RNase E action at a distance: degradation of target mRNAs mediated by an Hfq-binding small RNA in bacteria

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A major class of bacterial small RNAs (sRNAs), along with RNA-binding protein Hfq and endoribonuclease RNase E, acts on target mRNAs through base-pairing, leading to translational repression and rapid degradation of the mRNAs. In this issue of *Genes & Development*, Prévost and colleagues (pp. 385–396) demonstrate by using the well-characterized sRNA RyhB that RNase E cleavage at sites distal from the pairing region triggers degradation of target mRNAs. The study has provided an important insight into the initial events of sRNA-induced degradation of target mRNAs.

Regulation of gene expression by RNA molecules was known in prokaryotes far before the discovery of microRNAs (miRNAs) and siRNAs in eukaryotes. Since the serendipitous discovery of MicF, which is involved in the down-regulation of the expression of *ompF*, a gene that encodes a major outer membrane porin of *Escherichia coli* (Mizuno et al. 1984), a number of regulatory RNAs have been shown to be widely involved in the regulation of gene expression in bacteria. Among them, the Hfq-binding small RNAs (sRNAs), which are functionally analogous to eukaryotic miRNAs, have been most extensively studied (for reviews, see Waters and Storz 2009, Gottesman and Storz 2010). These sRNAs, which are generally transcribed under specific physiological conditions, act by pairing to target mRNAs and regulating their translation and stability, in many cases negatively, with the help of the RNA-binding protein Hfq. Hfq facilitates the pairing and stabilization of sRNAs, with the major outcome being translational repression and rapid degradation of the target mRNAs.

The mechanism by which sRNAs lead to translational repression is rather simple and fairly well understood. sRNAs mostly bind to the translation initiation region of target mRNAs and occlude the ribosome-binding site (RBS) (for reviews, see Waters and Storz 2009, Gottesman and Storz 2010). However, the mechanism of sRNA-mediated degradation of target mRNAs is apparently complicated and is only partially understood. Based on the studies on a few sRNAs, several notable features have been uncovered (for review, see Aiba 2007). First, a major endoribonuclease, RNase E, is responsible for the sRNA-mediated destabilization of target mRNAs (Massé et al. 2003; Vanderpool and Gottesman 2004; Kawamoto et al. 2005). Second, sRNAs paired with target mRNAs are degraded together in an RNase E-dependent manner (Massé et al. 2003). Third, Hfq, along with Hfq-binding sRNAs, binds to RNase E, resulting in the formation of specific ribonucleoprotein complexes that allow for the tethering of RNase E near the base-pairing region of target mRNAs (Morita et al. 2005). One of the unexplored questions is where and how the RNase E-dependent degradation of the mRNA–sRNA hybrid is initiated.

In this issue of *Genes & Development*, Prévost et al. (2011) use the well-characterized sRNA RyhB to show that RNase E cleavage at sites distal from the pairing region triggers RyhB-mediated rapid degradation of target mRNAs. When RyhB pairs with the target mRNAs, ribosome loading onto the translation initiation region is blocked, and ribosomes already engaged in translation are eventually cleared from the downstream cleavage sites. This allows RNase E tethered at the pairing region to attack the cleavage sites, leading to the subsequent rapid degradation of target mRNAs. The work has thus provided an important insight into the initial events of RyhB-induced degradation of target mRNAs. However, it remains to be seen whether RNase E action at a distance is a general mechanism for the mRNA degradation mediated by Hfq-binding sRNAs.

In this perspective, we first summarize the general properties of Hfq-binding sRNAs, the roles of Hfq in sRNA function, and the two major regulatory outcomes of base-pairing [translational repression and RNase E-dependent degradation of target mRNAs]. We also discuss how these regulatory outcomes are mediated by a ribonucleoprotein complex consisting of RNase E, Hfq, and sRNAs. Then, we discuss the work in the Prévost et al. (2011) study and its implications for the role of Hfq-binding sRNAs in mRNA degradation, focusing on how the translational repression...
mRNAs simultaneously, thereby increasing the local concentrations of two RNAs (Storz et al. 2004). Besides facilitating base-pairing, Hfq also contributes to sRNA function by stabilizing sRNAs (for review, see Gottesman 2004).

**Rnase E-dependent coupled degradation of target mRNAs and sRNAs**

Another major outcome of sRNA–mRNA pairing is the rapid degradation of target mRNAs. It is generally accepted that mRNA degradation in bacteria is initiated by endoribonucleolytic cleavage, resulting in mRNA fragments that are then degraded by 3′–5′ exonucleases (for reviews, see Belasco 2010; Carpousis 2007). Rnase E is a major endoribonuclease that is responsible for degradation of many RNAs in *E. coli* (for review, see Carpousis 2007). It forms the RNA degradosome, associating with polynucleotide phosphorylase [PNPase], RNA helicase B [RhlB], and the glycolytic enzyme enolase as the principle partners (for review, see Carpousis 2007). It is likely that both Rnase E and PNPase within the degradosome act in a concerted fashion during the degradation of RNAs, while RhlB unwinds the secondary structure of target RNAs to permit access of Rnase E and/or PNPase. The Rnase E polypeptide is composed of an N-terminal catalytic region and a C-terminal scaffold region responsible for the interaction with partner proteins (Carpousis 2007; Worrall et al. 2008).

sRNA-induced mRNA degradation was first observed in the study of RyhB and its target mRNAs. RyhB down-regulates a number of genes that encode for iron-binding proteins, such as *sodB*, which encodes for the superoxide dismutase enzyme in response to iron limitation (Massé and Gottesman 2002). RyhB causes a rapid degradation of target mRNAs in an Rnase E-dependent manner (Massé et al. 2003). The Rnase E-dependent rapid degradation of target mRNAs is probably a general feature of sRNA function (Morita et al. 2004, 2005). In addition, sRNAs were shown to be destabilized, depending on pairwise with target mRNAs, also in an Rnase E-dependent manner, suggesting that Rnase E somehow specifically attacks the sRNA–mRNA duplex. (Massé et al. 2003). Furthermore, it was shown that the C-terminal scaffold region of Rnase E is required for the coupled degradation of sRNAs and their target mRNAs (Massé et al. 2003; Morita et al. 2004).

**Ribonucleoprotein complexes consisting of sRNAs, Hfq, and Rnase E**

A key discovery for understanding the mechanism of sRNA-mediated degradation of target mRNAs was that Hfq associates with Rnase E through its scaffold region, and at least two sRNAs—SgrS and RyhB—associate with Rnase E through Hfq (Morita et al. 2005). SgrS is another Hfq-binding sRNA; it is induced in response to the accumulation of glucose-phosphate, resulting in the Rnase E-dependent destabilization of *ptsG* mRNA, which encodes for the glucose transporter IICB 

and RNase E-dependent degradation of target mRNAs mediated by sRNAs are interrelated.

**Hfq-binding sRNAs**

Hfq-binding sRNAs act through imperfect base-pairing with a single or multiple target mRNAs, regulating their translation and stability (for reviews, see Waters and Storz 2009; Gottesman and Storz 2010). In contrast to miRNAs that bind to the 3′ untranslated region (UTR) of target mRNAs, bacterial sRNAs bind to the 5′ UTR of target mRNAs (Gottesman 2005). The translation initiation region in the target mRNAs is most often used for base-pairing, and the pairing therefore occludes the RBS and results in translational repression—although, in a few cases, the pairing occurs upstream of the RBS (for review, see Waters and Storz 2009). The sRNAs can also activate expression of the target mRNAs in some cases, by disrupting an inhibitory secondary structure that sequesters the RBS (for reviews, see Waters and Storz 2009; Gottesman and Storz 2010). Although the potential base-pairing between sRNAs and target mRNAs ranges from 10 to 30 nucleotides (nt), more limited core-pairing appears to be sufficient for regulation (Waters and Storz 2009). For example, a 14-nt sequence within the 31-nt potential pairing region of SgrS is sufficient to inhibit the target *ptsG* translation (Maki et al. 2010), and only four single mutations in SgrS significantly affect the down-regulation of *ptsG* mRNA (Kawamoto et al. 2006).

The RNA-binding protein Hfq, which was originally identified as a host factor required for the in vitro replication of the RNA phage Qβ in *E. coli*, is extensively involved in the regulation of RNA metabolism (for review, see Valentin-Hansen et al. 2004). The involvement of Hfq in sRNA functions was first recognized during the study on translational repression of *rpoS* mRNA (encoding the alternative σ factor RpoS) by OxyS RNA that is induced in response to oxidative stress in *E. coli* (Zhong et al. 1998). Subsequently, Hfq was shown to bind to and be necessary for the regulatory activity of a number of sRNAs (for reviews, see Gottesman 2004; Valentin-Hansen et al. 2004). Hfq resembles the Sm and Sm-like proteins involved in splicing and mRNA degradation in eukaryotic cells in that it forms a donut-shaped structure consisting of six identical 11-kDa subunits (Moller et al. 2002; Zhang et al. 2002). Hfq has been demonstrated to stimulate sRNA–mRNA base-pairing in vitro in several cases, such as OxyS-filA (Zhang et al. 2002), Spot42-galK (Moller et al. 2002), MicA-ompA (Rasmussen et al. 2005), and SgrS-ptsG (Kawamoto et al. 2006). A recent study has demonstrated that, in vitro, Hfq is no longer required for translational inhibition once base-pairing is achieved (Maki et al. 2008). However, all Hfq-binding sRNAs examined to date require Hfq for their regulatory functions in vivo. It is highly likely that the Hfq-mediated rapid association of sRNAs with target mRNAs is necessary to efficiently block ribosome loading to the RBS. Hfq promotes base-pairing by either changing the RNA structures, acting as an RNA chaperone, thereby allowing accessibility of two complementary RNAs, or binding to a given sRNA and its target mRNA.
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Morita et al. 2003; Vanderpool and Gottesman 2004]. RyhB- and SgrS-mediated destabilization of target mRNAs is dependent on Hfq and the C-terminal scaffold region of RNase E [Kawamoto et al. 2005; Morita et al. 2004]. The physical association of sRNAs/Hfq with RNase E explains how the functional cooperation of sRNAs, Hfq, and RNase E is achieved. A recent study has established that the subregion of RNase E overlapping the RhkB-binding site is responsible for the interaction with Hfq, leading to the rapid degradation of target mRNAs mediated by sRNAs [Ikeda et al. 2011].

RNase E action at a distance

Prévost et al. [2011] addressed where and how the initial cleavage by RNase E occurs in sRNA–mRNA duplexes. By analyzing a series of sodB-lacZ transcriptional fusions in which various lengths of the 5′ portion of sodB transcript were fused to the lacZ, they defined a site, between +400 and +430, far downstream from the pairing region, that is required for the RyhB-mediated degradation of the target sodB mRNA. This mRNA degradation was shown to be dependent on both RNase E and Hfq, suggesting that RNase E tethered near the sodB RBS cleaves the region between +400 and +430. An AU-rich sequence within the 400–430 region of sodB is cleaved by RNase E in vitro, and in vitro cleavage no longer occurred when the AU-rich sequence was mutated. More importantly, the mutation eliminated the RNase E-dependent rapid degradation of a sodB-lacZ fusion mRNA in vivo. Furthermore, a short sodB-lacZ construct was RNase E-resistant but became sensitive to RNase E cleavage when the 30-nt section was added. This led Prévost et al. [2011] to conclude that the AU-rich sequence within the 400–430 region of sodB is sufficient to cause the RNase E-dependent rapid degradation mediated by RyhB/Hfq. They propose that RyhB induces an initial mRNA cleavage by RNase E at this distal site far downstream from the pairing region to destroy the target mRNA. Strictly speaking, it remains to be seen whether the cleavage at the distal AU-rich sequence by RNase E tethered at the base-pairing region is really an initial event for the rapid degradation of sodB mRNA mediated by RyhB, although it is surely the rate-limiting step.

mRNA cleavage at a distal site was observed in other RyhB target mRNAs, although the distance between the pairing site and the cleavage site varied depending on the target mRNA. This suggests that the mRNA cleavage at a distal site by RNase E is a general mechanism associated with sRNA-mediated rapid degradation of target mRNAs. Action at a distance is a widely used mechanism in the regulation of DNA and RNA functions. For example, it is well known that transcription of genes is regulated very often by transcription factors that bind to sites on the DNA at a considerable distance in both bacteria and eukaryotes [for review, see Ptashne 1986]. Translation of mRNAs is stimulated by proteins bound at its 3′ polyA tail and inhibited by protein components within RISC [RNA-induced silencing complex] tethered at the 3′ UTR through miRNAs in eukaryotes [for review, see Carthew and Sontheimer 2009]. The major impact of the study by Prévost et al. [2011] is that it demonstrated that endoribonucleases can also act at a distance.

Interplay between mRNA cleavage and translational repression

How are the translational repression and rapid degradation of target mRNAs mediated by sRNAs interrelated? Theoretically, these two events could affect each other. Namely, translational inhibition can lead to destabilization of an mRNA by exposing it to ribonucleases, while mRNA degradation should certainly cause loss of translation by reducing mRNA concentration. A previous study on SgrS and RyhB demonstrated several key features of the interrelationship between translational repression and sRNA-mediated mRNA degradation [Morita et al. 2006]. First, translational repression occurs without the RNase E-dependent degradation of mRNAs, and therefore is primarily responsible for the down-regulation of target mRNAs by sRNAs. Thus, the major role of the RNase E-dependent rapid degradation of mRNAs appears to be to make gene silencing irreversible and eliminate translationally inactive mRNAs, although mRNA degradation seems to contribute to the down-regulation of target mRNAs without blocking translation in one case [Pfeiffer et al. 2009]. In addition, translational repression itself—by either Hfq/sRNAs or a drug—is not sufficient for degradation, and recruitment of RNase E to the base-pairing region by Hfq/sRNAs is required for the RNase E-dependent rapid degradation of target mRNAs. The study by Prévost et al. [2011] confirmed and extended these previous conclusions. They showed, by introducing a stop codon far upstream of the cleavage site, that blocking translation alone is not sufficient to induce rapid degradation. In other words, RyhB-mediated rapid mRNA degradation is an active outcome by a concerted action of RNase E, Hfq, and RyhB, rather than a passive secondary effect of blocking translation.

The work by Prévost et al. [2011] has provided an additional insight into the relationship between the translational repression and mRNA degradation mediated by sRNAs. This derives from the analysis of a series of SodB-LacZ translational fusions. In contrast to the transcriptional fusions discussed above, all translational fusions were sensitive to RyhB-mediated degradation, even when the fusions did not harbor the identified RNase E site between +400 and +430. This could be explained by assuming the presence of RNase E-sensitive sites within the LacZ ORF. If so, why are the additional cleavage sites resistant to RNase E in the sodB-lacZ transcriptional fusions? Although Prévost et al. [2011] did not extensively address this point in their study, the results provide further insight into the mechanism of RyhB-mediated mRNA cleavage at a distance. It is highly likely that the RyhB-mediated cleavage of target mRNAs at a distance occurs only when the translating ribosomes, which are expected to protect mRNAs from RNase attack, are cleared from the RNase E site. It is important to note that ribosomes are eventually cleared from the lacZ portion in SodB-LacZ translational fusions when RyhB inhibits the
new ribosome loading to the sodB RBS. On the other hand, the inhibition of sodB translation by RyhB does not affect the translation of the lacZ portion in transcriptional fusions because the lacZ translation occurs independently of sodB translation. Thus, the RNase E sites in the lacZ portion of the sodB-lacZ transcriptional fusions are protected by translating ribosomes. This is probably why the cleavage site within lacZ is resistant to RNase E. In other words, the RyhB-mediated translational inhibition is necessary for RyhB-mediated mRNA cleavage. We emphasize that RNase E recruited to the target mRNA by Hfq/sRNAs—but not free RNase E—can efficiently attack the cleavage sites. Altogether, the data presented in the study by Prévost et al. (2011) lead to the model shown in Figure 1. When RyhB is induced, it associates with Hfq and therefore RNase E, resulting in a specific ribonucleoprotein complex. The ribonucleoprotein complex acts on the target mRNAs: Hfq facilitates the pairing and RNase E is tethered at the pairing region. The pairing inhibits new ribosome loading, while translating ribosomes protect the downstream RNase E site from the attack by RNase E. Whether RNase E tethered at the pairing region, but not freestanding RNase E/degradosome, efficiently attacks the RNase E site after the translating ribosomes have passed through. Whether RNase E tethered at the base-pairing region directly attacks the paired RyhB remains to be studied.

Figure 1. Model for RNase E action at a distance. (A) Hfq binds to the C-terminal scaffold region of RNase E. When RyhB is induced, it associates with Hfq and therefore RNase E, resulting in a specific ribonucleoprotein complex. (B) The ribonucleoprotein complex acts on the target mRNAs: Hfq facilitates the pairing and RNase E is tethered at the pairing region. The pairing inhibits new ribosome loading, while translating ribosomes protect the downstream RNase E site (shown in red line) from the attack by RNase E. (C) RNase E tethered at the pairing region, but not freestanding RNase E/degradosome, efficiently attacks the RNase E site after the translating ribosomes have passed through. Whether RNase E tethered at the base-pairing region directly attacks the paired RyhB remains to be studied.

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