RNA enables the material interpretation of genetic information through time and space. The creation, destruction and activity of RNA must be well controlled and tightly synchronized with numerous cellular processes. We discuss here the pathways and mechanism of bacterial RNA turnover, and describe how RNA itself modulates these processes as part of decision-making networks. The central roles of RNA decay and other aspects of RNA metabolism in cellular control are also suggested by their vulnerability to sabotage by phages; nonetheless, RNA can be used in defense against phage infection, and these processes are described here. Salient aspects of RNA turnover are drawn together to suggest how it could affect complex effects such as phenotypic diversity in populations and responses that persist for multiple generations.

Introduction

While transcripts are transient entities in the lifetime of a cell, their limited lifespans serve as a means of manipulating the strength and duration of gene expression. The adjustment of transcript turnover sustains homeostasis and is a key aspect of the rapid adaptation to environmental changes. In *Escherichia coli*, protein-encoding RNAs have on average overall chemical half-lives of 7–8 min, but individual transcripts each have their own characteristic lifetime that can be modulated according to growth stage and stress response. The modulation of transcript lifetime is affected by protective binding partners and the accessibility to the machinery of RNA decay that include ribonucleases, their accessory proteins and helper RNAs that catalyze RNA turnover.

The general scheme for degradation of RNA and the key participating enzymes for the Gram-negative *Escherichia coli* and the Gram-positive *Bacillus subtilis* are shown in the schematic in Figure 1. These bacterial species are thought to have diverged at least 1 billion years ago, and each is considered representative in genomic character for many of the currently known bacteria. In both *E. coli* and *B. subtilis*, the principal degradation pathways begin with endoribonuclease cleavage that destines the transcript to a fate of complete destruction. The cleavage(s) made by an endoribonuclease can be followed by degradation catalyzed by exoribonucleases.

Despite the similarity in the overall pathways of *E. coli* and *B. subtilis*, surprisingly few of the participating enzymes are common to both species. It seems remarkable that entirely different enzymes, with unrelated structures or evolutionary origins, have evolved with analogous function in the two organisms. This likely reflects the biological importance of the pathway and the power of convergent evolution to meet the cellular demands for RNA metabolism. In *E. coli* and many other proteobacteria, one of the key enzymes in the decay pathway is the hydrolytic endoribonuclease RNase E, which is highlighted in the *E. coli* decay pathway shown in the left panel of Figure 1. In *B. subtilis*, RNase Y serves a functionally analogous role to *E. coli* RNase E, as shown in the right panel of Figure 1.

Although RNase E cleaves RNA internally, it often interacts with the 5' end of its many substrates and prefers those harboring a monophosphate group. In bacteria, primary transcripts are transcribed with a 5' terminal triphosphate group, which can confer protection from RNase E attack. However, a pyrophos- phophydroxylase (RppH) can remove the pyrophosphate from the 5' end of the RNA, leaving a monophosphorylated RNA that can be rapidly degraded by RNase E. The *B. subtilis* ribonuclease RNase J also has a preference for substrates with a 5' mono- phosphate. A homolog of the RppH pyrophosphohydrolase in *B. subtilis* appears to play an analogous role to decap the triphosphate group of transcripts. It seems likely that 5' end recognition is also important in the decay pathway in *B. subtilis* although the structural basis for the ribonuclease activation has an entirely different stereochemical origin; this parallel seems yet another remarkable case of evolutionary convergence.

Following the initiating endoribonuclease cleavage(s), exoribonuclease activities in *E. coli* are provided by 3' to 5' exoribonucleases such as the phosphorolytic polynucleotide phosphorylase (PnPase), or the hydrolytic enzymes RNase R or RNase II. Exoribonuclease cleavages in *B. subtilis* are mediated by the hydrolytic RNase J in the 5' to 3' direction, and by PNPase, with the opposite polarity. These processes are assisted by other enzymes, such as poly(A) polymerase (PAP), which adds 3'-terminal poly-adenine tails that become sites for processive degradation. Poly-adenine tails may also help initiate degradation, as polyadenylation affects the lifetime of many transcripts in *E. coli* and potentially helps to control levels of tRNA.

Moreover, in *B. subtilis*, poly(A) and heteropolymeric tails seem
to mark the RNA for degradation as well, although the enzyme responsible for adding the tails has not yet been identified.27 In considering the degradation pathways of the two representative bacteria, it is interesting to note that the archaea harbor some bacterial-like enzymes such as RNase J and some eukaryotic-like RNA metabolic assemblies, such as the exosome (described further below). The archaeal pathways will not be described here, except to draw occasional parallels between the bacterial components.

Several of the enzymes depicted in Figure 1 play additional functional roles, as they are also involved in processing precursors of structured RNA. For instance, *E. coli* RNase E also takes part in RNA maturation by cleaving precursors of structured RNAs.8 This activity reveals a more controlled operation mode for the ribonuclease, in which the cleavage products are not marked for destruction but rather processed further to form a functional RNA molecule. How this discrimination is achieved is an intriguing puzzle. The processing activity of RNase E is a dominating aspect of its in vivo activity, and it has been suggested that the enzyme may catalyze as many processing as degradation cleavages during exponential growth stage.8 Processing activities are also likely to be a dominating mode for RNase Y and RNase J in *B. subtilis* and other species.

A multi-enzyme assembly for RNA degradation. In many bacterial species, a multi-enzyme assembly known as the RNA degradosome acts as a key machinery of RNA decay and processing.28 The components of the degradosome cooperate to degrade RNA transcripts and to process precursors of structured RNAs. In *E. coli* and several other bacteria, the core of the complex is formed by RNase E, which was mentioned above. The protein composition of the degradosome assemblies differs between evolutionarily divergent bacterial species, and some degradosome assemblies include enzymes such as exoribonucleases and metabolic enzymes, but they often include a DEAD-box helicase that directly associates to an endoribonuclease.29

The exoribonuclease PNPase, mentioned above as an important contributor to prokaryotic RNA degradation is a canonical component of the *E. coli* RNA degradosome (Fig. 2). There are interesting structural and functional parallels between PNPase with the archaeal and eukaryotic exosomes. All share a common evolutionary origin with the phosphorolytic enzyme RNase PH and have chamber-like quaternary architecture that forms a central channel containing the active site. In both PNPase and the exosome, the entrance to the central channel narrows to form an aperture wide enough only for single-stranded RNA substrate to enter, suggesting a necessity for structured RNA substrates to be unwound prior to processing by the degradative machinery.

The RNA helicase (RhlB) as part of the *E. coli* RNA degradosome, aids the processing of RNA substrates by the assembly.30–32 The unwinding or remodelling activities of RhlB are required to improve the efficiency of mRNA degradation, since the ribonuclease activities of RNase E and PNPase are specific for single-stranded RNA (Fig. 2). The ATPase activity of RhlB is required to facilitate the degradation by PNPase of structured RNA transcripts with a repetitive element that forms stable stem-loop structures.30–33 The helicase binds to a site in RNase E that is flanked by RNA-binding domains, and these perhaps help to capture substrates for the helicase, as does also an arginine-rich tail on the C terminus of RhlB.29

Riboregulation: RNA turnover mediated by small regulatory RNAs. Small regulatory RNAs (sRNAs) are an important class of trans-acting regulators of post-transcriptional gene regulation that can modulate transcript lifetimes, rapidly and with specificity (Fig. 3). They appear to be ubiquitous throughout the bacterial lineages, and mediate numerous processes, including membrane stress response, carbon regulation, virulence effects, biofilm formation and countless other complex behaviors.34,35 Many sRNAs act by impeding translation and increasing likelihood for encounters with a ribonuclease with subsequent turnover, but some also have entirely the opposite effect, and act by exposing the ribosome-binding site to favor translation initiation;36 this presumably protects the RNA against ribonuclease activity.

sRNAs can target multiple transcripts, and some transcripts are the targets for several different sRNAs.34 They emulate elements of complex logic circuits, and help to build post-transcriptional networks that are as rich in complexity as the well-characterized transcriptional regulatory networks.37 In many bacteria, a key facilitator of sRNA function in these regulatory networks is the RNA chaperone protein Hfq.38 The protein belongs to the extensive Sm-family, characterized by a conserved fold and propensity to assemble into a ring-like oligomeric architecture (Fig. 4). Hfq can interact with sRNA and target transcripts through three different binding surfaces on this ring: proximal, distal and circumferential. These interactions help to mediate pairing of sRNAs to targets. The limited cellular quantity of the protein means that RNAs must compete for binding Hfq in sRNA-mediated riboregulation, with implications for inter-connecting different regulatory networks.39

In *E. coli*, RNase E appears to be the dominant enzyme responsible for sRNA-mediated response and for degradation of the sRNAs.40 Another enzyme that was shown to influence sRNAs that are not associated with Hfq is PNPase, and this exoribonuclease is responsible for their degradation in stationary phase.41 However, PNPase has been identified in a genetic screen to be linked to Hfq-mediated sRNA activity in *E. coli* and, surprisingly, it affects the stability of some sRNAs in vivo by protecting them from degradation by other ribonucleases.42 It seems paradoxical that a ribonuclease might confer a protective effect on an sRNA in vivo, but there is precedent for example in RNase II playing a protective role for other RNAs by removing 3′-poly(A) tails.43 As PNPase is a processive exonuclease, it might be expected to drive the degradation of the substrate once it has loaded, but perhaps in the case of structured sRNAs it might trap them into protective complexes. This mechanism awaits experimental confirmation.

sRNAs are also present in *B. subtilis*, however the study of the networks involving these regulatory molecules in this organism are still at an early stage, and the roles of sRNAs are not yet as well defined as in *E. coli*. However, it has already been shown in *B. subtilis* that a small RNA and a variety of potential RNA chaperones are involved in iron sensing.44 The *B. subtilis* RNase Y may also be involved in the turnover of cis-acting regulatory RNA,
The cleavage of some RNA substrates in *E. coli* is not significantly influenced by the status of their 5’ end, these are recognized and cleaved probably by RNase E through an internal entry pathway, which does not require enzyme activation by the monophosphate bound in the 5’ binding pocket. The processes are assisted by the C-terminal half of RNase E, which is a mostly unstructured region that contains two RNA-binding domains important for the substrate recognition and alignment on the enzyme.

The 3’ end of a transcript is also important for affecting its lifetime. In *E. coli*, the decay of many mRNAs is affected by polyadenylation, and the half-lives of certain mRNAs are found to be boosted in the absence of polyadenylation. The poly(A) tails are added by poly(A) polymerase or another enzyme with such activity (yellow ellipse) in *B. subtilis*, contributing to the processivity of RNA turnover in these bacteria. The stem loop structure on the 3’ end of the transcripts at the top are rho-independent transcription terminators.

**Figure 1.** A billion years of bacterial RNA degradation. The general scheme of RNA degradation pathways in *Escherichia coli* (left panel) and *Bacillus subtilis* (right panel). The key enzymes involved in the different steps are listed in parentheses, and represented in brown (endonucleases) or gray (exoribonucleases). RNase J has both endo- and exo-activities, and it is depicted with the concentric gray and brown circles. “Decapping” of the 5’ terminal triphosphate to leave a 5’ monophosphate may convert some transcripts into preferred substrates for RNase E in *E. coli* and RNase J in *B. subtilis*. At the 3’ end of transcripts, poly(A) tails are added by poly(A) polymerase or another enzyme with such activity (yellow ellipse) in *B. subtilis*, contributing to the processivity of RNA turnover in these bacteria. The stem loop structure on the 3’ end of the transcripts at the top are rho-independent transcription terminators.
be involved in some cases, as the polyadenylation of the antisense RNA I in vivo may recruit PNPase and lead to decay of that RNA. Polyadenylation and linked degradation in bacteria has an interesting parallel with surveillance in eukaryotes, where the helicase-bearing TRAMP complex recognizes aberrant transcripts, certain non-coding RNA and snoRNAs, tagging them with poly(A) tails and directing them to the exosome.\textsuperscript{67} Perhaps a somewhat similar process occurs in bacteria, with the poly(A) polymerase marking RNAs with the poly(A) tails, the RNA unwinding role being fulfilled by the DEAD box helicase RhlB and cleavage is associated with RNase E together with PNPase in the RNA degradosome assembly.

**RNA turnover and energy economy.** Under starvation conditions, *E. coli* enters the stationary phase, where growth is slow, metabolism is less active overall and there is a general downregulation of both transcription and translation.\textsuperscript{68} Counterintuitively, translation of some metabolic enzymes must increase as cells enter stationary phase: this enables the bacterium to use a broader range of carbon and nitrogen sources. The starvation phenomenon of persistence, in which sub-populations in an isogenic population are more resistant to the action of antibiotic.\textsuperscript{83} This phenomenon is termed confers immunity to a population of bacteria by triggering death of infected cells. It is not presently known what triggers the switch.\textsuperscript{71} This switch is affected by acetylation of a specific lysine in RNase R. Metabolic enzymes in *Salmonella* are also regulated by lysine acetylation,\textsuperscript{79} suggesting that this modification could be the downstream effect of a signaling pathway. We speculate that there is a likely link of cell division and ribonuclease pathway, sensitive to the metabolic state of the cell.\textsuperscript{74} We speculate that there is a likely link of cell division and ribonuclease pathway, sensitive to the metabolic state of the cell.\textsuperscript{74}

**Figure 2.** The *Escherichia coli* RNA degradosome, and facilitated substrate delivery to the exoribonuclease component, polynucleotide phosphorylase (PNPase, blue). A portion of the PNPase ring has been removed to illustrate the central chamber that harbors the active site. An ATP-dependent DEAD box helicase RhlB (green), another degradosome component, is unwinding the RNA stem-loop (red), providing a single-stranded RNA for PNPase. Both enzymes are associated with RNase E (purple). For clarity, other degradosome components, such as enolase, were omitted.

RNA turnover can proceed either through hydrolysis or phosphorylation. The latter is more useful for shunting into DNA precursors for cell division, and would be expected to occur close to the onset of DNA replication. It is interesting to note that levels of the hydrolytic exoribonuclease RNase R decreases in stationary phase,\textsuperscript{70,71} but they increase in stress conditions.\textsuperscript{72} This switch is affected by acetylation of a specific lysine in RNase R. Metabolic enzymes in *Salmonella* are also regulated by lysine acetylation,\textsuperscript{79} suggesting that this modification could be the downstream effect of a signaling pathway. We speculate that there is a likely link of cell division and ribonuclease pathway, sensitive to the metabolic state of the cell.\textsuperscript{74} The speculative link is supported by the functionally important role of ribonucleases for controlling certain cell division genes.\textsuperscript{75-77} Perhaps in spore-forming species such as *Streptomyces* and *Bacillus*, this proposed linkage might be important for energy conservation for cells as they enter the vegetative storage stage.

**RNA is manipulated by phage in infection, but also used in host defense.** Given the role of RNA metabolism in the regulation of complex genetic networks, it might seem unsurprising that this pathway can be targeted by phage during infection in numerous ways. For instance, RNase E is directly manipulated through phosphorylation by a kinase encoded by T7 phage, and the modification changes the ribonuclease activity to selectively stabilize phage-encoded transcripts.\textsuperscript{8,78} Recently, bacteriophage T4 polynucleotide kinase has been found to destabilize host mRNAs.\textsuperscript{71}

Small RNAs are used in host defense against phage infection. The Type III toxin-antitoxin system comprises a toxic endoribonuclease and a non-coding RNA that inhibits the enzyme.\textsuperscript{79-82} In the ToxIN system of *Pectobacterium atrosepticum*, the RNA forms a pseudoknot structure from a repetitive element that is processed by the ToxN ribonuclease. Thus, the ToxI antitoxins inhibit their own parent-processing enzymes and consequently protect the cell from a harmful general ribonuclease. This system confers immunity to a population of bacteria by triggering death of infected cells. It is not presently known what triggers toxin activation—whether foreign metabolite, RNA or abnormal state. Other types of toxin-antitoxin systems contribute to the phenomenon of persistence, in which sub-populations in an isogenic population are more resistant to the action of antibiotic.\textsuperscript{83}

Parallels can be drawn between ToxIN proteins and Type I Cas6 ribonucleases of the anti-viral CRISPR/Cas systems, which provide adaptive immunity against phage infection (CRISPR, clustered regularly interspaced short palindromic repeats; Cas, CRISPR-associated proteins). Like the ToxIN protein, the Cas6 enzyme also remains associated with its own catalytic product following cleavage of the repetitive CRISPR transcript substrates into crRNAs.\textsuperscript{84-86} Binding of crRNAs to their Cas6-processing proteins, such as enolase, were omitted.

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transcripts that are poorly bound by the ribosomes.\textsuperscript{90} However, recent evidence shows that certain transcripts in \textit{E. coli} may in fact be transcribed at some distance from the site of translation,\textsuperscript{91} and in these cases there may be other means of protecting the RNA against ribonuclease attack. RNase E and the degradosome are compartmentalized to the cytoplasmic membrane.\textsuperscript{92} This spatial separation from transcription sites has been proposed to provide a time delay for transcripts to be recognized and tagged by sRNAs before they reach the degradosome for rapid destruction.\textsuperscript{8}

Within its generation time, each \textit{E. coli} cell synthesizes an estimated 430,000 transcripts. How do sRNAs find their targets in such a thicket? The number of potential pairwise interactions is astronomical—taking by analogy the potential interactions in the protein interactome, for a sRNA to find a target by exhaustive search would require improbable rates of discriminating encounters.\textsuperscript{93} We suggest that the problem is more acute in RNA compared with protein, because of the propensity of even random RNA to fold into semi-compact states.

Figure 3. Schematic of small RNA (sRNA)-mediated regulation of gene expression. In the top panel, the sRNAs (red) are transcribed in trans to the target gene (green). A cognate seed region that pairs to a recognition site in the target RNA can relieve repression of translation by secondary structure that masks the ribosome-binding site (RBS, or Shine Dalgarno sequence). In this way, sRNAs can activate gene expression (left side). Conversely, they can mask the RBS (ribosome binding site), thereby preventing translation initiation and making the naked RNA vulnerable to ribonuclease attack, as shown in the middle panel. sRNA can also bind within the coding sequence of the target mRNA and the sRNA 5' end can allosterically activate RNase E to cleave target mRNA (right side).
Does this decay arise from stochastic access to the transcripts, or from a defined process that is precise and accurate? The published decay rates suggest that the processes may be multi-exponential, and the deviation of decay rates from single-exponential imply that the process of RNA turnover is not entirely random, and can arise from numerous processes that affect gene expression—for instance, the hypothetical possibility of counting translation rounds for each transcript, or a synchronized delay in the onset of turnover. Another key question concerning transcript decay is the extent of variation in the lifetime for a particular type of transcript. It seems likely that the detailed analysis of the key degradation parameters may be very illuminating for understanding the underlying processes.

**Summary and Perspective**

When the transcriptional inhibitor rifampicin is added to cultures, many transcripts in *E. coli* are found to decay with exponential timecourse. Does this decay arise from stochastic access to the transcripts, or from a defined process that is precise and accurate? The published decay rates suggest that the processes may be multi-exponential, and the deviation of decay rates from single-exponential imply that the process of RNA turnover is not entirely random, and can arise from numerous processes that affect gene expression—for instance, the hypothetical possibility of counting translation rounds for each transcript, or a synchronized delay in the onset of turnover. Another key question concerning transcript decay is the extent of variation in the lifetime for a particular type of transcript. It seems likely that the detailed analysis of the key degradation parameters may be very illuminating for understanding the underlying processes.

RNA metabolism and sRNA-mediated regulation can give rise to complex phenotypes. sRNAs have multiple targets, and some target transcripts have multiple sRNA regulators. This results in networks of sRNAs forming a complement to the networks controlling transcription and mediated by transcription factors.

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**Figure 4.** The structure and proposed RNA-binding modes of the RNA chaperone Hfq. (A) Crystal structures revealing the ring-like architecture of the hexameric Hfq protein and RNA binding to the distal (left and central panels, *E. coli*) and proximal faces (right panel, *Salmonella typhimurium*) of the ring. (B) C-terminal tails emanate from the core, based on small angle X-ray solution scattering. These tails have some weak conservation in families and are likely to be flexible. They may help to capture RNA and accommodate species with complex folds or serve for interaction with other proteins. (C) A proposed binding model for sRNA (red) in which the 3' poly(U) tail is engaged on the proximal side of the Hfq hexamer (purple), and the body of the sRNA is engaged on the circumferential rim of the ring. The mRNA recognition region, here located on the 5' end of the sRNA, is proposed to be able to peel off to mediate cognate base pairing with the target mRNA.
These networks have, for instance, multi-output feedforward loops that can act to reduce leaky expression and maintain repression of target genes with changing nutrient conditions. One striking feature of sRNA-mediated regulation is the sensitivity of the response curve to the ratio of regulatory RNA to targets. As the ratio approaches unitary stoichiometry, sRNA effects are no longer controlled, and how targets are recognized with specificity and speed.

Genomes seem to bear more information than is encoded as binary bits, as the information acts back on itself. That is, the act of generating a transcript affects the subsequent generation of that transcript itself—and this can be achieved either directly or indirectly. This makes the system self-referential and, accordingly, the information is much richer than the sequence pattern itself. The use of sRNAs in riboregulation enriches and expands this capacity for self-information. Genetic information is thus not only in the digital composition, but also in its base-pairing and fold of cognate matches to RNA. Establishing the full information content of a system like the genome remains a challenging but important problem for experiment and theory, and the mechanisms of RNA metabolism and riboregulation are likely to be a key aspect to address this challenge in the future.

Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

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References
1. Grunberg-Manago M. Messenger RNA stability and its role in control of gene expression in bacteria and phages. Annu Rev Genet 1999; 33:193-227; PMID:10690408; http://dx.doi.org/10.1146/annurev.genet.33.1.193.
2. Miller C, Schwab B, Maier K, Schulz D, Dümske S, Zacher B, et al. Dynamic transcriptome analysis measures rates of mRNA synthesis and decay in yeast. Mol Syst Biol 2011; 7:458; PMID:21206491; http://dx.doi.org. 10.1038/msb.2010.112.
3. Mohanty BK, Kukkani SR. Analysis of the function of Escherichia coli polA (polyA) polymerase I in RNA metabolism. Mol Microbiol 1999; 34:1094-108; PMID:10594833; http://dx.doi.org/10.1046/j.1365-2958.1999.01673.x.
4. Selinger DW, Saxena RM, Cheung KJ, Church GM, Rosenow C. Global RNA half-life analysis in Escherichia coli reveals positional patterns of tran- script degradation. Genome Res 2003; 13:216-23; PMID:12566699; http://dx.doi.org/10.1101/ rgr.921663.
5. Kuner F, Ogusawara N, Moser I, Albertini AM, Alloni G, Azevedo V, et al. The complete genome sequence of the gram-positive bacterium Bacillus subtilis. Nature 1997; 390:249-56; PMID:9384377; http://dx.doi.org/10.1038/36786.
6. Carpousis AJ. The RNA degradome of Escherichia coli: an mRNA-degrading machine assembled on RNase E. Annu Rev Microbiol 2007; 61:71-87; PMID:17447862; http://dx.doi.org/10.1146/annurev.micro.61.080706.093440.
7. Carpousis AJ, Luisi BF, McDowall KJ. Endonucleolytic initiation of RNA decay in Escherichia coli. Prog Mol Biol Transl Sci 2009; 85;91-135; PMID:19215771; http://dx.doi.org/10.1016/S0079-6603(08)00083-9.
8. Mackie GA. RNase E: the interface of bacterial RNA processing and decay. Nat Rev Microbiol 2013; 11:45-57; PMID:23241849; http://dx.doi.org/10/nrmic26936.
9. Stead MB, Marshall S, Mohanty BK, Mitra J, Pena Castillo L, Ray D, et al. Analysis of Escherichia coli Rnase E and Rnase III activity in vivo using tiling microarrays. Nucleic Acids Res 2011; 39:3188-203; PMID:21149258; http://dx.doi.org/10.1093/nar/gkq1242.
10. Durand S, Gilet L, Bessières P, Nicolais P Condon C. Three essential ribonucleases-RNase Y, J1, and III-control the abundance of a majority of Bacillus subtilis mRNAs. PLoS Genet 2012; 8:e1002520; PMID:22421379; http://dx.doi.org/10.1371/journal. pgen.1002520.
11. Lehnik-Habrink M, Lewis RJ, Mader U, Stülke J. RNA degradation in Bacillus subtilis: an interplay of essential endo- and exoribonucleases. Mol Microbiol 2012; 84:1005-17; PMID:22568516; http://dx.doi.org/10.1111/j.1365-2958.2012.08072.x.
12. Shahbahian K, Jamalli A, Zig L, Putzer H. RNase Y, a novel endoribonuclease, initiates ribonuclease turn-over in Bacillus subtilis. EMBO J 2009; 28:3523-33; PMID:19779461; http://dx.doi.org/10.1002/embj.2009.283.
13. Yao S, Bechhofer DH. Initiation of decay of Bacillus subtilis rpoP mRNA by endoribonuclease RNA Y. J Bacteriol 2010; 192;3279-86; PMID:20418391; http://dx.doi.org/10.1128/JB.00230-10.
14. Mackie GA. Ribonuclease IV is a 5′-end-depen- dent endonuclease. Nature 1998; 395;720-3; PMID:9790196; http://dx.doi.org/10.1038/27246.
15. Celenkin H, Deana A, Belasco JG. Initiation of RNA decay in Escherichia coli by 5′-phosphatase removal. Mol Cell 2007; 27;79-90; PMID:17612492; http://dx.doi.org/10.1016/j.molcel.2007.05.038.
16. Deana A, Celenkin H, Belasco JG. The bacterial enzyme RppH triggers messenger RNA degradation by 5′ phosphatase removal. Nature 2008; 451;355-8; PMID:18202662; http://dx.doi.org/10.1038/nature06475.
17. Mathy N, Bénard I, Pellegrini O, Daou R, Wen T, Condon C. 5′-to-3′ exoribonucleolytic activity in bacteria: role of RNase J1 in RNA maturation and 5′ stability of mRNA. Cell 2007; 129;681-92; PMID:17512465; http://dx.doi.org/10.1016/j.cell.2007.02.051.
18. Richards J, Liu Q, Pellegrini O, Celenk H, Yao S, Bechhofer DH, et al. An RNA pyrophospho- hydrolase triggers 5′-exonucleolytic degradation of mRNA in Bacillus subtilis. Mol Cell 2011; 43;940-9; PMID:21925382; http://dx.doi.org/10.1016/j.molcel.2011.07.023.
19. Li de la Sierra-Gallay I, Zig L, Jamalli A, Putzer H. Structural insights into the dual activity of RNase J. Nat Struct Mol Biol 2009; 15;206-12; PMID:18204464; http://dx.doi.org/10.1038/nsmb.1376.
20. Condon C. What is the role of RNase J in mRNA turnover? RNA Biol 2010; 7;316-21; PMID:20458164; http://dx.doi.org/10.4161/rna.7.3.11193.
21. Durand S, Richard G, Bonnem F, Uzan M. Bacteriophage T4 polynucleotide kinase triggers degradation of mRNAs. Proc Natl Acad Sci USA 2012; 109;7073-8; PMID:22499790; http://dx.doi. org/10.1073/pnas.1119802109.
22. Joanny G, Le Derout J, Bréchemier-Baey D, Labas V, Vinh J, Réguier P, et al. Polyadenylation of a functional mRNA controls gene expression in Escherichia coli. Nucleic Acids Res 2007; 35;2494- 502; PMID:17395638; http://dx.doi.org/10.1093/nar/gkm120.
23. Xu F, Lin-Chao S, Cohen SN. The Escherichia coli polB gene promotes adenylylation of antisense tRNA of ColE1-type plasmids in vivo and degradation of RNAI decay intermediates. Proc Natl Acad Sci USA 1993; 90;6756-60; PMID:7688127; http://dx.doi.org/10.1073/pnas.90.14.6756.
24. Hajnolstor E, Braun E, Haugel-Nielsen J, Réguier P. Polyadenylation destabilizes the rpoP mRNA of Escherichia coli. Proc Natl Acad Sci USA 1995; 92;973-7; PMID:7732015; http://dx.doi.org/10.1073/pnas.92.9.9737.
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