RNAs actively cycle on the Sm-like protein Hfq

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Hfq, a protein required for small RNA (sRNA)-mediated regulation in bacteria, binds RNA with low-nanomolar $K_d$ values and long half-lives of complexes (>100 min). This cannot be reconciled with the 1-2 min response time of regulation in vivo. We show that RNAs displace each other on Hfq on a short time scale by RNA concentration-driven (active) cycling. Already at submicromolar concentrations of competitor RNA, half-lives of RNA–Hfq complexes are ~1 min. We propose that competitor RNA associates transiently with RNA–Hfq complexes, RNAs exchange binding sites, and one of the RNAs eventually dissociates. This solves the “strong binding–high turnover” paradox and permits efficient use of the Hfq pool.

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The Escherichia coli protein Hfq, identified as a host factor required for replication of bacteriophage Qb [Franze de Fernandez et al. 1968], is conserved and contains the Sm motif typical of eukaryotic Sm and Lsm proteins. Hfq tightly binds RNA with relaxed specificity. When absent, bacteria show pleiotropic effects that can be attributed to Hfq’s function in bacterial small RNA (sRNA) regulation of gene expression [Romby et al. 2006; Brennan and Link 2007; Waters and Storz 2009]. Numerous Hfq–sRNA and Hfq–mRNA interactions have been validated in large-scale screens [Zhang et al. 2003; Sittka et al. 2008] and studied in detail [Geissmann and Touati 2004; Vecerek et al. 2010]. Hfq can increase sRNA–target RNA association rates [Soper and Woodson 2008], stabilize sRNAs [Urban and Vogel 2007], and act as a chaperone by unfolding of an sRNA or mRNA structure [Geissmann and Touati 2004].

The homohexameric Hfq ring displays two faces: proximal and distal. Hfq–RNA interactions show a preference of U-rich for proximal and A-rich RNA sequences for distal face binding [de Haseth and Uhlenbeck 1980a; Mikulecky et al. 2004]. Simultaneous binding may occur on both sides as well, which could facilitate intramolecular base-pairing and regulation [Rajkowitsch and Schroeder 2007].

[Keywords: Hfq, sRNA, antisense RNA, post-transcriptional regulation, RNA chaperone, Lsm protein]

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Structures of Hfq from E. coli, Staphylococcus aureus, and Pseudomonas aeruginosa have been determined by X-ray crystallography [Schumacher et al. 2002; Sauter et al. 2003; Nikulin et al. 2005]. Two co-crystal structures support two distinct binding surfaces: In S. aureus Hfq, AU-G RNA is bound around the inner rim of the proximal face [Schumacher et al. 2002], and E. coli Hfq has oligo-A bound on the distal face [Link et al. 2009].

This study addresses the question of how stable Hfq–RNA binding can be reconciled with Hfq’s function in sRNA-mediated regulation in vivo. Many RNAs bind tightly to Hfq, with $K_d$ values ranging from sub- to mid-nanomolar [Arluison et al. 2004; Geissmann and Touati 2004; Lease and Woodson 2004; Mikulecky et al. 2004; Sittka et al. 2008; Soper and Woodson 2008; Hopkins et al. 2009; Holmqvist et al. 2010]. Thus, if binding-competent RNAs were in molar excess, almost all Hfq would be bound to RNAs. Hfq–RNA dissociation rate constants in vitro are too low to be compatible with a biologically relevant time scale; half-lives of complexes are in the range of a generation time. If newly induced sRNAs typically could access free Hfq after its dissociation from bound RNAs, their activity should be severely delayed. Yet, the time frame from induction of an sRNA to a significant regulatory effect is short [1–2 min] [Massé et al. 2003], and hence sRNAs can acquire Hfq rapidly. This highlights a paradox, with Hfq being tightly sequestered by the intracellular pool of RNAs, contrasted by the need of new sRNAs to rapidly access Hfq. We considered here a conventional cycling model [dissociative/passive] (Fig. 1A) and associative/active cycling (Fig. 1B). In model A, newly synthesized RNA (Fig. 1A, in red) can only bind Hfq after the resident RNA (Fig. 1A, in blue) has dissociated, i.e., the rate of binding of the incoming RNA is limited by the Hfq–RNA dissociation rate constant and is not affected by the concentration of the free RNA. In model B, free RNA transiently binds the Hfq–RNA complex, whereupon one of the RNAs eventually dissociates. Thus, the dissociation rate of the bound RNA is a function of the concentration of the free RNA (Fig. 1B). This would render cycling much more rapidly, and the intracellular pool of binder RNAs would rapidly equilibrate on Hfq. The two models are distinguishable, since the RNA exchange in model A is limited by the dissociation rate constant of Hfq–RNA, whereas exchange in model B increases in proportion to the free RNA concentration.

Results and Discussion

Hfq binds model sRNAs and mRNAs with high affinity

We chose six RNAs from E. coli to determine their binding characteristics, using three different approaches. These were MicA, MicC, and MicF, and their targets, ompA, ompC, and ompF mRNA. Their regulatory interactions are known, as is their Hfq dependence in vivo [Andersen and Delithas 1990; Chen et al. 2004; Rasmussen et al. 2005; Udekwu et al. 2005].

RNA binding at increasing concentrations of Hfq (hexameric Hfq is Hfq throughout this paper) at 37°C was assayed by gel-shift analysis. Supplemental Figure S1 indicates tight Hfq–RNA binding, with $K_d$ values ranging from 0.9 to 4 nM [Table 1]. Binary Hfq–RNA complexes were formed first, and supershifted complexes

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appeared at higher Hfq concentrations [Supplemental Fig. S1]. The K_d values of RNA–Hfq complexes are often subnanomolar [e.g., Geissmann and Touati 2004; Sittka et al. 2008] to mid-nanomolar [e.g., Lease and Woodson 2004; Mikulecky et al. 2004]. For MicA and ompA RNA, additional methods were used for comparison. Surface plasmon resonance (SPR) was employed to assess binding kinetics of either MicA or ompA RNA to Hfq. K_d values and association and dissociation rate constants were calculated [Supplemental Fig. S2]. The association rate constant k_a for both RNAs was »10^6 M^{-1} sec^{-1}, and the dissociation rate constants k_d were 3 × 10^{-4} sec^{-1} to 6 × 10^{-4} sec^{-1} [Supplemental Table S1].

We then used nitrocellulose filter binding to determine dissociation rates and assess competition. MicA* or ompA RNA* (asterisk indicates labeled RNA) was incubated with increasing concentrations of Hfq6, and retention of bound RNA* was monitored. This gave K_d values of 1.7 ± 0.1 nM [MicA–Hfq6] and 1.5 ± 0.5 nM [ompA–Hfq6], similar to values obtained by gel shift and SPR [Table 1].

In these assays, MicA–Hfq6 binding was essentially complete before 20 sec at the concentrations used (1 nM MicA* or ompA RNA*, 10 nM Hfq6). Dissociation was measured by preforming MicA*–Hfq6 or ompA RNA*–Hfq6 complexes, followed by >250-fold dilution and filtering after a further incubation for different time periods at 37°C. Both MicA–Hfq6 and ompA–Hfq6 complexes dissociated slowly, at k_d values of »0.7 × 10^{-4} sec^{-1} [Table 1]. This corresponds to half-lives of >150 min.

RNAs promote concentration-dependent dissociation of Hfq-bound RNA

By filter binding, we asked whether unlabeled competitor RNA in high excess could promote faster dissociation of RNA* from complexes with Hfq. Figure 2, A and B, shows time courses of dissociation. In addition to the RNAs mentioned above, RNA*–Hfq6 was challenged by IstR-1 [Darfeuille et al. 2007], poly-U, and poly-A. Competitor RNA (0.75 μM) was added to 0.75 nM RNA*–Hfq6 complexes, incubated for the times indicated, and filtered. All Mic RNAs, poly-U, ompC, and ompA RNA, caused rapid dissociation of MicA* from Hfq [Fig. 2A], with >50% release within 2–5 min. IstR-1, a poor Hfq binder, had no significant effect, and neither did ompF RNA and poly-A. Similarly, rapid dissociation of ompA RNA* from Hfq was induced by all RNAs but IstR-1 [Fig. 2B]. Note that, although ompF RNA and poly-A failed to compete with MicA* [Fig. 2A], they did dissociate ompA RNA* [Fig. 2B], likely indicating differences in face preference.

The chaser RNA concentration dependence of dissociation was measured by competition experiments on RNA*–Hfq complexes [Fig. 2C,D]. The dissociation rates of RNA*–Hfq complexes increased as a function of competitor RNA concentration, and half-lives of RNA*–Hfq complexes decreased from >150 min (absence of chaser) to »1 min [highest concentrations of chaser RNAs] [Table 1]. These results are incompatible with model A (Fig. 1A) and show that RNAs drive dissociation of RNA*–Hfq complexes much faster than expected from the first-order dissociation rate constants determined in the absence of competitor.

Since RNA*–Hfq dissociation rates are a function of chaser RNA concentration, the reaction is second order. At the high molar excess of chaser used, the reaction follows pseudo-first-order kinetics. Second-order rate constants can be estimated by plotting the apparent k_d against the concentration of chaser RNA [Supplemental Fig. S3]; initial slopes of the curves reflect the association rate of competitor RNA to RNA*–Hfq complexes. The plateaus indicate saturation of binding and a slow rearrangement of RNAs on Hfq before one of them dissociates [see below].

Displacement of Hfq-bound RNAs involves at least two steps

The competitor concentration dependence [Fig. 2C,D] (Table 1) can be approximated with straight lines in the log-log graphs. There are, however, deviations from the assumption of pseudo-first-order kinetics: (1) an initial delay in the decay curves, (2) dissociation rates saturate at micromolar concentrations of competitor, and (3) Hfq-binding plateaus at a nonzero value [Supplemental Fig. S4C]. We developed a mathematical model of the RNA exchange mechanism in which the competitor RNA binds the stable Hfq–RNA* complex in a second-order reaction [Supplemental Fig. S4]. The initially weakly bound RNA can either dissociate rapidly or gradually displace strongly bound RNA* in a series of reversible first-order reactions. When RNA* reaches its weakly bound state, it either dissociates or reverts into a more stably bound conformation. This kinetic model accurately reproduces all aspects of the chase experiment. In particular, the intrinsic consecutive displacement steps limit how fast Hfq can cycle, and explain the initial delay in the dissociation curves and the saturation at high competitor concentration. The nonzero plateau levels reflect the equilibrium binding of RNA* to Hfq, which shows that Hfq does not drive exchange irreversibly in one direction.

The kinetic parameters that fitted the experimental data to the model give quantitative insights into the mechanism; i.e., when MicA RNA* is in the weakly bound state, it dissociates from Hfq at a rate of 0.06 sec^{-1}. However, in the absence of competitor, it only spends <1% of its time in this state, resulting in an overall dissociation rate of <10^{-4} sec^{-1} [Supplemental Fig. S4]. Thus, the chaser RNA concentration dependence supports an active exchange model [Fig. 1B]: The release of the resident RNA...
<table>
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<tr>
<th>RNA</th>
<th>K_d (nM)</th>
<th>Gel-shift assays</th>
<th>Filter binding assays</th>
<th>Surface plasmon resonance</th>
<th>K_d [10^{-4} sec^{-1}]</th>
<th>t_{1/2} [min]</th>
<th>K_d [10^{-4} sec^{-1}]</th>
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*aMean values of at least two independent experiments, each based on at least six independent data points.

*bK_d values were obtained by data fitting to a one-site direct-binding formula [Supplemental Material].

*cThe k_d values were calculated as in the Supplemental Material.

*dHalf-lives of complexes, t_{1/2}, were calculated as ln2/k_d. (n.d.) Not determined.
UNOCUPIED SUBUNIT (Fig. 1C), and dissociation of the
itor RNA gains initial access to only one transiently
binding states (from one to six site occupancy). Compet-
binding assays were performed as described. MicA∗–Hfq (A) or ompA
RNA∗–Hfq (B) complexes were preformed (0.75 nM RNA/7.5 nM
Hfq), and competed by adding a 1000-fold excess of unlabeled RNAs.
The graphs show the retention of labeled RNA by Hfq as a function
of time, using the competitor RNAs indicated. (C,D) Competition of
MicA∗–Hfq by MicA, MicF, and ompC RNA (C), and of ompA∗ RNA–
Hfq by ompA RNA, ompC RNA, and MicC (D), except that compet-
itor RNA was added at the concentrations indicated. (E) MicA∗–ompA
RNA–Hfq complexes (0.75 nM MicA∗, 7.5 nM ompA, 7.5 nM Hfq),
were competed as in A and B. (F) Competition as a function of MicA
concentration was analyzed as in C and D. The dissociation rate
constants and half-lives of complex are shown in Table 1.

is driven by association of other (or the same kind of)
RNAs in a concentration-dependent manner. Based on
published structures, the Hfq hexamer has six identical
binding sites on each surface. High affinity is obtained by
contacts at multiple binding sites (Link et al. 2009), as
also seen for protein chaperones (Randall and Hardy
1995). Hfq–RNA complexes represent an ensemble of
binding states (from one to six site occupancy). Compet-
itor RNA gains initial access to only one transiently
unoccupied subunit [Fig. 1C], and dissociation of the

resident RNA occurs from a state in which only one
monomer–RNA contact remains. We envision several
binding intermediates with different subsite occupancies
for the two RNAs that are simultaneously present on
Hfq. Stepwise exchange of RNA elements on Hfq sub-
units involves low ΔG barriers, since binding energies of
both transiently bound RNAs per site should be nearly
equivalent. The observed RNA exchange is not external
energy-driven (Supplemental Table S2).

In the model in Figure 1C, RNA exchange on Hfq is
fully reversible, as required for a mechanism that operates
at equilibrium. The high turnover of binding-competent
RNAs in vivo will, however, drive the net displacement
of any given RNA bound to Hfq, such that the population
of bound RNAs changes with the intracellular RNA pool.
Based on our results, many competitor RNAs at 100–200
nM will drive dissociation, and thereby exchange, from
half-lives of >150 min to 1–2 min, a relevant time frame
in vivo. Passive cycling cannot achieve this and is limited
by the first-order dissociation rate constant of RNA–Hfq
(Supplemental Fig. S5).

RNA concentration-driven cycling is most effective
when the total Hfq concentration is lower than the total
concentration of Hfq-binding sites on RNAs. The sum of
all Hfq binders in vivo must be a significant fraction of
the entire RNA pool; deep sequencing identified >800
Hfq coimmunoprecipitation-enriched mRNAs [Sittka
et al. 2008], and many if not most sRNAs bind Hfq
[Wassarman et al. 2001; Zhang et al. 2003; Sittka et al.
2008]. Some individual sRNAs reach micromolar con-
centrations [Altuvia et al. 1997]. Hfq also binds 16S rRNA
[de Haseth and Uhlenbeck 1980b] and poly-A tails on
mRNAs [Arluison et al. 2004], and many RNAs in E. coli
carry “Hfq aptamer motifs” [Lorenz et al. 2010]. Thus, the
many binding-competent RNAs, even at moderate copy
numbers per cell, are expected to saturate Hfq.

What are the consequences of passive or active cycling?
Pulse induction of the Hfq-dependent sRNA RybB down-
regulates many targets. Already, at 1 min, their mRNA
levels were <50% [Papenfort et al. 2006]. If we assume an
intracellular Hfq concentration of 1 μM and a generation
time of 50 min in the passive model, dissociation can
release only a negligible fraction of Hfq from bound RNAs,
and de novo synthesis of free Hfq will add the equivalent
of 20 nM within 1 min. Thus, the induced sRNA would
encounter little free Hfq. Furthermore, the sRNA needs to
compete for free Hfq against a great excess of other newly
synthesized RNAs and against free unbound RNAs. Thus,
obtaining a significant fraction of Hfq–RybB complex,
sufficient for multitarget inhibition in the time frame
observed, is not plausible. Note that this conclusion is
also valid at higher Hfq concentrations. In active cycling,
induced RybB exchanges with bound RNAs on the entire
Hfq pool and “equilibrates” in the minute range to permit
a significant fraction of RybB to acquire Hfq.

The above scenarios are necessarily oversimplified.
Association and dissociation rates of RNA–Hfq pairs vary.
Some RNAs have strong distal or proximal face prefer-
ences, which may result in simultaneous binding of two
RNAs, rather than competition [de Haseth and Uhlenbeck
1980a, Mikulecky et al. 2004; Sun and Wartell 2006;
Rajkowitsch and Schroeder 2007; Vecerek et al. 2008; Link
et al. 2009]. In the model [Fig. 1C], we assume that each
exchange occurs on one or the other face at a time and
requires the presence of several identical binding sites.
Hfq binds stably to, and increases the association rate of, Mic–omp mRNA complexes

The three Mic RNAs require Hfq for regulation of their targets. All matching combinations of Mic and omp RNAs were allowed to form base-paired complexes in the absence or presence of increasing concentrations of Hfq6, followed by gel-shift analysis. For all RNA pairs, ternary complexes were stable on gels and were distinguishable from RNA*–RNA duplexes by slower mobility (Fig. 3A; Supplemental Fig. S6A,B). When ompA-M6 mRNA, inactive in MicA binding and regulation (Udekwu et al. 2005), was used as target, ternary complexes were not formed (data not shown). Thus, Hfq stably interacts with base-paired RNA–RNA complexes.

Hfq enhances sRNA–target RNA binding (Møller et al. 2002, Zhang et al. 2002; Afonyushkin et al. 2005; Kawamoto et al. 2006; Soper and Woodson 2008; Holmqvist et al. 2010). We monitored association of the Mic*–omp RNA pairs ±10 nM Hfq6. From the time courses in Figure 3B (and Supplemental Fig. S6C,D), second-order binding rate constants were calculated. In the absence of Hfq, RNA duplex formation was slow (MicA*–ompA RNA: 7 × 104 M−1 sec−1), but inclusion of Hfq promoted ternary complex formation within 20 sec, which corresponds to an ∼50-fold increase in sRNA–mRNA association rate (MicA*–ompA RNA: 3.5 × 106 M−1 sec−1; less for Mic–ompF) (Supplemental Fig. S6D), in agreement with results reported (Soper and Woodson 2008).

Competitor RNAs readily displace Hfq from RNA duplexes

We asked whether Hfq can be actively displaced from ternary complexes as from binary complexes. Preformed Mic*–omp or Mic*–omp–Hfq6 complexes were competed with increasing concentrations of Mic RNAs, followed by gel-shift analysis (Fig. 3C; Supplemental Fig. S6E,F). These results show that (1) Mic*–omp complexes formed in the absence of Hfq are stable; even a great molar excess of unlabeled Mic RNA only slightly increases the level of free Mic*, (2) Hfq increases RNA–RNA association rate rather than prevents dissociation, and (3) increasing concentrations of unlabeled Mic RNAs scavenge Hfq from the ternary complexes, leaving the Mic*–omp RNA complexes mostly intact.

RNA-driven displacement on ternary complexes was quantitatively assayed by filter binding. Figure 2, E and F, and Table 1 show that ternary complexes were very stable and that dissociation was competitor concentration-dependent (somewhat less effective than on binary complexes). Apparent k4 values ranged from ∼0.4 × 10−4 sec−1 (no competitor) to ∼2.3 × 10−4 sec−1 [at 7 μM MicA], representing a ∼50-fold increase in dissociation rate and a decrease in half-life from ∼290 to ∼5 min. Thus, active exchange of ternary complexes contributes to overall cycling.

Cycling occurs in many protein–protein and RNA–protein complexes. However, the realization that cycling occurs does not answer the question of how it works and whether the properties of the mechanism can account for the biological constraints in vivo. The mechanism of cycling of RNAs on Hfq described here is fundamentally different from conventional (passive) cycling. Even though observations in line with ours have been published (e.g., Leake and Woodson 2004, Afonyushkin et al. 2005), previous work did not consider that rapid cycling is promoted by RNA concentration-driven displacement.

It will be a challenge to address the validity of the active cycling model in vivo. Overexpression of an sRNA changes the repertoire of Hfq-associated RNAs (Papenfort et al. 2009), in agreement with our model. We plan to obtain more quantitative data on Hfq-bound/free RNAs under sRNA induction conditions and address cycling kinetics in vivo. Finally, it is tempting to speculate that cycling of RNA on eukaryotic Lsm proteins follows the model proposed in this study.

Materials and methods

Chemicals, reagents, and oligodeoxyribonucleotides

Chemicals and reagents were purchased from Sigma-Aldrich or GE Healthcare. Oligodeoxyribonucleotides (Supplemental Table S3), poly-A, and poly-U RNAs were from Sigma-Genosys.

Purification of Hfq

Purification of His6-tagged Hfq from E. coli strain BL21(DE3)pLys with plasmid pTE607 (a kind gift of Thomas Elliott, West Virginia Science Center) is described in detail in the Supplemental Material. The final Hfq fractions were stored in buffer A (50 mM Tris-HCl at pH 7.5, 1 mM EDTA, 5% glycerol + 50 mM NH4Cl and 0.1% Triton X-100) at 4°C. Hfq concentrations are given as hexamers throughout this paper.
RNA synthesis and labeling

The ompA, ompC, and ompF RNAs comprise the 5′-most 172, 216, and 214 nucleotides of the three mRNAs, respectively. All RNAs were transcribed from PCR-generated DNA templates carrying a T7 promoter. For primers, see Supplemental Table S3. In vitro RNA synthesis was done by T7 RNA polymerase [Ambion], and purification of the RNAs followed the protocols described [Darfeuille et al. 2007]. When applicable, RNAs were dephosphorylated and 5′-end-labeled with γ-[32P]ATP or γ-[33P]ATP (PerkinElmer).

Gel mobility shift assays

Gel mobility shift assays were carried out with in vitro transcribed RNAs and preparations of Hfq in TMN buffer (20 mM Tris-acetate at pH 7.6, 100 mM Na-acetate, 5 mM Mg2-acetate) for 5 min at 37°C. For details on the specific conditions used, and on gel analyses and calculations, see the Supplemental Material.

Filter binding assays

Reactions conditions and buffers were as described for gel-shift assays, and details of the assay protocol can be found in the Supplemental Material.

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References


Filter binding assays

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