Membrane association of the bacterial riboregulator Hfq and functional perspectives

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Hfq is a bacterial RNA binding protein that carries out several roles in genetic expression regulation, mainly at the post-transcriptional level. Previous studies have shown its importance in growth and virulence of bacteria. Here, we provide the direct observation of its ability to interact with membranes. This was established by co-sedimentation assay, cryo-transmission electron (cryo-TEM) and atomic force (AFM) microscopies. Furthermore, our results suggest a role for its C-terminus amyloidogenic domain in membrane disruption. Precisely, AFM images of lipid bilayers in contact with Hfq C-terminus fibrils show the emergence of holes with a size dependent on the time of interaction. Cryo-TEM observations also show that liposomes are in contact with clusters of fibrils, with occasional deformation of the vesicles and afterward the apparition of a multitude of tiny vesicles in the proximity of the fibrils, suggesting peptide-induced breakage of the liposomes. Finally, circular dichroism spectroscopy demonstrated a change in the secondary structure of Hfq C-terminus upon interaction with liposomes. Altogether, these results show an unexpected property of Hfq and suggest a possible new role for the protein, exporting sRNA outside of the bacterial cell.

The bacterial Hfq protein was identified in Escherichia coli as an abundant RNA-binding protein about fifty years ago. Today Hfq is recognized as the core component of a global post-transcriptional network that facilitates the imperfect base-pairing of small regulatory noncoding RNA (sRNA) with trans-encoded mRNA targets. Consequently, Hfq induces the repression or activation of translation, with important related consequences for RNA stability. This mode of regulation is of primary importance for the adaptation of bacteria to the changing environment, and thus for the control of cell division and for the virulence of pathogenic species. The ability of Hfq to interact with DNA has also been demonstrated and it has been described as one of the E. coli nucleoid associated proteins (NAP) in vivo. Nevertheless, the fraction of Hfq within the nucleoid represents only 10–20% of the total protein concentration, while approximately 50% of the protein is found in close proximity of the membrane. This makes sense as the majority of Hfq-regulated sRNA targets encode membrane proteins.

Regarding its molecular structure, E. coli Hfq forms an Sm-fold in its N-terminal region (NTR, 2/3 of the 102 amino acid residues protein). This fold consists of a five-stranded antiparallel β-sheet capped by an N-terminal α-helix. The β-sheets from six monomers interact with each other to assemble in a toroidal structure with two different surfaces. It appears that the distal face and the edge of the protein are involved in both DNA and RNA binding, while the proximal face (on which the α-helix is exposed) seems to be involved in RNA binding only. Besides its Sm-like domain, other regions of Hfq such as the C-terminal region (CTR) also play a role in nucleic acid recognition and binding. The Hfq three-dimensional structures of various bacteria have been resolved, but until now all lack the CTR, so the way this Hfq region folds remains enigmatic.

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Structurally, the CTR seems to extend outside the Sm-core and appears to be intrinsically disordered. This domain is also presumably important for protein stability. Recently, it has been shown that the CTR region of *E. coli* Hfq contains an amyloid sequence. This allows the Hfq protein to self-assemble, a feature common to Hfq and Sm proteins.

In this work, we focus our attention on a new unexplored feature of Hfq, namely its ability to bind to membranes. This finding opens new perspectives in Hfq-dependent riboregulation that will be discussed herein.

**Results**

**Hfq CTR and NTR domains both interact with lipid bilayers.** In order to have a broad view of the ability of Hfq to interact with lipid bilayers, several lipids were first tested for potential interactions by co-sedimentation assays. In all cases, we observed that the protein alone at the concentration specified did not sediment in the conditions used. Results are presented in Fig. 1. The wild-type Hfq hexamer has a pronounced affinity for dioleoylphosphatidylglycerol (DOPG) polar lipids, which are present in the natural inner membrane of *E. coli* and therefore in the *E. coli* polar extract (EPE) mixture, partially explaining the good affinity for the EPE small unilamellar vesicles (SUVs) (Fig. 1a). Only a small interaction is observed with dioleoylphosphatidylcholine (DOPC), an example of less polar lipids. This result is however not surprising considering the highly hydrophilic and positively charged *E. coli* Hfq surface. A comparable behavior is obtained with truncated Hfq-NTR65.

We also investigated the Hfq-CTR38 propensity to interact with SUVs, knowing its tendency to aggregate into amyloid fibers and the natural property of such a domain to interact with membranes. Results with the 38 amino-acid peptide corresponding to Hfq CTR tail are presented in Fig. 1c, showing a significant binding of CTR peptide to EPE lipids. Note that this interaction is promoted by the presence of NaCl, and the opposite is observed for the NTR domain alone, thus suggesting a different mode of interaction.

**Clusters of Hfq are in direct contact with lipid bilayers.** As seen by Cryo-electron microscopy (cryo-TEM), SUVs formed from EPE or DOPG lipids are round-shaped with a mean diameter ~ 120 nm (Fig. 2a and b). The membrane bilayer with the expected thickness of 4.1 ± 0.3 nm is clearly observed (in agreement with previous reports). Liposomes were sparsely distributed over the grid and are mostly found isolated. Mixing SUVs with full-length WT Hfq induces SUVs clustering. The liposomes are decorated with proteins, leading to a significant
Figure 2. Cryo-microscopy images and tomography of wild-type Hfq bound to SUVs. (a) Liposomes composed of 100% EPE lipids show a clean bilayer. (b) Same as (a) but with DOPG lipids. (c) EPE liposomes incubated with 10\(\mu\)M Hfq. (d) DOPG liposomes incubated with 10\(\mu\)M Hfq. (e) and (f) Same as (c) and (d) with lower magnification, respectively. (g) Cryo-tomography of Hfq and DOPG SUVs showing a bilayer completely covered by proteins, which eventually form aggregates visible on the surface. (h) Tomography of double DOPG liposomes showing that only the outer membrane (2) is decorated by Hfq, and is thicker. The inner membrane (1) has an undecorated bilayer. (i) Grey intensity profile is displayed according to the red arrow on image (h), showing the thickness difference between liposome 1 and 2. Scale bars: 20 nm.
increase of membrane thickness: 7.6 ± 1.0 nm or 8.2 ± 1.2 nm for EPE or DOPG liposomes, respectively (Fig. 2c–f). A subtraction between the respective membrane thickness of uncoated and coated liposomes gives a layer of protein measuring approximately 3–3.5 nm for EPE. As Hfq hexamer measures about 3 nm in height, this suggests that hexamers are lying horizontally on the membrane. Note that the protein layer is not completely flat, reflecting a complex arrangement, and that the bilayer is sometimes no longer visible, suggesting that Hfq may sometimes affect the membrane surface. Furthermore, proteins cover almost all the DOPG liposomes, showing some thicker stacks, whereas the EPE SUVs are not always decorated (Fig. 2e). This is consistent with the apparent higher affinity for DOPG observed by co-sedimentation. In order to evaluate the global shape and protein coating of SUVs, we also performed cryo-tomography. 3D reconstruction gives very thick liposomes, normally shaped and fully decorated by proteins, in accordance with TEM results (Fig. 2g). Occasionally, a small SUV is trapped inside a larger one coated by Hfq (Fig. 2h), allowing clear observation of the different thicknesses; measurement of the membranes thickness along the red arrow gives approximately 3.87 nm for the inner liposome and 6.82 nm for the outer one, in accordance with measurements obtained on projections (Fig. 2i).

To prove that Hfq promotes liposome-liposome contacts, we then used C-terminal His-tagged Hfq in the presence of Ni²⁺-functionalized gold beads, to denote the position of the proteins (Fig. 3a). Note that in this condition, the His-tagged protein concentration is three times lower than the wild-type Hfq concentration used in Fig. 2. Control experiments showing beads and liposomes without protein (Fig. S1a) and beads weakly bound to the natural histidines of the wild-type protein (Fig. S1b) are provided as supplementary data. As seen, gold beads are not randomly coating liposomes but clustered, suggesting that proteins are organized in clusters (possibly fibers) at the liposome-liposome interfaces. This observation is consistent with the partial decoration of liposomes, indicating that Hfq protein tends to bind preferentially to protein-lipid clusters already formed (Fig. 3b and highlight in Fig. 3c). This observation may suggest that Hfq binding to membranes is cooperative.

In order to confirm this result, we then used atomic force microscopy (AFM) in solution to observe Hfq behavior on supported lipid bilayers. As expected, after a short incubation time on an EPE bilayer, proteins are distributed over the surface in small aggregates, compatible with hexamers regrouped in clusters (Fig. 4a). The aggregates have an average height of 2.7 ± 0.2 nm, obtained from measuring the profile of 20 different aggregates.
Figure 4. AFM images of wild-type Hfq at 2 µM (a,b,d,e), Hfq-NTR$_{65}$ at 2 µM (c,f), or Hfq-CTR$_{11}$ at 176 µM (g-l) in solution on an EPE supported bilayer. (a) After 5 min incubation, Hfq binds on the top and forms groups of several hexamers. (b) After 1 h incubation, the bilayer is affected and 1–2 nm deep holes appear, containing undetermined material inside. (c) After 1 h incubation, no trace of Hfq-NTR$_{65}$ is visible and the membrane is clean. (g) Clean bilayer before adding Hfq-CTR$_{11}$. (h) After 30 min incubation, small 4 nm deep holes start to form. (i) After 1 h incubation, holes have grown to several hundreds of nm wide, containing undetermined material inside. (d, e, f, j, k and l), height profiles of grey lines on images a, b, c, g, h and i, respectively. Protein concentrations were chosen in order to have similar mass concentrations.
than full length Hfq, Hfq-NTR 65 was able to regroup liposomes, but no visible protein decoration is observed on SRCD spectra were obtained for freshly dissolved Hfq-CTR38 in solution and Hfq-CTR 38 incubated with SUVs the Hfq-CTR is interacting with SUVs, we performed Synchrotron Radiation Circular Dichroism (SRCD). The were synthetized according to Waltz algorithm predictions34 (see Table S1 for sequences). Self-assembly of Hfq was identified precisely. To that end, various peptides corresponding to overlapping sequences in the CTR protein adsorbs weakly on the membrane and is easily washed away when rinsing with buffer before imaging, or undergo reorganization and holes formation even after a long incubation time. It suggests that the truncated protein adsorbs weakly on the membrane and is easily washed away when rinsing with buffer before imaging, or pushed by the AFM tip. In order to understand which part of the protein is involved in the hole-formation observed with the full-length protein, we then investigated the effects of the amyloidogenic Hfq-CTR, which is also able to interact with EPE SUVs (Fig. 1c).

Identification of the amyloid region in Hfq-CTR. First, the amyloid region within CTR domain of Hfq was identified precisely. To that end, various peptides corresponding to overlapping sequences in the CTR were synthesized according to Waltz algorithm predictions34 (see Table S1 for sequences). Self-assembly of these peptides was tested by cryo-TEM and positive results are summarized in Fig. S4. This allowed us to determine that a short sequence of 11 amino acids residues is responsible for the self-assembly, precisely the region SAQNTSAQQDS. We also confirmed by FTIR spectroscopy35 that this small peptide self-assembles into amyloid fibrils. This 11 amino-acid residues peptide will be referred to as Hfq-CTR 11.

Hfq-CTR 11 amyloid fibers trigger liposome breaking and membrane disruption. The same experiments carried out previously with the native protein were repeated with Hfq-CTR 11. In this case, fibrils can be clearly observed with Cryo-TEM (peptide concentration 440 µM, i.e. 0.5 g.L−1) (Fig. 5). Stacks of fibrils decorated with liposomes form along a specific direction, different for each cluster (Fig. 5a). Liposome deformations at the liposome-fibril contact regions are frequent (red arrows Fig. 5b and c). Furthermore, many small liposomes of 10–30 nm appear (blue arrows Fig. 5), resulting from the rupture of larger SUVs. Note that, although the cryo-TEM indicates an interaction between fibrils and liposomes, it is not possible to determine from individual projections whether overlap occurs in the same plane of the sample. Therefore, cryo-electron tomography has been used to reconstruct the 3D structure of the interaction between fibrils and liposomes (Fig. 5d and e). Tomography images directly confirm contacts between the two components. Furthermore, analysis of the liposome morphology also demonstrates that some of the liposomes are deformed by fibrils (Fig. 5d) and small 10–30 nm liposomes are also observed (Fig. 5e).

Then, the effect of Hfq-CTR 11 on EPE mica-supported lipid membranes was observed by AFM. Before protein incubation, they presented a roughness under 1 nm (Fig. 4g and i). After addition of Hfq-CTR 11, small holes clearly appear (Fig. 4h) that grow with incubation time (Fig. 4i). The measured depth of these holes is 4 nm, which means that unlike the native protein, the Hfq-CTR 11 peptide removes the whole bilayer (Fig. 4k and l), a result compatible with the SUV deformation and splitting observed in TEM. Again, we assume that protein/lipid aggregates are likely present in the holes formed (Fig. 4i). But in this case no aggregated protein is found adsorbed on the intact membrane area.

Interaction with membrane induces a conformational change in Hfq CTR. Finally, to assess how the Hfq-CTR is interacting with SUVs, we performed Synchrotron Radiation Circular Dichroism (SRCD). The SRCD spectra were obtained for freshly dissolved Hfq-CTR38 in solution and Hfq-CTR 38 incubated with SUVs (Fig. 6). The respective secondary structure compositions were determined using Bestsel and are reported in Table 1. As expected from our previous observations36, the peptide alone is partially unstructured or folded in β-sheets (38%). When incubated with EPE liposomes, the percentage of β-sheets decreases to 33%, while when incubated with DOPG liposomes, the percentage of β-sheets is reduced to 28% with a small increase of turns and apparition of 5.6% α-helices (classified as “distorted helix” category according to Bestsel). In the later case, it is likely that only a fraction of the peptide is interacting the membrane and that more than 5–6% helices is formed in the population bound to membrane. Note that the helices formed seem to be unstable without the interaction of the peptides with the polar phosphoglycerol lipid heads.

Discussion
Our results demonstrate that Hfq has a propensity to interact with membranes. Importantly, the native protein used for all experiments here was exempt of post-translational modification such as lipid addition. This has been confirmed by mass spectrometry analysis (Fig. S5). Thus, such a modification is not needed for interaction of the protein with a bilayer membrane in vitro, even if the existence of this modification has been reported previously36.

Full length Hfq affects the bilayer organization, but not HfqNTR 65. Surprisingly, after longer incubation times, the membrane exhibits a drastic reorganization. Lipid domains appear, which is comprehensible when a positively charged protein interacts with anionic phospholipids present in the EPE33. However, we also observed the presence of numerous unexpected holes with a diameter around 100 nm and 1–2 nm in depth (Fig. 4b). As the membrane treated in the same condition in the absence of Hfq does not present such a reorganization on the same timescale, this indicates that Hfq affects the membrane integrity. Note that some undetermined material is still present inside the deep holes, which does not correspond to the ~4 nm thickness of the bilayer (Fig. 4e). We assume this could be Hfq or Hfq-lipids aggregates resulting from the membrane reorganization.

We then performed the same set of experiments with Hfq-NTR 65. Cryo-TEM revealed that, to a lower extent than full length Hfq, Hfq-NTR 65 was able to regroup liposomes, but no visible protein decoration is observed on the surface (Fig. S3). Furthermore, AFM with Hfq-NTR 65 gives different results than with Hfq-WT, closer to a


For Hfq-membrane interaction, two distinct actions may be described. First, a relatively weak, reversible binding with the torus lying on the membrane is observed. The affinity of the native protein for polar lipids suggests that binding is mainly driven by hydrogen bonding and electrostatic interactions. The importance of weak electrostatic interactions between basic residues and acidic lipids could be considered as a mean for proteins to be transiently released in the cytoplasm and could explain why Hfq was not identified in the membrane fraction in large-scale proteomic studies.

The self-assembly of Hfq through its C-terminal tails at the bilayer surface also provides a possible explanation for the superstructures observed previously in vivo in close proximity of the inner membrane. The C-terminal tails strengthen the protein-lipid interaction. As membrane association is lost when this CTR is

Figure 5. Cryo-TEM images and tomography of Hfq-CTR11 incubated with EPE SUVs. (a) Hfq-CTR11 peptides aggregate to form amyloid fibrils and regroup large clusters of liposomes. (b) A high proportion of small liposomes of 10–30 nm (blue arrows) are present, possibly coming from SUVs disruption by Hfq-CTR11. (c) Multiple contacts between liposomes and fibers create SUV deformations (red arrows). (d) Liposome deformation as seen by cryo-tomography. (e) Small liposomes (10–30 nm) as seen by cryo-tomography. Scale bars: 200 nm for (a), 50 nm for (b,c) and 20 nm for (d,e).
deleted, we assumed this part of the protein was essential for the interaction with membrane in vivo. Our present results with Hfq CTR region confirm this assumption. This result is also in agreement with previous works on the ability of fibrils including amyloid fibrils to interact with membranes, also in the case of peptide-forming fibers. The second action we have observed is a drastic change in the membrane organization with longer incubation times. This rearrangement is induced by the C-terminal amyloidogenic portion of the protein. AFM measurements suggest that the full-length protein remove one monolayer of the membrane, in agreement with the recent work of Pyne et al. Taking into account that monolayer poration has already been reported for other proteins, one possibility is that similarly to Tilamin, Hfq-induced monolayer-pores may cause local membrane rupture. Nevertheless, in the case of Tilamin, monolayer poration induces progressive removal of the outer membranes and bacterial death. More likely in the case of Hfq, the protein induces transient pores in vivo and not membrane disintegration. We also confirm here the ability of amyloid regions to disrupt and distort membranes.

Finally, the secondary structural change of Hfq-CTR observed by SRCD suggests that the amyloid-like region could be partially converted into a α-helix, possibly a trans-membrane helix as reported earlier for other amyloids. Another possibility would be that an amphipathic helix lying horizontally in the acidic phosphate-heads layer forms, such as that observed for RNase E or RNase H. This would be in agreement with the flexibility of Hfq CTR and with its composition of both basic/polar (H, Q, N, T) and non-polar (A, Y) amino acid residues. Alternatively, a tilted helix may also form as in the case of Tilamin, resulting in its insertion in the monolayer. These possibilities however need to be investigated further, for instance using orientated circular dichroism (OCD) to observe the orientation of the helix inside the membrane.

In conclusion, the key finding of this work is that Hfq is able to affect biological membranes, as shown in AFM and cryo-TEM. This finding may support a possible mechanism by which Hfq could contribute to the export of RNA outside the bacteria or in the periplasmic space. To date, known exported molecules for quorum sensing were limited to small molecules. Nevertheless bacterial RNAs have been identified extracellularly. Among those RNAs, small noncoding RNA are found. Here we propose a new possible role for Hfq in sRNA export, with important consequences for bacterial communication.

Materials and Methods

Protein expression and purification. Wild type Escherichia coli Hfq was purified as described previously. Hfq-NTR (residues 1–65) was purified from over expressing BL21(DE3)Δhfq/pLATE11-hfqNTR cells. The pLATE11-hfqNTR expression vector was constructed according to the manufacturer protocol.
(Thermofisher). As the his-tag potentially modifies Hfq properties	extsuperscript{31}, a sequence encoding the Tobacco Etch Virus (TEV) protease site (ENLYFQG) was inserted between the N-terminal His-tag and Hfq sequence. For the purification of Hfq-NTR\textsubscript{65}, cells from post-induction cultures (IPTG 1 mM, 3 h induction) were resuspended on ice in 20 mM Tris HCl pH 7.5 containing 0.5 M NaCl and 20 mM imidazole; the protein was eluted with a gradient of imidazole (20–500 mM) in the same buffer. The eluted protein was dialyzed in the appropriate buffer and TEV digestion was carried out according to the manufacturer’s instructions (Thermofisher). After digestion, Hfq-NTR\textsubscript{65} was subsequently purified with a HiTrap SP HP 1 mL column (GE Healthcare). Before injection on the cation exchange column, Hfq-NTR\textsubscript{65} was diluted five times in equilibration buffer (50 mM HEPES pH 8) to reduce salt concentration. The resin was equilibrated with the same buffer and the protein was eluted with a gradient of NaCl (0–1 M). The cleaved His-tag, non-digested Hfq-NTR\textsubscript{65} and other remaining contaminants are eliminated with this cation exchange column.

Hfq C-terminus peptides were obtained from Proteogenix (France). Fibrils of synthetic peptides were prepared from dissolved peptide in deionized water at 0.5 g.L\textsuperscript{-1}. Fibrils were usually observed after one week. Note that kinetics of assembly can be greatly affected by sample batch-to-batch variability. Several factors contribute to this variability, including the presence of salts in batches and the presence of various amounts of pre-formed aggregates in samples.

**Preparation of small unilamellar vesicles (SUVs).** Lipids purchased from Avanti Polar Lipids (Alabaster, AL, USA) were dissolved at 10 mg.mL\textsuperscript{-1} in chloroform/methanol 1:1 (v/v) on ice. The lipids used were E. coli Polar Extract (abbreviated EPE, ref 100600P), 1,2-dioleoyl-sn-glycero-3-phosphocholine (abbreviated DOPC, ref 859375) and 1, 2-dioleoyl-sn-glycerol-3-phospho-(1’-rac-glycerol) (abbreviated DOPG, ref 840475). The solvent was slowly evaporated with N\textsubscript{2}(g), forming a lipid film that was then dried for 30 min under N\textsubscript{2} flow. Lipids were hydrated in SUV buffer (10 mM Tris pH 7.5 containing 100 mM NaCl). Vesicles formed during slow stirring at room temperature over 30 min. The solution was then passed 31 times through a 0.2 μm polycarbonate filter (Avanti Mini Extruder) to obtain SUVs. Samples were kept at 4 °C and used for subsequent experiments within one week. For liposomes used in CD experiments, the protocol was identical but Tris-HCl was replaced by phosphate buffer (10 mM sodium phosphate buffer pH 7.5 with 100 mM NaCl) and using 0.1 μm polycarbonate filter for extrusion.

**Co-sedimentation assay.** Gradual concentrations of SUVs (100, 300 and 600 μM lipids) were mixed with proteins in SUV buffer with or without NaCl and left 20 min at 22 °C before being ultracentrifuged 1 h at 100 000 g at the same temperature in a Beckman TL-100 ultracentrifuge with a Beckman TLA100 rotor. 10% of total sample volume were recovered from the supernatants and pellets which were then loaded on polycrylamide gels for SDS-PAGE. Gels were stained with R-250 or G-250 Coomassie Brilliant Blue.

**AFM imaging in solution.** SUV suspension diluted at 0.4 g.L\textsuperscript{-1} in SUV buffer supplemented with 2 mM CaCl\textsubscript{2} was incubated on a freshly-cleaved mica surface for one hour at 37°C. The sample was then rinsed ten times with SUV buffer to remove excess SUVs. Supported bilayers were checked by AFM to confirm the extent of surface coverage and the quality of the lipid bilayer on the mica before protein incubation.

The proteins were added to the surface at a given concentration at room temperature for various times (from 5 min to several hours). Incubations were done in a sealed box to avoid evaporation. The sample was then rinsed ten times to remove unbound proteins and imaged in the same buffer. Protein concentrations were 2 μM for both native Hfq and Hfq-NTR\textsubscript{65} and 176 μM for Hfq-CTR\textsubscript{11} peptide (1136 g/mol). Protein concentrations were chosen in order to have similar mass concentrations.

In this work, commercial Olympus rectangular silicon nitride cantilevers (RC800PSA) and two different commercial AFMs were used: an Agilent technologies 5500 microscope (Santa Clara, CA, USA) operated in the tapping mode at a resonance frequency of 15 kHz and 0.73 N.m\textsuperscript{-1} spring constant cantilever and a Nanotec microscope (Madrid, Spain) operated in the jumping mode with an applied force <100 pN\textsuperscript{52}. After the sample was installed on the AFM stage, about 30 min were required for the system to reach thermal equilibrium before commencing the AFM scans. We imaged 5 to 10 positions at different resolutions to ensure consistency among the observations.

Before obtaining the height profiles from the AFM images, the tilt of the surface and the line to line noise were removed using the plane and flattening filters provided in the WSxM free software (www.wsxmsolutions.com)\textsuperscript{53}. The height profiles were all obtained from line scans perpendicular to the slow scanning direction.

**Cryo-Electron microscopy (cryo-TEM).** SUVs in SUV buffer (0.2 g.L\textsuperscript{-1} final concentration) and protein in the same buffer were mixed and left at room temperature for 20 min. When necessary, 5 nm Ni-NTA-NanoGold beads (NanoProbe, Yaphank, NY) were added 2 minutes before freezing at a final concentration of 4 nM in order to label His-tagged proteins. The high clustering observed in cryo-TEM images originates from the fact that Ni-NTA gold beads possess multiple Ni-NTA moieties that can bind several liposomes together. Protein concentrations were 10 μM for both Hfq-WT and Hfq-NTR\textsubscript{65}, and 440 μM for the Hfq-CTR\textsubscript{11} peptide. Protein concentrations were chosen in order to have similar mass concentrations.

5 μL of samples were then deposited on air plasma-cleaned EM grids (Lacey carbon films on 300 mesh copper grids). The excess was blotted with a filter paper and the grid was immediately plunged into a liquid ethane bath cooled with liquid nitrogen. Images were recorded on a Gatan Ultrascan camera using a 2200FS transmission electron microscope.
electron microscopy from JEOI Ltd operated at 200 kV. High magnification images were taken over large sections at random locations, in order to avoid statistical bias. Tomographic series were taken from regions of interest covering an angular range from −60° to + 60°. Statistics on membrane thickness were calculated from n = 20 pictures.

Images were processed with ImageJ software and tomogram reconstructions were calculated by the OS-SART method using the TomoJ plugin. Stitching and quantitative analysis of the images were also carried out in ImageJ.

**Synchrotron Radiation Circular Dichroism (SRCD).** Measurements and data collection were carried out on DISCO beam-line at the SOLEIL Synchrotron (Gif-sur-Yvette, France) at 1 nm steps at 1.2 integration time, between 260 and 180 nm were performed in triplicate for the samples as well as for the baselines (consisting of buffer alone or buffer + SUVs). (+)-camphor-10-sulfonic acid (CSA) was used to calibrate amplitudes and wavelength positions of the SRCD experiment. Data-analyses including averaging, baseline subtraction, smoothing, scaling and standardization were carried out with CDTool. Secondary structure content was determined using BestSel. Normalized root-mean-square deviation (NRMSD) indicated the most accurate fit for each spectrum; values of < 0.15 were considered significant.

**Data availability.** All data generated or analyzed during this study are included within this article and the Supplementary information or are available from the corresponding author upon request.

**References**

1. Shapiro, L., Franze de Fernandez, M. T. & August, J. T. Resolution of two factors required in the Q-beta-RNA polymerase reaction. *Nature* **220**, 478–480 (1968).
2. Gottesman, S. & Storz, G. RNA reflections: converging on Hfq and its accomplices. *J Biol Chem* **288**, 7996–8003 (2013).
3. Zambrano, N. *et al.* Involvement of Hfq protein in the post-transcriptional regulation of E. coli bacterial cytoskeleton and cell division proteins. *Cell Cycle* **8**, 2470–2472 (2009).
4. Grippenland, J. *et al.* RNAs of bacterial virulence. *Nat Rev Microbiol* **8**, 857–866 (2010).
5. Azam, T. A. & Ishihama, A. Twelve species of the nucleoid-associated protein from Escherichia coli. Sequence recognition specificity and DNA binding affinity. *J Biol Chem* **274**, 33105–33113 (1999).
6. Wulcz, C. *et al.* Eukaryotic Lsm proteins: lessons from bacteria. *Nat Struct Mol Biol* **12**, 1031–1036 (2005).
7. Brennan, R. G. & Link, T. M. Hfq structure, function, and ligand binding. *Curr Opin Microbiol* **10**, 125–133 (2007).
8. Updegrove, T. B., Correia, J. J., Galletto, R., Bujalowski, W. & Wartell, R. M. E. coli DNA associated with isolated Hfq interacts with Hfq's distal surface and C-terminal domain. *Biochim Biophys Acta* **1799**, 588–596 (2010).
9. Kauer, E. Structure and RNA-binding properties of the bacterial Lsm protein Hfq. *RNA* **10**, 610–618 (2013).
10. Jiang, K. *et al.* Effects of Hfq on the conformation and compaction of DNA. *Nucleic Acids Res* **43**, 4332–4341 (2015).
11. Malabirade, A. *et al.* Condensation and compaction of DNA mediated by the C-terminal domain of Hfq. *Nucleic Acids Res* doi:10.1093/nar/gkj268 In press (2017).
12. Schumacher, M. A., Pearson, R. F., Möller, T., Valentin-Hansen, P. & Brennan, R. G. Structures of the pleiotropic translational regulator Hfq and an Hfq–RNA complex: a bacterial Sm-like protein. *Enmb* **21**, 3546–3556 (2002).
13. Sauter, C., Basquin, J. & Suck, D. Sm-like proteins in Eubacteria: the crystal structure of the Hfq protein from Escherichia coli. *Nucleic Acids Res* **31**, 4091–4098 (2003).
14. Nikulin, A. *et al.* Structure of Pseudomonas aeruginosa Hfq protein. *Acta Crystallogr D Biol Crystallogr* **61**, 141–146 (2005).
15. Sauer, E. & Weichenrieder, O. Structural basis for RNA 3′-end recognition by Hfq. *Proc Natl Acad Sci USA* **108**, 13065–13070 (2011).
16. Stanek, K. A., Patterson-West, J., Randolph, P. S. & Mura, C. Crystal structure and RNA-binding properties of an Hfq homolog from Vibrio cholerae in complex with Qrr1 sRNA. *Acta Crystallogr D Struct Biol* **73**, 294–315 (2017).
17. Vincent, H. A. *et al.* The low-resolution solution structure of Vibrio cholerae Hfq in complex with Qrr1 sRNA. *Nucleic Acids Res* **40**, 8698–8710 (2012).
18. Reich-Fransden, S. M. *et al.* Structural insights into the dynamics and function of the C-terminus of the E. coli RNA chaperone Hfq. *Nucleic Acids Res* (2011).
19. Arluison, V. *et al.* The C-terminal domain of Escherichia coli Hfq increases the stability of the hexamer. *Eur J Biochem* **271**, 1258–1265 (2004).
20. Fortas, E. *et al.* New insight into the structure and function of Hfq C-terminus. *Biochim Biophys Acta* **1805**, 35 (2015).
21. Mura, C., Phillips, M., Koshukhovsky, A. & Eisenberg, D. Structure and assembly of an augmented Sm-like archaeal protein 14-mer. *Proc Natl Acad Sci USA* **100**, 4539–4544 (2003).
22. Arluison, V. *et al.* Three-dimensional structures of Frullbar Sm Proteins: Hfq and Other Sm-like Proteins. *J Mol Biol* **356**, 86–96 (2006).
23. Herrara, A. L., Tomich, J. M. & Prakash, O. Membrane Interacting Peptides: A Review. *Curr Protein Pept Sci* **17**, 827–841 (2016).
Author Contributions

A.M. and V.A. conceived the original idea and designed the experiments. A.M. purified proteins. A.M. and J.S. performed T.E.M. experiments. A.M., V.A. and F.W. performed SRCD experiments. A.M. and V.A. wrote the manuscript. All authors analyzed results and commented on the manuscript.

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Additional Information

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