). To characterize the CTDs dynamics, we calculated root-mean-square deviation values for chosen individual CTD residues (SI Appendix, Figs. S5 and S6). Additional information is provided in SI Appendix.

Data, Materials, and Software Availability. MD simulation trajectories and Energy Resolved mass spectra raw data have been deposited in the Johns Hopkins Data Archive Dataverse Network (https://doi.org/10.7283/t/11RT6G00) (51).

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