

rRNA Mimicry in RNA Regulation of Gene Expression

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ABSTRACT The rRNA is the largest and most abundant RNA in bacterial and archaeal cells. It is also one of the best-characterized RNAs in terms of its structural motifs and sequence variation. Production of ribosome components including >50 ribosomal proteins (r-proteins) consumes significant cellular resources. Thus, RNA *cis*-regulatory structures that interact with r-proteins to repress further r-protein synthesis play an important role in maintaining appropriate stoichiometry between r-proteins and rRNA. Classically, such mRNA structures were thought to directly mimic the rRNA. However, more than 30 years of research has demonstrated that a variety of different recognition and regulatory paradigms are present. This review will demonstrate how structural mimicry between the rRNA and mRNA *cis*-regulatory structures may take many different forms. The collection of mRNA structures that interact with r-proteins to regulate r-protein operons are best characterized in *Escherichia coli*, but are increasingly found within species from nearly all phyla of bacteria and several archaea. Furthermore, they represent a unique opportunity to assess the plasticity of RNA structure in the context of RNA-protein interactions. The binding determinants imposed by r-proteins to allow regulation can be fulfilled in many ways. Some r-protein-interacting mRNAs are immediately obvious as rRNA mimics from primary sequence similarity, others are identifiable only after secondary or tertiary structure determination, and some show no obvious similarity. In addition, across different bacterial species a host of different mechanisms of action have been characterized, showing that there is no simple one-size-fits-all solution.

Despite the many roles for RNA as a regulator in eukaryotes, archaea, and bacteria, the rRNA is the most abundant cellular RNA and the size of the rRNA outstrips nearly all other functional RNAs. Furthermore, the ribosome is also composed of >50 ribosomal proteins (r-proteins), the majority of which directly contact the rRNA, forming specific interactions with RNA (1).

Since most regulatory RNAs in bacteria appear to be relatively recent inventions (2–5), they most certainly have evolved in the context of abundant rRNA and r-proteins, and thus have been shaped by them. Many regulatory RNA structures contain portions that bear strong resemblance to motifs within the rRNA. Some of this similarity is due to the role that rRNA plays in our understanding of RNA structure, and in other cases it is due to interaction with an r-protein. This review will first illustrate the role of the ribosome in our understanding of RNA structures generally and subsequently examine how r-proteins may interact with RNA outside the ribosome to act in a regulatory capacity.

THE rRNA AS A SOURCE OF RNA STRUCTURAL MOTIFS

The rRNA plays an outsized role in our general understanding of RNA structure. Despite more than a decade since publication of the initial high-resolution ribosome structures and significant growth in the number and diversity of RNA structures in the Protein Data Bank, the rRNA still represents a significant proportion of the three-dimensional structure information available for RNA and RNA-protein complexes. Of the 3,692

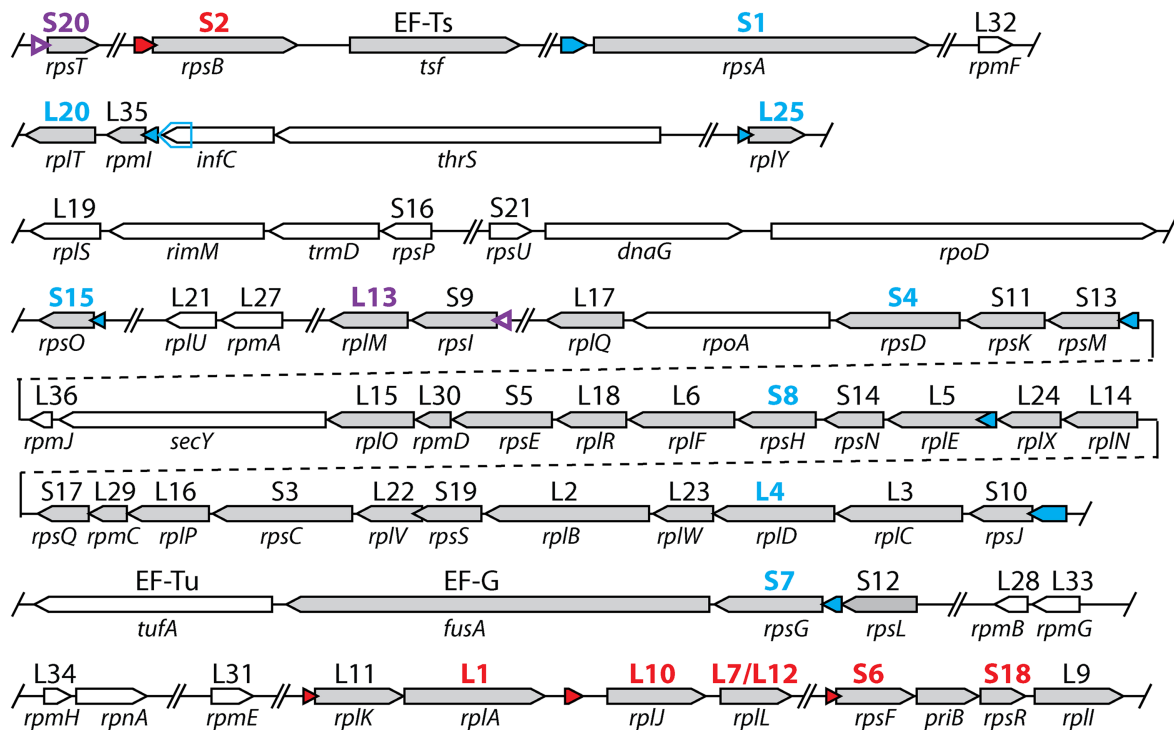
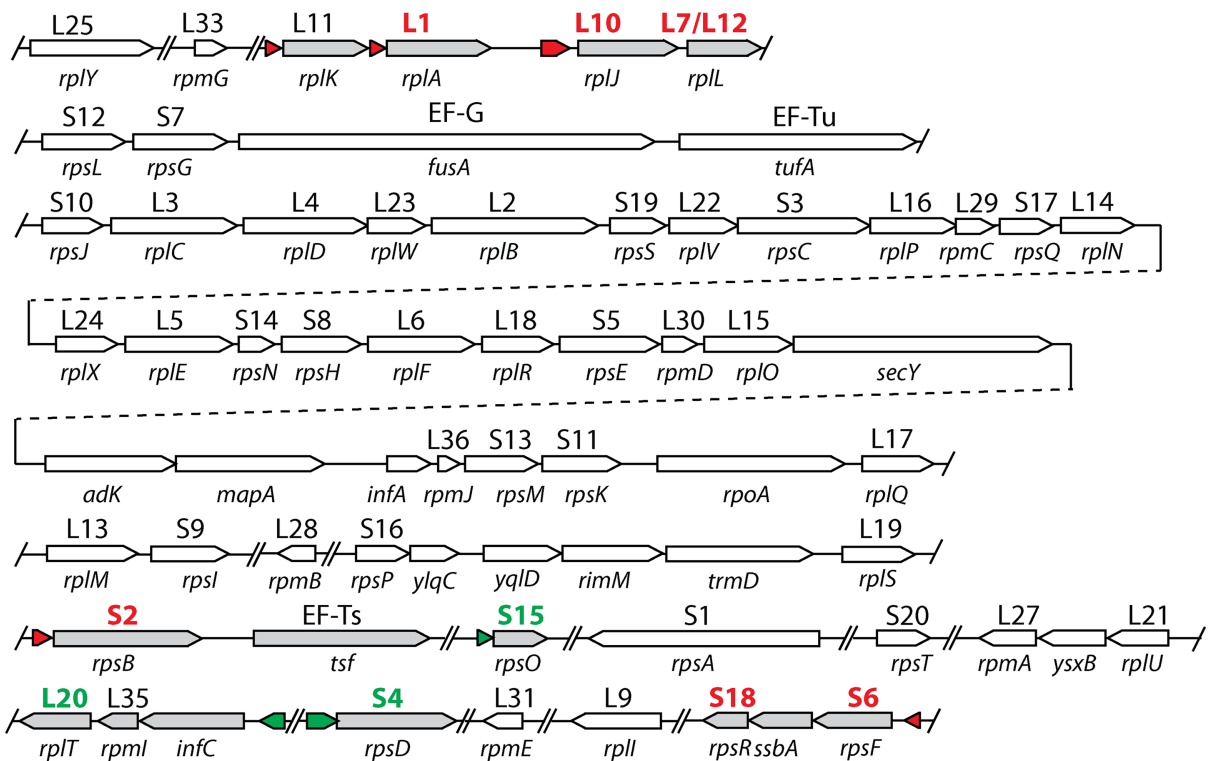
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structures containing RNA, 1,082 contain segments derived from the rRNA or otherwise associated with the ribosome. The ribosome has also significantly influenced the development of RNA structure descriptions (6). Many recurring RNA structure motifs, such as kink-turn (k-turn), loop-E, and loop-C motifs (7–11), were first recognized in the context of the ribosome, and our knowledge of the sequences that may fold into many such features is heavily influenced by rRNA alignments (12–14). These structural motifs form the basis of not only the rRNA but many other structured RNAs including riboswitches (15), T boxes (16), as well as other catalytic RNAs such as the group I and II introns (17, 18). Several reviews specifically addressing the roles such motifs play in RNA structure are available (19–21).

r-PROTEINS AS AUTOGENOUS REGULATORS

Many r-proteins have secondary functions (22, 23) as negative regulators of their own synthesis. r-Proteins and other protein components necessary for translation can account for up to 40% of cellular proteins (24) and 41% of active translation in actively growing cells in rich medium (25). Thus, maintaining stoichiometry among the >60 ribosome components is essential for efficient resource utilization, and the mRNA structures responsible for implementing regulation are only one of several regulatory layers. In *Escherichia coli*, over half of the r-protein operons are regulated by autogenous regulatory mechanisms where an individual r-protein will bind to a portion of its own transcript to inhibit transcription or translation. Often the mRNA will take a structure that bears significant similarity to the rRNA; however, there are several different paradigms for RNA-protein recognition that are embodied by the mRNA structures that mediate r-protein autogenous regulation.

Discovery of r-Protein Autogenous Regulatory mRNA Structures

The mRNA structures enabling regulation of r-protein synthesis in *E. coli* were among the first mRNA regula-

tory sites discovered. Many distinct *E. coli* examples were described based on similar observations and using the same experimental approaches. Initial studies demonstrated that overexpression of specific r-proteins resulted in inhibited synthesis of entire r-protein operons (26, 27), and that these effects were operon specific (28). Using *in vitro* transcription/translation systems as well as reporter gene assays, the inhibitory properties for several r-proteins including L1, L4, S4, S7, S8, and the L10(L12)₄ complex were uncovered (29–33). Most mechanisms involve inhibition of translation (30, 34–37); however, alterations to the mRNA decay rate (27, 38–40) and attenuation (premature transcription termination) mechanisms also occur in conjunction with translational inhibition (41, 42).

In many cases, mimicry between the mRNA regulatory sites responding to an r-protein and its rRNA binding site was proposed as soon as a DNA sequence became available (e.g., S4, S7, S8, L1, L4, and L10 [35, 43–46]). However, demonstration of direct RNA-protein contacts that such similarity would imply lagged behind the speculation considerably (47–49). In several cases, proposed similarities were merely the result of sequence gazing, and it has become apparent that the rRNA and mRNA binding sites do not have structural similarity (e.g., S4 and L4) (50, 51). In other cases, the initially observed similarity between the mRNA and rRNA was verified when three-dimensional structural data became available (e.g., S8 and L1) (52, 53).

Since the initial discoveries of r-protein autogenous mRNA structures in *E. coli*, an additional 9 mRNA structures responding to r-proteins (S1, S2, S15, S20, L19, L20 [2 sites], and L25) (54–60) or r-protein complexes (S6:S18) (61–63) have been described in *E. coli*, and today there are a total of 15 r-protein-interacting mRNA structures described in *E. coli* (Fig. 1A; Table 1). Many have been extensively characterized, but for others the mechanisms of action, or even whether a direct RNA-protein interaction occurs, remain undetermined. With some exceptions, the complement of r-proteins and organization of r-protein operons are largely conserved across bacterial species (64, 65). However, many of the

FIGURE 1 Diagrams of r-protein operons from *E. coli* (A) and *B. subtilis* (B). Genes are shown in the order in which they appear in the genome and to scale. Gray genes are subject to r-protein autogenous regulation; white genes have no described autogenous regulation. Colored arrows represent r-protein RNA binding structures. Red arrows indicate structures that are widely distributed to many bacterial phyla, blue arrows indicate RNA structures that are confined to *Gamma*proteobacteria, green arrows indicate RNA structures confined to *Firmicutes*, and purple arrows indicate presumed r-protein binding sites where no explicit RNA secondary structure has been described. For each operon the effector protein is colored to match the RNA site with which it interacts.

TABLE 1 Summary of r-protein-interacting mRNAs that allow regulation of r-protein genes

Binding partner	Regulated genes	Position	Species distribution ^a
L1 ^{b,c,d,e}	<i>rplA</i> , <i>rplK</i> , ^f <i>rplP1</i> ^f	Varied	Archaea/Bacteria
L4 ^{b,c,d,e}	<i>rpsJ</i> , <i>rplC</i> , <i>rplD</i> , <i>rplW</i> , <i>rplB</i> , <i>rpsS</i> , <i>rplV</i> , <i>rpsC</i> , <i>rplP</i> , <i>rpmC</i> , <i>rpsQ</i>	<i>rpsJ</i> 5' UTR	Gammaproteobacteria
L10/L10(L12) ₄ ^{b,c,d,e}	<i>rplJ</i> , <i>rplL</i>	<i>rplJ</i> 5' UTR	Bacteria
L13 ^b	<i>rplM</i> , <i>rpsI</i>	<i>rplM</i> 5' UTR	<i>Escherichia coli</i>
L20 ^{b,c,d,e}	<i>rpmI</i> , <i>rplT</i>	<i>infC</i> 5' UTR	Firmicutes
L20 ^{b,c,d,e}	<i>rpmI</i> , <i>rplT</i>	<i>infC-rpmI</i> intergenic	Gammaproteobacteria
L20 ^{b,c,d,e}	<i>rpmI</i> , <i>rplT</i>	<i>infC</i> coding/ <i>infC-rpmI</i> intergenic	<i>Escherichia coli</i>
L25 ^{b,d}	<i>rplY</i>	<i>rplY</i> 5' UTR	Gammaproteobacteria
S1 ^{b,c,d,e}	<i>rpsA</i>	<i>rpsA</i> 5' UTR	Gammaproteobacteria
S2 ^{b,d}	<i>rpsB</i>	<i>rpsB</i> 5' UTR	Bacteria
S4 ^{b,c,d,e}	<i>rpsM</i> , <i>rpsK</i> , <i>rpsD</i> , <i>rplQ</i>	<i>rpsM</i> 5' UTR	Gammaproteobacteria
S4 ^{b,d}	<i>rpsD</i>	<i>rpsD</i> 5' UTR	Firmicutes
S6:S18 ^{b,c,d}	<i>rpsF</i> , <i>rpsR</i> , <i>rplL</i> ^f	<i>rpsF</i> 5' UTR	Bacteria
S7 ^{b,c,d}	<i>rpsL</i> , <i>rpsG</i> , <i>fusA</i>	<i>rpsL-rpsG</i> intergenic	Gammaproteobacteria
S8 ^{b,c,d,e}	<i>rplN</i> , <i>rplX</i> , <i>rplE</i> , <i>rpsN</i> , <i>rpsH</i> , <i>rplF</i> , <i>rplR</i> , <i>rpsE</i> , <i>rpmD</i> , <i>rplO</i> , <i>secY</i> , <i>rpmJ</i>	<i>rplX-rplE</i> intergenic	Gammaproteobacteria
S15 ^{b,c,d,e}	<i>rpsO</i>	<i>rpsO</i> 5' UTR	Gammaproteobacteria
S15 ^{c,d}	<i>rpsO</i>	<i>rpsO</i> 5' UTR	Firmicutes
S15 ^{c,d}	<i>rpsO</i>	<i>rpsO</i> 5' UTR	<i>Thermus thermophilus</i>
S15 ^{b,c,d}	<i>rpsO</i>	<i>rpsO</i> 5' UTR	Alphaproteobacteria
S20 ^{b,c}	<i>rpsT</i>	<i>rpsT</i> 5' UTR	<i>Escherichia coli</i>

^aWhere a single species is listed for distribution, either no structure is available or no comparative genomic work has been conducted for the RNA and only the species of characterization is given.

^bRegulation demonstrated using *in vitro* transcription/translation system or reporter gene assays.

^cDirect RNA-protein interaction demonstrated *in vitro*.

^dStructure of mRNA binding site characterized.

^eMechanism of action known.

^fMay only be regulated in some species.

structures allowing regulation in *E. coli* are not widely distributed to organisms outside of a few orders of gammaproteobacteria (66–71). Furthermore, most enterobacterial endosymbionts appear to have lost these structures during the course of genome reduction (71, 72). The only organism with significant study of r-protein regulation other than *E. coli* is the Gram-positive model bacterium *Bacillus subtilis*. This organism shares the mRNA binding sites that interact with r-proteins L1, L10, S2, and S6 with *E. coli* (Fig. 1; Table 1), but the other 11 structures known in *E. coli* are not apparent in *B. subtilis* or its relatives. Alternative regulatory structures that respond to S4, L20, and S15 have been described (73–75) (Fig. 1B; Table 1).

With the growing number and diversity of sequenced bacterial genomes, comparative genomics has also proved to be a powerful approach for discovery. The combination of RNA-specific homology search tools (76) and the availability of RNA structural families corresponding to most known r-protein-responsive structures (71, 75, 77) enables accurate annotation of these structures in bacterial genomes. In addition to characterized mRNA structures, hundreds of novel putative *cis*-regulatory

mRNA motifs have been identified in bacterial genomes, many of which are associated with r-proteins or bear resemblance to the rRNA (78–82). The low cost of sequencing has also enabled the direct discovery of regulatory RNAs through comparative transcriptomics (4), 5'-end sequencing (83), and RNA-protein immunoprecipitations (84). However, relatively few such motifs have been experimentally validated.

The S8-Interacting mRNA Structure: A Prototype r-Protein *cis*-Regulatory RNA

The mRNA segment bound by r-protein S8 to regulate the *spc* operon is the prototype mRNA binding motif that embodies all the properties initially hypothesized for all mRNA structures bound by r-proteins. S8 is a primary rRNA binding protein that interacts with the rRNA early during ribosome assembly. The interaction site for S8 on the mRNA is within the intergenic region between *rplX* and *rplE* (encoding L24 and L5) and the coding region of *rplE* (Fig. 1A). S8 inhibits translation of several proteins following the protein binding site (L5, S14, S8, L6, L18, S5, L20, and L15), and there is

evidence that the two genes upstream of the S8 binding region (*rplN* and *rplX*, encoding L14 and L24) are also downregulated in response to S8 due to increased mRNA degradation (40, 85).

The initial observed sequence similarity between the rRNA and mRNA binding sites for S8 extends to shared secondary structure (35, 86–88) and three-dimensional structure (52, 86) (Fig. 2). The S8 binding site consists of an internal loop. The motif centers on two internal Watson-Crick base pairs that are separated from the rest of the pairing element by bulged bases on either side, although many of the base identities are not strongly conserved in the case of the rRNA (Fig. 2B). S8 itself directly contacts the minor groove of the internal loop. Structures of the mRNA and rRNA are directly superimposable (Fig. 2C) (52). The major difference between the rRNA and mRNA binding sites is an additional bulged base in the mRNA structure a few nucleotides away from the S8 recognition sites (orange). While this base decreases binding affinity by about 10-fold, it does not directly interact with S8. Despite a highly conserved rRNA-S8 interface across all bacteria (88–92) and archaea (93), the S8-responsive regulatory RNA structure observed in *E. coli* is narrowly distributed to a few orders of *Gammaproteobacteria* (71). What, if any, regulation occurs in other organisms has not yet been characterized, and the causes of the narrow distribution are unclear. The phylogenetic distribution of the S8-interacting mRNA structure is similar to those of many r-protein mRNA regulators identified in *E. coli*, suggesting that similar selective constraints influenced the evolution of all the regulatory structures. The preponderance of known structures in *E. coli* is likely due to a significant discovery bias. Ribosome assembly and stoichiometry is by far the best studied in *E. coli*. Similar regulators may be present but as of yet unidentified in other bacteria. The narrow distribution displayed by most of the *E. coli* structures makes it more difficult to utilize comparative genomic approaches for discovery, and it is likely that several of the characterized motifs in *E. coli* would not be easily rediscovered using state-of-the-art comparative genomic tools.

The L10(L12)₄-Interacting Regulatory Structure: Homologous Binding Sites, Different Mechanisms of Action

The L10(L12)₄-interacting mRNA structure also represents a mimic of the rRNA (47, 48, 94–96) and participates in the regulation of translation initiation in *E. coli* (34, 97), directly impacting only *rplJ* and *rplL* (Fig. 1A). Sequence similarity between the mRNA and rRNA

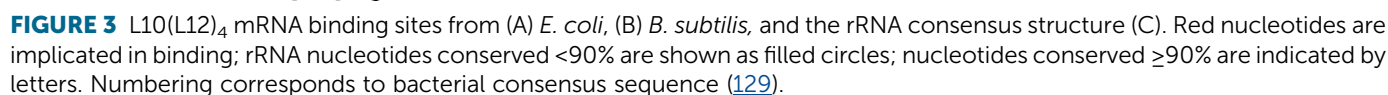
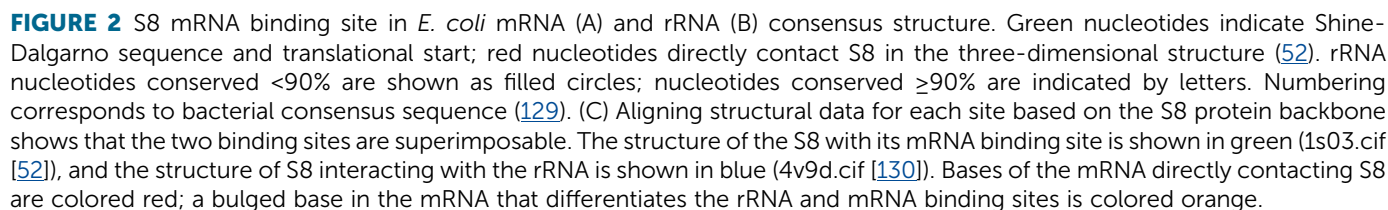
binding sites has been described (46, 68), but the L10 (L12)₄ complex is typically not resolved in ribosome crystal structures and three-dimensional data for an mRNA-L10(L12)₄ complex are not available. The L10 (L12)₄ binding site consists of a k-turn motif that is four base pairs away from an internal loop containing a pair of adenosines. In the rRNA the internal loop is a multi-stem junction (Fig. 3), while in the mRNA the structure is often a bulge, but may be a multi-stem junction (71). In both the rRNA and the mRNA the adenosines are highly conserved, and mutating them reduces binding affinity substantially (68).

In contrast to the S8-interacting mRNA structure, the RNA structure responsible for interacting with L10 in *E. coli* is widely conserved throughout many bacterial species (68, 71, 80). However, the mechanism of action is not the same across all species. In many Gram-positive species, the L10-interacting structure is followed by an intrinsic transcription terminator (80), and the mechanism of regulation in *B. subtilis* is regulation of transcription termination (98). Thus, r-protein binding structures are similar to riboswitches where homologous sensor domains may utilize different mechanisms of action in diverse species (99).

The L1-Interacting mRNA Structure: Convergence on the Same Binding Determinants

Like the L10- and S8-responsive mRNA structures, the L1-interacting mRNA structure shows obvious similarity to the rRNA (45, 53, 100), and examples of the L1 recognition site are found across nearly all bacterial phyla (71) as well as archaea (101–103). The binding determinants for L1 are often accommodated in a short hairpin of <30 nucleotides and consist of a base-paired region containing an asymmetric internal loop closed by a noncanonical A•G pairing (Fig. 4A). In three dimensions this corresponds to two canonical helices, one of which is capped by the noncanonical A•G pair, that are separated by a sharp turn (53). Diverse L1 homologs are able to interact with an example of the mRNA binding site from *Methanococcus vannielii* (103), and structural data show that the rRNA and mRNA sites are nearly superimposable (53).

In *E. coli*, the binding site is within the 5' untranslated region (5' UTR) of the transcript encoding both *rplK* and *rplA* and L1 regulates translation initiation of L11 and L1 (45) (Fig. 4). Surprisingly, in the archaea *M. vannielii*, *Methanocaldococcus jannaschii*, and *Methanococcus thermolithotrophicus*, the L1 binding site appears ~30 nucleotides inside the coding region for L1 and regulates



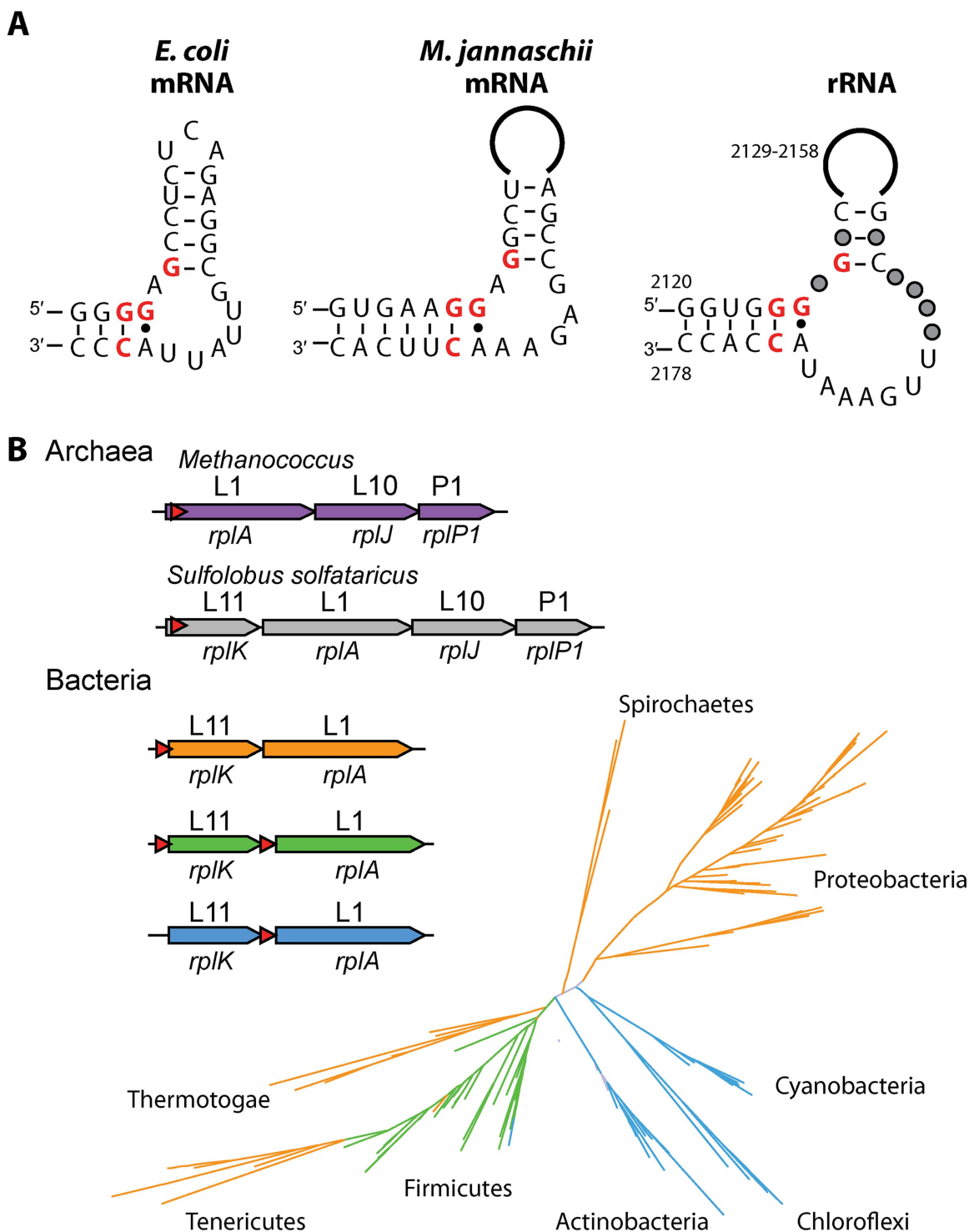


FIGURE 4 (A) L1-interacting mRNA structures from *E. coli* and *M. jannaschii*, and the L1 rRNA binding site (bacterial consensus). Red nucleotides directly contact L1 in the three-dimensional structure (53). rRNA nucleotides conserved <90% are shown as filled circles; nucleotides conserved ≥90% are indicated by letters. Numbering corresponds to bacterial consensus sequence (129). (B) Diagrams indicating the genomic position of L1 mRNA binding sites in two Archaea clades (several *Methanococcus* species and *Sulfolobus solfataricus*) and in bacteria. Proteins encoded by *rplJ* and *rplP0* are homologous. Bacterial genomic positions of L1 binding site are mapped to a 16S rRNA tree.

translation of L1, L10, and P1 (homolog of L12 [104]), and the gene encoding L11 occurs elsewhere in the genome (101, 103). In *Sulfolobus solfataricus*, the L1 binding site is found within the L11 coding region, which directly precedes genes encoding L1, L10, and P1 (Fig. 4B) (103).

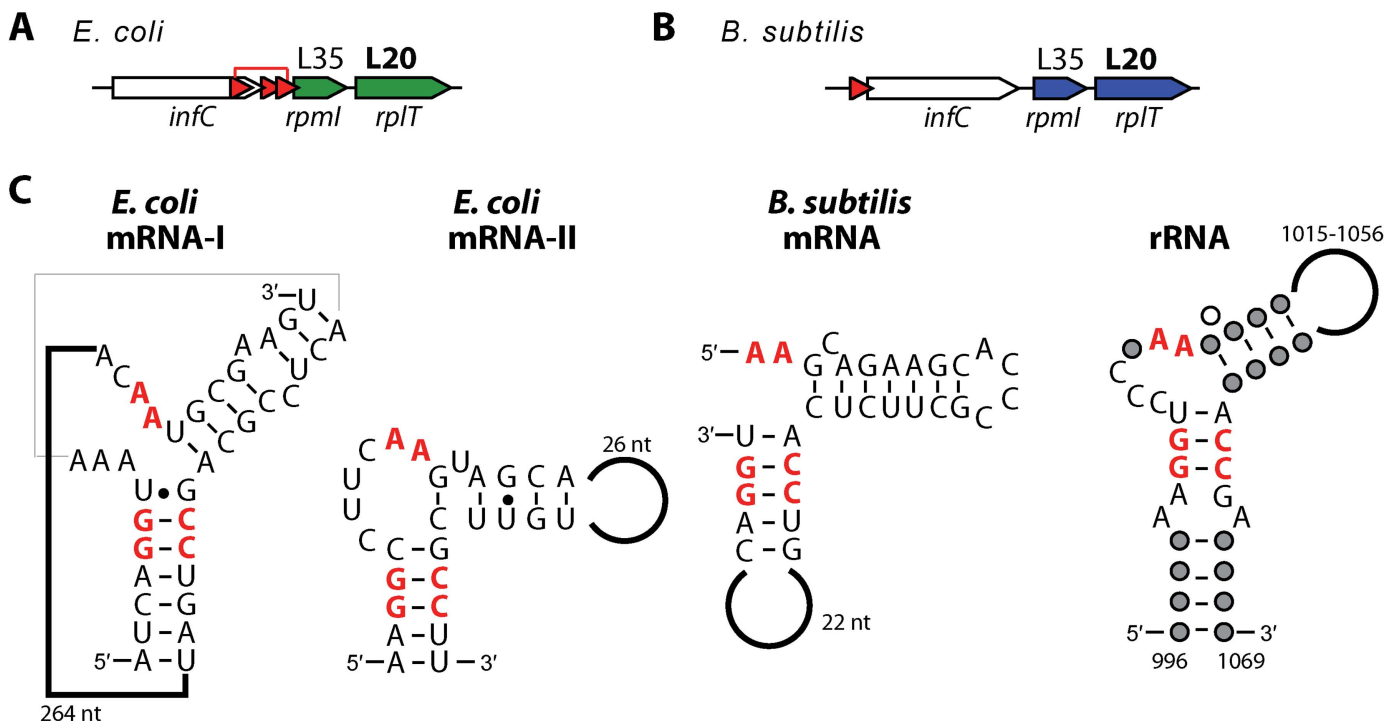
In addition to examples that have been explicitly examined, a systematic homology search for L1 binding sites in bacterial genomes identified the site within transcripts encoding L1 and L11 in many bacterial species (71). However, like the examples identified in *Archaea*, the location of the binding site relative to the coding regions is not consistent. In *Cyanobacteria*, *Actinobacteria*, and *Chloroflexi*, the L1 binding site precedes *rplA*, typically between *rplK* and *rplA*. In *Proteobacteria*, *Spirochaetes*, *Thermotogae*, and *Tenericutes*, the binding site precedes *rplK*, presumably to regulate both *rplA* and *rplK*. Furthermore, in many species of *Firmicutes*, L1 binding sites appear preceding both *rplA* and *rplK*. In *Geobacillus kaustophilus*, both sites are capable of binding L1 *in vitro* (71). Interestingly, there is evidence

of loss for each individual binding site within species scattered throughout *Firmicutes* (Fig. 4B). The combination of the wide distribution and changing position of the L1 binding site relative to the regulated genes suggests that the site may have evolved convergently in many species.

L20-Interacting mRNA Regulatory Structures: Diverse Scaffolds Support the Same Binding Determinants

In addition to cases where there is a single mRNA binding site that mimics the rRNA, there are also cases where homologous r-proteins interact with distinct mRNA secondary structures in different bacterial species. Three L20-interacting mRNA structures are known, two in *E. coli* and one in *B. subtilis*. Each structure mimics the rRNA, but uses a different arrangement of secondary structure to support the necessary bases in the correct geometry required for recognition (Fig. 5). In *E. coli*, two L20-responsive mRNA structures control the IF3 operon (*infC*, *rpmI*, and *rplT*, encoding IF3, L35, and

FIGURE 5 Diagram of *infC* operons showing genomic positions of L20-interacting mRNA structures (red arrows) in *E. coli* (A) and *B. subtilis* (B). Genes regulated by the RNA structure are colored. (C) L20-interacting mRNA structures from *E. coli* (mRNA-I and mRNA-II) and *B. subtilis* and the consensus rRNA L20 binding site. Red nucleotides are important for L20 interaction. rRNA nucleotides conserved <90% are shown as filled circles; nucleotides conserved ≥90% are indicated by letters. Numbering corresponds to bacterial consensus sequence (129).



L20). One structure is found within the intergenic region between *infC* and *rpmI* (70) (Fig. 5A), and consists of a relatively straightforward bulged stem-loop where the binding site includes a pair of adenosines within the bulge and a set of consecutive G-C base pairs just after the closing base pair of the loop (Fig. 5C, mRNA-II). This arrangement is the closest mimic of the rRNA. The second structure comprises a pseudoknot formed by long-range interactions between a sequence within *infC* and a sequence adjacent to the start of the *rpmI* coding region (54, 70) (Fig. 5C, mRNA-I). In this structure, the pair of adenosines is found in the single-stranded region just prior to the 3'-most portion of the pseudoknot. Both of these structures are required for full translational repression of the operon *in vivo*, and L20 binds independently to each (70, 105). A high-quality alignment and phylogenetic distribution is only available for the mRNA structure preceding *rpmI*. The pseudoknotted binding site is challenging to identify using RNA-specific homology search tools (106) due to its significant overlap with coding sequence, long-range interactions, and pseudoknotted structure. However, the structure preceding *rpmI* is narrowly distributed to *Gammaproteobacteria*.

In addition to the two L20-responsive structures in *E. coli*, L20 binds to a regulatory structure in *B. subtilis* that precedes *infC* (Fig. 5B and C). While this structure shares many features with the L20-interacting structures identified in *E. coli*, the binding features present near the multi-stem junction are supported by a different arrangement of secondary structure, and the ordering of the elements with respect to one another in a linear sequence is distinct. A potential intrinsic transcription terminator follows this mRNA structure, and the mechanism of action is L20-induced structural change resulting in early transcription termination (74). In this structure, the conserved adenosines are in a single-stranded region just 5' of the first hairpin and the G-C pairs in the second hairpin. This structure is found in most *Firmicutes* (75, 80), although more frequently in the class *Bacilli* than in *Clostridia*, and the transcription attenuation mechanism appears conserved in these organisms. While *infC* is part of this operon, in *B. subtilis* IF3 levels are decoupled from those of L20 and L35 through the presence of a second upstream promoter. The transcript produced from this promoter is cleaved by RNase Y and only allows translation of L20 and L35 translation (107).

The three L20 sites all present the same effective binding geometry for L20 recognition (Fig. 5C). Both *E. coli* structures are capable of interacting with an L20

homolog from *Aquifex aeolicus* to repress gene expression from *rpmI*'-'*lacZ* reporter constructs (108), and the *E. coli* homolog is able to stimulate premature transcription termination during *in vitro* assays with the *B. subtilis* mRNA structure (74). Thus, the L20 regulators serve as an example of how the same three-dimensional geometry may be supported by different arrangements of secondary structure elements. This example also illuminates how challenging identification of common binding sites may be. Despite similar binding determinants, the distinct arrangements of the necessary recognition elements make automatic detection difficult or impossible.

S15-Interacting Regulatory Structures: Diverse Binding Determinants Produce Diverse Structures

r-Protein S15 also regulates gene expression using multiple distinct mRNA binding sites in diverse bacterial species. To date, four different S15-interacting mRNA structures spanning several bacterial phyla have been experimentally characterized (109–112) and several additional putative structures identified (112) (Fig. 6). Each of these structures directly precedes and controls expression of *rpsO*, the gene encoding S15. In *E. coli* the mechanism of action is through entrapment of the translation initiation complex (110), but in other species the mechanism has not explicitly been characterized.

The structures share very little in the way of a single recognizable sequence or structural motif. This is partially due to the bidentate nature of the S15 binding site on the rRNA. S15 recognizes two portions of the 16S rRNA: a multi-stem junction and a stem containing a slight defect characterized by a G•U/G-C set of base pairs directly adjacent to the junction (Fig. 6). The mRNA regulatory structures that interact with S15 often only partially mimic this binding site. For example, the mRNA from *Thermus thermophilus* includes a three-stem junction formed by the bases of three adjacent pairing elements. However, the pairing elements themselves show no evidence for the G•U/G-C defect recognized by S15 in the rRNA. In contrast, mRNA structures described from *E. coli*, *Rhizobium radiobacter* (formerly *Agrobacterium radiobacter*), and *Geobacillus stearothermophilus* (previously *Bacillus stearothermophilus*) include more obvious mimics of the G•U/G-C elements, and require this element for interaction. In several cases the mRNAs have additional recognition elements that are necessary but do not directly mimic the rRNA (Fig. 6) (109, 113, 114).

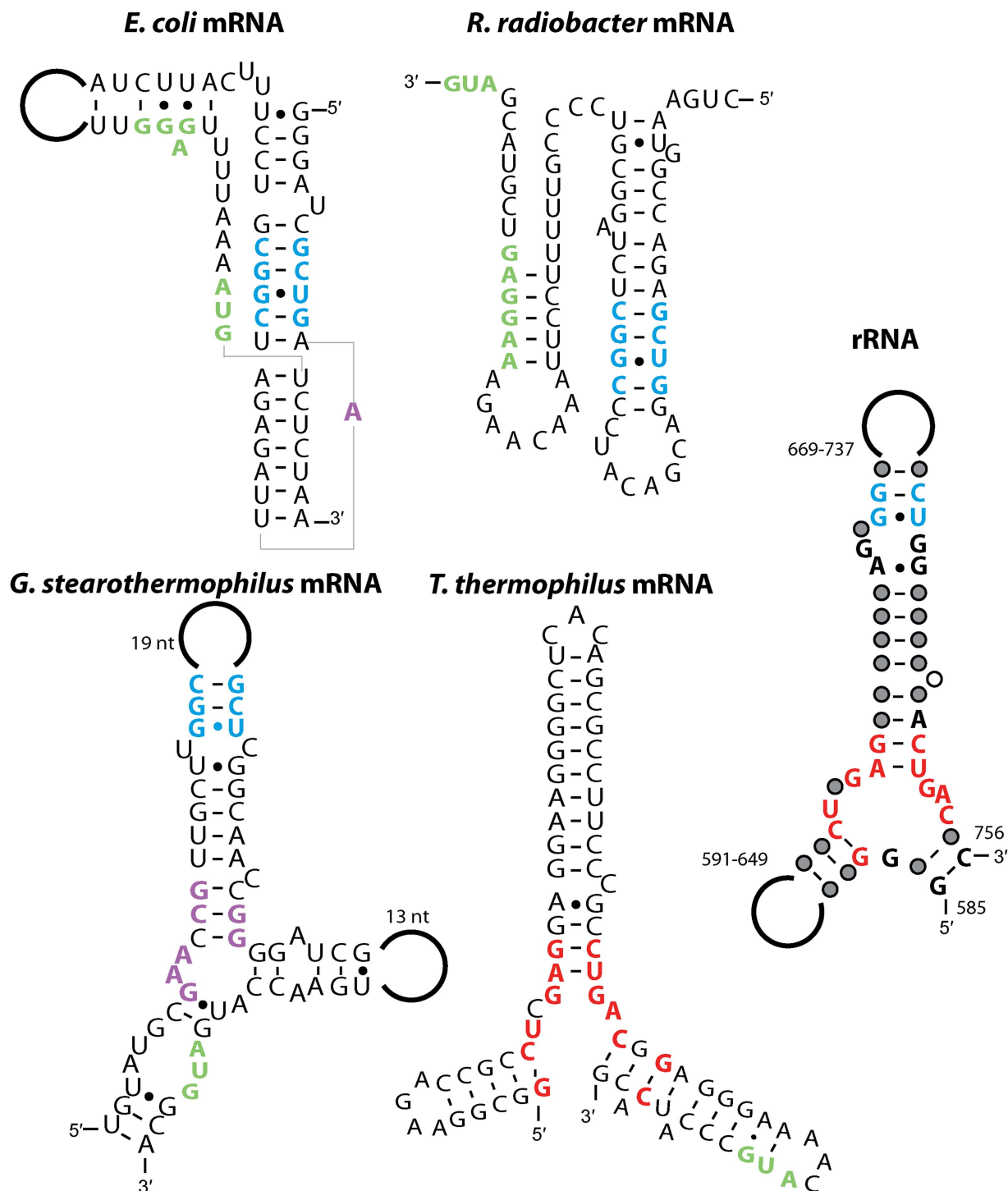


FIGURE 6 S15-interacting mRNA structures in different bacterial phyla and the consensus S15 rRNA binding site. Red nucleotides correspond to the rRNA three-stem junction and its direct mimics. Blue nucleotides correspond to G•U/G-C helix imperfection in the rRNA binding site and its mimics in mRNA structures. Purple nucleotides are important for S15 recognition but do not directly correspond to any rRNA motif. Green nucleotides correspond to Shine-Dalgarno or translational start sequences. rRNA nucleotides conserved <90% are shown as filled circles; nucleotides conserved ≥90% are indicated by letters. Numbering corresponds to bacterial consensus sequence (129).

The differences between the mRNAs are sufficiently large such that specificity of interaction has been reported (113, 115). For example, the S15 homolog from *G. kaustophilus* does not interact with the mRNA structure originating from *E. coli* and the S15 homolog from *T. thermophilus* does not interact with several of the mRNA structures containing only the G•U/G-C motif and no mimic of the three-stem junction (113, 115). Mutagenesis studies indicate that the same face of S15 appears to be used for interaction (115, 116). However, in *E. coli* different S15 amino acids are implicated in rRNA and mRNA binding (114, 117). Furthermore, selective recognition of the *Geobacillus* and *E. coli* mRNA structures may be traced to specific amino acids that are differentially conserved in S15 homologs originating from organisms containing RNAs of each type (113). Thus, the diversity of S15-interacting structures is due not only to the bidentate recognition site that may allow a larger set of potential interaction partners but also to differences in the protein homologs that change recognition. These findings suggest that despite very similar rRNA recognition sites across all bacteria, the r-proteins and their mRNA binding sites are influencing each other's evolution.

BEYOND AUTOGENOUS REGULATION: L7Ae

Archaeal r-protein L7Ae participates in processes well beyond its role in the ribosome. L7Ae interacts with k-turn and k-loop motifs as a component of the ribosome (1), RNase P (118), the C/D box and H/ACA box snoRNPs responsible for site-selective 2'-O-methylation (119, 120), and in mammals an L7Ae homolog binds to the U4 snRNP of the spliceosome (121). There is no r-protein that directly corresponds to L7Ae in prokaryotes. Two L7Ae homologs in *B. subtilis* exist, and both bind to k-turns (122), but their biological function is unknown. The role of the k-turn as a fundamental RNA structural building block has already been discussed. L7Ae specifically recognizes this motif, and therefore has a role in many RNA complexes, primarily to stabilize RNA structure.

A recent RNA immunoprecipitation sequencing (RIP-seq) study of L7Ae in *Sulfolobus acidocaldarius* identified several mRNA fragments in addition to the expected interaction partners (84). Many of these mRNA fragments contained sequences corresponding to the consensus sequences for a k-turn, suggesting a biologically relevant interaction. Among the mRNAs identified are those encoding L7Ae, Nop5, and fibrillarin (other components of the snoRNP), a hypothetical DNA binding protein, and a hypothetical glycosyltransferase. Subse-

quent reporter gene assays and phylogenetic analysis showed that L7Ae negatively regulates the transcript encoding L7Ae in *S. acidocaldarius* and several other diverse *Archaea* species. The presence of k-turn motifs preceding several genes and L7Ae interaction with these motifs suggests that L7Ae may regulate not only its own synthesis but also synthesis of its interaction partners in snoRNPs, Nop5 and fibrillarin (84).

ENGINEERED r-PROTEIN-RESPONSIVE REGULATORY RNA SYSTEMS

r-Protein binding motifs have also been used to create synthetic regulatory systems. Repressing systems designed for eukaryotic cells were created by placing the L7Ae binding site near the translational start site (123), allowing L7Ae to prevent translation initiation. Systems with the L7Ae binding site within the coding region proved more effective than those where the binding site was placed in the 5' UTR, both in an *in vitro* translation system and within HeLa cells. Activating systems in which L7Ae binding removes a *trans*-acting RNA to prevent translation also proved effective *in vitro*. In addition, L7Ae-mediated activation was achieved in HeLa cells by adding an L7Ae binding site to a synthetic shRNA (short hairpin RNA); thus L7Ae binding prevented shRNA-mediated mRNA degradation (124). These examples demonstrate how the L7Ae protein binding site may be easily transferred to an alternative context and harnessed for gene expression in a modular manner.

Indeed, creation of synthetic regulatory systems responding to r-proteins within cells appears to be facile in comparison to the creation of many types of synthetic regulators where the transition from *in vitro* to *in vivo* can be challenging (125). Several synthetic regulatory systems responding to r-protein S15 have also been created (126). Unlike the L7Ae examples, these regulators were created through *in vitro* selection of RNA aptamers interacting with r-protein S15 from *G. kaustophilus* rather than transplantation of a known binding site. One striking observation from this work is that even without explicit selection for regulation, a high proportion of aptamers enable regulation when positioned correctly relative to the start codon. A second finding is that r-protein S15 can interact with a wide variety of different binding sites (127). This observation is echoed in previous work where *in vitro* selection to r-protein S8 yielded both aptamers similar to the natural RNA binding partners as well as those showing substantial differences (128).

CONCLUDING REMARKS

Regulatory RNA structures displaying motifs found in the rRNA are commonly identified. While in some cases similarity may be due to shared RNA tertiary structure motifs, in other cases structural similarity can imply a shared r-protein binding partner. Many r-proteins have a secondary role as negative regulators of their own synthesis, and while it was postulated that all such regulatory structures would resemble the rRNA, this has proved true only in some cases. This review illustrates a range of different regulatory mRNA structures that display similarity to the rRNA, but it is by no means exhaustive. While the mRNA structures controlling r-protein synthesis in *E. coli* remain the best characterized, r-protein-responsive mRNA structures hail from nearly all species of bacteria and several archaea. From these examples it is apparent that r-protein-responsive mRNA structures can be direct and obvious mimics of the rRNA, but they do not have to be. Many r-protein-interacting mRNA structures display no similarity to their cognate rRNA sites (e.g., *E. coli* S4 regulator), while others share only partial similarity. Second, very similar binding sites can appear in diverse organisms, but they may use alternative mechanisms to regulate gene expression or display different positioning relative to regulated genes. Third, due to the structural plasticity of RNA, a geometrically similar binding site may be displayed in several very different manners. Finally, from the diversity of natural regulatory mRNA structures, *in vitro* selection of aptamers, and design of r-protein-responsive regulatory mechanisms, it is clear that the sequence space that allows for r-protein binding and subsequent gene regulation may be quite large. This conclusion combined with the lack of knowledge of r-protein regulation outside of *E. coli* suggests that many r-protein-responsive mRNA structures, including those not directly associated with r-protein operons, remain undiscovered or unverified.

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